No Effects of Cognitive Remediation on Cerebral White Matter in Individuals at Ultra-High Risk for Psychosis—A Randomized Clinical Trial

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Background: Individuals at ultra-high risk for psychosis (UHR) present with subtle alterations in cerebral white matter (WM), which appear to be associated with clinical and functional outcome. The effect of cognitive remediation on WM organization in UHR individuals has not been investigated previously.

Methods: In a randomized, clinical trial, UHR individuals aged 18 to 40 years were assigned to treatment as usual (TAU) or TAU plus cognitive remediation for 20 weeks. Cognitive remediation comprised 20 x 2-h sessions of neurocognitive and social-cognitive training. Primary outcome was whole brain fractional anisotropy derived from diffusion weighted imaging, statistically tested as an interaction between timepoint and treatment group. Secondary outcomes were restricted to five predefined region of interest (ROI) analyses on fractional anisotropy, axial diffusivity, radial diffusivity and mean diffusivity. For significant timepoint and treatment group interactions within these five ROIs, we explored associations between longitudinal changes in WM and cognitive functions/clinical symptoms. Finally, we explored dose-response effects of cognitive remediation on WM.

Results: A total of 111 UHR individuals were included. Attrition-rate was 26%. The cognitive remediation group completed on average 12 h of neurocognitive training, which was considerably lower than per protocol. We found no effect of cognitive remediation on whole-brain FA when compared to treatment as usual. Secondary ROI analyses revealed a nominal significant interaction between timepoint*treatment of AD in left medial...
leminiscus (P=0.016) which did not survive control for multiple comparisons. The exploratory test showed that this change in AD correlated to improvements of mental flexibility in the cognitive remediation group (p=0.001). We found no dose-response effect of neurocognitive training on WM.

**Conclusions:** Cognitive remediation comprising 12 h of neurocognitive training on average did not improve global or regional WM organization in UHR individuals. Further investigations of duration and intensity of cognitive training as necessary prerequisites of neuroplasticity-based changes are warranted.

**Clinical Trial Registration:** ClinicalTrials.gov, identifier NCT02098408.

**Keywords:** ultra-high risk for psychosis, cognition, white matter, clinical trial, cognitive remediation, diffusion-weighted imaging

**INTRODUCTION**

Individuals at ultra-high risk (UHR) for psychosis present with cognitive impairments (1), intermediate between the marked cognitive deficits observed in patients with manifest psychotic disorders, and unaffected healthy controls (2–4). The cognitive impairments in UHR individuals are present across multiple domains (5), with small to large effect sizes compared to healthy controls (6).

Moreover, magnetic resonance imaging studies have revealed widespread, but subtle alterations in WM organization in UHR individuals, typically measured as lower fractional anisotropy (7). Fractional anisotropy (FA) is the currently most widely applied index of WM organization (8), sensitive to a broad range of neurobiological substrates, such as axonal density, axonal cross-section, myelin and crossing fibers (9). These WM-alterations have been associated with severity of cognitive deficits (10) and functional outcome (11), and appear to convey a liability for transition to frank psychosis (12). In a previous cross-sectional study of the current sample (10), we performed multivariate analyses of baseline data on associations between WM and cognition, comparing UHR individuals to healthy controls. At a whole-brain level, we identified an association between globally higher fractional anisotropy with a pattern of better cognitive functions associated with WM-neuroplasticity as a result of cognitive interventions in psychiatric patients, and to our knowledge none in UHR individuals. Cognitive remediation in UHR individuals is of particular interest, since these individuals may hold greater potential of brain plasticity, compared to patients with manifest psychotic disorders (19).

Cognitive remediation is a structured method of cognitive training, which is defined as “a behavioral training-based intervention that aims at improving cognitive processes (attention, memory, executive function, social cognition or metacognition) with the goal of durability and generalisation” (20). Thus, cognitive remediation targets selected cognitive domains and explicitly aims to improve social and neurocognitive functions in a manner that impacts real life functioning (21). There is meta-analytic evidence for the effectiveness of cognitive remediation in schizophrenia, with medium effect-sizes on global cognition and functional outcome (22), and the effects appear durable (20). Cognitive remediation is emphasized as an enhancing supplement to standard treatment, more effective when embedded in a broader psychiatric rehabilitation program (20). Studies are sparse in UHR-populations. A recent systematic review (23), examining six studies on cognitive remediation with UHR individuals provided some preliminary evidence for the effectiveness of cognitive remediation on cognition [e.g improvement of verbal memory (ES 0.61) (24) and processing speed (ES 0.50) (25)], and aspects of functional outcome. However, methodological considerations were raised, such as low sample sizes and high attrition rates, precluding any solid conclusions to be drawn.

In this randomized clinical trial (Function and Overall Cognition in Ultra-high risk States, the FOCUS-trial), we aimed to evaluate the effect of comprehensive neurocognitive and social cognitive remediation in UHR individuals on cerebral WM organization, as assessed by fractional anisotropy. We hypothesized, that 20 weeks of...
cognitive remediation in addition to treatment as usual (TAU) would increase whole-brain fractional anisotropy in UHR individuals allocated to the intervention, when compared to UHR individuals allocated to TAU. Second, we investigated the effects of cognitive remediation on FA and additional WM-measures in a priori ROIs. The regions were selected as the five regions our previous study had shown differentially associated to cognitive functions in UHR individuals. Next, we planned exploratory correlation-analyses between changes in WM-measures in the predefined regions, which showed significant interaction-effects on time×group and changes in clinical symptoms and cognitive functions. We expected correlations between changes in regional WM and changes in clinical symptoms/cognitive functions to be different, when comparing UHR individuals receiving cognitive remediation versus TAU. Finally, we explored a dose-response effect of cognitive remediation on WM.

METHODS

The current study is a sub study of a randomized, assessor-blinded, parallel-group, superiority clinical trial comparing TAU plus 20 weeks of intensive cognitive remediation with TAU (the FOCUS-trial, registered at ClinicalTrials.gov NCT 02098408). The main trial protocol has been published (26), and the implications of the main results have been discussed in a separate paper (27). The study was carried out at the Mental Health Centre Copenhagen and Mental Health Centre Glostrup, University of Copenhagen, Denmark. Baseline data on the relationship between cognition and WM from the FOCUS-trial have been published elsewhere (10). Here, we analyzed longitudinal data from UHR individuals at two timepoints, pre-treatment (i.e. at baseline) and post-treatment (i.e. at 26-weeks follow-up). All participants provided informed consent prior to inclusion into the study. The study protocol was approved by the Committee on Health Research Ethics of the Capital Region Denmark (study: H-6-2013-015) and the Danish Data Protection Agency (RHP-2014-009-02670).

Procedures

Following baseline assessments, participants were randomly assigned to TAU or TAU plus cognitive remediation for 20 weeks. The randomisation was centralised and computerised with a concealed randomisation sequence carried out by the Copenhagen Trial Unit (CTU). Randomisation was stratified by current use of antipsychotic medication (yes/no) and IQ score (≤100/>100). Block size was blinded. Group allocation was concealed until the statistical analyses of the data had been completed. Investigators blinded to group allocation conducted assessments at 26-weeks (post-treatment) at a site remote from the intervention. Participants were instructed not to disclose their allocation prior to assessments. In case of failure of concealment, the assessment would be conducted by another research assessor. Assessors were psychologists and medical doctors with extensive training in the assessment instruments. Efficacy-analyses on WM were conducted by a blinded researcher.

UHR Individuals

UHR individuals were recruited from psychiatric in- and outpatient facilities in the catchment area of Copenhagen. Individuals were help-seeking individuals aged 18 to 40 years, who fulfilled one or more of the UHR criteria as assessed by the Comprehensive Assessment of At-Risk Mental State (CAARMS) (28): attenuated psychotic symptoms, and/or brief limited intermittent psychotic symptoms, and/or trait and vulnerability state along with a significant drop or sustained low functioning for the past year. Exclusion criteria were a history with a psychotic episode of more than one week of duration; psychiatric symptoms which were explained by a physical illness with psychotropic effect or acute intoxication; a diagnosis of a serious developmental disorder, or currently receiving methylphenidate.

Assessments

The CAARMS interview and The Structured Clinical Interview for DSM-IV Axis I Disorders (SCID-I) and part of the Structured Clinical Interview for DSM-IV Axis II Disorders (SCID-II) (29, 30) were used to diagnostically assess all UHR individuals. Level of general psychiatric symptoms was measured with the Brief Psychiatric Rating Scale Expanded Version (BPRS) (31), the level of negative symptoms was measured with the Scale for the Assessment of Negative Symptoms (SANS) (32), and the level of depressive symptoms was measured with the Montgomery-Åsberg Depression Rating Scale (MADRS) (33). Inter-rater reliability was assessed using intra-class correlations for the outcome measure SANS, BPRS, and MADRS in 12 interviews (ICC ratings from 0.96 to 0.99).

The third version of the Danish Weschler Adult Intelligence Scale (WAIS-III) (34) provided estimates of current level of intelligence. Cognitive functions were assessed using selected tests from the Brief Assessment of Cognition in Schizophrenia battery (BACS) (35): list-learning, digit sequencing, verbal fluency, and symbol coding, as well as tests from the Cambridge Neuropsychological Test Automated Battery (CANTAB) (36): Stockings of Cambridge (SOC Problems solved in minimum moves) and Intra-extradimensional set shifting test (IED Total errors adjusted, lower is better). For a detailed overview on cognitive domains and tests, see Supplementary Table S1.

Image Acquisition and Processing

MRI scans were acquired on a 3 Tesla scanner (Philips Healthcare, Best, the Netherlands). Diffusion-weighted images (DWIs) were acquired using single shot spin-echo echoplanar imaging (EPI) sequence with 30 noncollinear diffusion-weighted [b = 1,000 s/mm²] directions and one non-diffusion weighted b=0 s/mm². Two DWI scans were acquired, and the latter in an opposite phase encoding direction, enabling correction for susceptibility distortions (37). Details on image acquisition and processing are provided in Supplementary Text S2. Tools from the FSL software library (5.0.10 (38) and Mrtrix3 (www.mrtrix.org) were used for image processing. Motion parameters were extracted to correct for head motion. Maps of fractional anisotropy (FA), axial diffusivity (AD), radial diffusivity (RD) and mean diffusivity (MD) were calculated.
Tract-based spatial statistics (TBSS) (39) was used to create skeleton maps using a threshold of 0.2. Using the JHU DTI-based white matter atlas labels (40), we computed the mean FA values in five WM-ROIs from skeletonized data. ROIs were selected a priori and constituted the five regions where we in our previous cross-sectional study of baseline data had identified aberrant associations between WM and cognitive functions: fornix, medial lemniscus bilaterally, left superior cerebellar peduncle, and left uncinate fasciculus (10). MRI quality metrics were assessed by visual inspection and calculated from each subject using a quality assessment method described in Roalf et al. (41) (Table S3).

**Intervention**

The experimental intervention was delivered by a senior clinical psychologist with a specialization in psychotherapy and consisted of manualized neurocognitive and social cognitive remediation. The integrative approach targeting both neurocognitive and social cognitive deficits was based on the assumption, that neurocognitive and social cognitive remediation may work synergistically to increase the transfer of cognitive remediation gains to participants’ real-world functioning (42).

Duration was 2 h once a week for a total of 20 weeks in a group-setting, combined with individual home-training. In addition to the group training, the participants received 12 individual sessions designed to maximize transfer of the effect of the cognitive training to the daily lives of the participants. The approach in individual sessions were cognitive behavioral therapy and targeted everyday challenges in relation to the UHR individual’s specific cognitive deficits.

The neurocognitive remediation was performed using the Neuropsychological Educational Approach to Cognitive Remediation (NEAR) (43). Exercises from the web-based ScientificBrainTrainingpro.com and supplementary exercises from Brainhq.com were trained individually on computers for 1 h in the group-setting, followed by a group discussion aiming at relating the cognitive exercises to real world activities (bridging). UHR individuals were instructed to train the neurocognitive exercises at home to achieve the recommended dosage of 2 h per week of neurocognitive training (40 h in total). Homework was monitored in the training programs and motivation was addressed ongoing in the individual sessions, by sending text reminders and offering support from the group therapist. The therapist personalized the neurocognitive remediation based on the cognitive domains with the most severe impairments, as assessed by the UHR individuals baseline neurocognitive performance. The social cognitive training was performed using the Social Cognition and Interaction Training (SCIT) manual (44), which addresses several key social-cognitive domains. The therapist had attended a SCIT training course and received ad hoc supervision by Dr. David Roberts, first-author of the SCIT-manual (45). Adherence-rating to the treatment manual was conducted according to the SCIT Fidelity Scale (44) by an external rater.

Both the cognitive remediation group and the control group received TAU, which consisted of the treatment offered by the psychiatric services. TAU involved monitoring of medication, regular contact to primary practitioner delivering supportive counselling, and frequently a more regular psychotherapeutic intervention, such as social skills training, psycho-education, family-groups, supportive group-therapy, or individual therapeutic sessions with a psychologist, but not targeted cognitive remediation.

**Outcomes**

The primary outcome was whole brain fractional anisotropy (FA) as an effect of interaction between timepoint and treatment-group (‘cognitive remediation versus TAU’).

Secondary outcomes were measures of white matter: FA, axial diffusivity (AD), radial diffusivity (RD), and mean diffusivity (MD) in the five ROIs fornix, medial lemniscus bilaterally, left superior cerebellar peduncle, and left uncinate fasciculus, as an effect of interaction between timepoint and treatment-group (‘cognitive remediation versus TAU’).

Planned explorative tests were correlation analyses between WM-changes in the predefined ROIs, which showed significant between-group difference on secondary outcomes, and clinical and cognitive changes as compared between treatment-groups (‘cognitive remediation versus TAU’). Moreover, as motivated by the potential dose-dependent effect of cognitive remediation on WM (17), we post-hoc tested a dose-response relationship between the proportion of cognitive remediation and change in whole-brain and regional WM.

**Statistical Analyses**

The main trial planned to enroll 126 UHR individuals. Power-calculation estimating sample-size was based on cognitive outcomes for the main trial (26).

Distribution of continuous data (clinical, cognitive and demographic data) was tested for normality, and group differences were tested using general linear modelling (GLM) co-varied for age and gender. Group differences in ordinal data (tobacco smoking, alcohol, drug use, and comorbidity) were tested using the Mann–Whitney U test or Fisher’s exact test. Nominal data were tested using Pearson’s χ² test. Group differences on baseline whole brain and regional FA were tested using univariate GLM and co-varied for age, gender, and relative and absolute head motion in the scanner.

Main analyses of treatment effect of cognitive remediation on primary and secondary outcomes were conducted using linear mixed models with repeated measurements and an unstructured covariance matrix as recommended for RCTs (46), assessing the interaction term between timepoint and group (cognitive remediation versus TAU). Covariates were age, gender, IQ, antipsychotic medication at baseline, and relative and absolute head motion in the scanner. Analyses were conducted according to the intention-to-treat principle, analyzing all participants, who completed MRI-scanning in the groups they were assigned to by randomization. Missing data were handled implicitly by the linear mixed modelling by full information maximum likelihood. Results were corrected for multiple comparisons using Bonferroni correction, and the significance threshold for the between-group analyses was set at P ≤ 0.001 (0.05/(1 whole brain + 5 regions) x 4 WM-measures x 2 treatment-groups)) (47).
Planned exploratory correlation analyses between changes in WM-measures on secondary outcomes and changes in clinical symptoms and cognitive functions, respectively, were conducted with bivariate correlation analyses using Pearson’s r or Spearman’s rho as appropriate. In order to examine correlations specifically related to a potential treatment effect, testing was conducted on the predefined ROIs, which had shown an interaction-effect on time*group. Change-scores were calculated as: [numerical follow-up score – numerical baseline score]. Testing was performed with and without extreme outliers. If the main results remained unaffected, outliers were included in the analyses. Results were corrected for multiple comparisons using Bonferroni correction, and the significance threshold was set at $P \leq 0.001 (0.05/(1 \text{ ROI} \times 6 \text{ cognitive tests} \times 4 \text{ WM-measures} \times 2 \text{ treatment-groups})).$

Exploratory post-hoc analyses to test potential dose-response effect of cognitive remediation at the follow-up were conducted with linear mixed models with repeated measurements and an unstructured covariance matrix, with covarying for number of attended neurocognitive training-hours. All analyses were covaried for age, gender, IQ, antipsychotic medication, and head motion in scanner (relative/absolute).

All analyses were performed using IBM SPSS Statistics for Windows, Version 25.0, Armonk, NY, and the significance level was set to 0.05 (two-sided).

**RESULTS**

The main trial results have been described elsewhere (no effect of cognitive remediation on primary outcome of global neurocognition, or secondary outcomes of social- and occupational functioning, negative- or depressive symptoms) (27). In short, a total of 146 UHR individuals were assigned to either TAU or TAU + cognitive remediation (see Figure 1). Of these, 113 were MRI-scanned, and included in the current study.

**FIGURE 1** | Flowchart of the FOCUS trial. CR, cognitive remediation; MRI, magnetic resonance imaging; TAU, treatment as usual; UHR, ultra-high risk.
Two MRI-scans were excluded due to poor image quality and extreme values. Thus, 111 UHR individuals were included in the analyses (Figure 1). Twenty UHR individuals discontinued the cognitive remediation intervention (18% attrition rate). Further nine UHR individuals did not complete follow-up scans due to practical issues (exceeded timeframe, scanner breakdown), resulting in a total of 82 UHR individuals completing MRI-scans at follow-up. The cognitive remediation group received significantly less TAU with a mean of 20 h within the 26 weeks intervention period compared to the TAU group (mean, 26 h). The cognitive remediation group completed a mean of 12 h of neurocognitive training. Fidelity-rating to the SCIT treatment corresponded to an excellent adherence to the SCIT therapy manual, and inter-rater reliability on the clinical outcome measures showed excellent agreement (ICC ratings from 0.96 to 0.99). No adverse events were reported relating to the intervention.

Cognitive remediation and TAU-groups did not differ significantly on sociodemographic variables (Table 1), clinical symptoms or cognitive functions (Table 2). We found no significant difference on whole brain or regional FA between UHR individuals allocated to cognitive remediation or TAU on both timepoints (Figure 2). At 26-weeks follow-up, remission from UHR-status was similar in both cognitive remediation and TAU-groups (16% and 18% respectively); as well as conversion-rates to psychosis (9% and 12% respectively) (Table 2).

The main mixed modelling analyses revealed no effect of the cognitive remediation on the primary outcome of whole-brain FA, with no significant interaction between timepoint*treatment (P=0.81, Table 3). Secondary outcome on regional WM-measures revealed a nominal significant interaction between timepoint*treatment of AD in left medial lemniscus (P=0.016, Table 3), which did not survive Bonferroni-correction for multiple comparisons.

Exploratory analyses of associations between change in WM-measures in the region with significant-between-group differences on secondary outcomes, and change in cognitive functions revealed a significant negative correlation between WM (AD increase) in left medial lemniscus, and changes in CANTAB IED (total errors adjusted decreased) (P=0.001, CC =0.54) for the cognitive remediation group, but not for TAU (Figure 3). No other correlations survived Bonferroni correction (See Supplementary Tables S4 and S5 for details).

In the exploratory analyses of dose-response effects in the cognitive remediation group, we found no effect when adding number of neurocognitive training hours as a covariate to WM-changes.

### DISCUSSION

To our knowledge, this is the first randomized, clinical trial to evaluate the effect of cognitive remediation on white matter in UHR individuals. Contrary to our main hypothesis, we found no effect of cognitive remediation on the primary outcome of whole-brain FA. On our secondary outcome a nominal significant effect on one predefined white matter region (left medial lemniscus) did not survive adjustment for multiple comparisons. The lack of global and regional WM-changes as a result of cognitive remediation we found are in line with the results from the main trial (27), which did not result in improvements in global measures of cognition, function, or clinical symptoms, but marginal effects on exploratory outcomes (i.e. emotion recognition). Nor did our dose-response analyses on completed hours of neurocognitive training in the cognitive remediation group support an effect on FA in whole brain or ROIs.

The result could, beyond the conclusion of no effect of cognitive remediation on WM, be explained by the low number of neurocognitive training sessions. The average of 12 h of neurocognitive training is considerably lower than the recommended dosage (25–30 h) assumed necessary for driving cognitive improvements (48). A controlled, randomized trial on
patients with schizophrenia reported the neuroplastic effects of cognitive remediation (increased FA in corpus callosum), which was correlated to cognitive executive functions (18). However, this intervention group was exposed to 40 1-h sessions of neurocognitive training with a higher frequency, which is a considerably larger dose than the UHR individuals in the current trial. The question of sufficient dosage to induce consistent WM-changes might be particularly important when evaluating neuroplasticity-based treatment. A multistage model for associations between learning and cerebral changes would explain how rapid improvements in learning and behavior may be linked to transient and elastic WM-changes (49), which tend to return to baseline at subsequent follow-ups (50). Consolidation and automatization of training and learning, which would be accompanied by more robust neuroplastic reorganization, would appear within a longer time-frame (51). Indeed, several studies have demonstrated a sustained effect on WM-measures, suggesting a temporal dynamic in progressive WM-reorganization associated with increased skills and new learning (52–54). However, the underlying biological mechanisms involved in these time-dependent variations are not straightforward, as multiple cellular

### TABLE 2 | Clinical and cognitive data at baseline and follow-up for UHR individuals.

| Variable                  | UHR-CR Mean (S.D./No. (Percent)) | UHR-TAU Mean (S.D./No. (Percent)) |
|---------------------------|----------------------------------|-----------------------------------|
|                          | Baseline (N = 54) | Follow-Up (N = 43)    | Baseline (N = 57) | Follow-up (N = 48) |
| Medication                |                    |                       |                    |                       |
| Antipsychotic-naive       | 34 (64.2%)     | 16 (37.2%)           | 29 (50.9%)        | 12 (25.0%)           |
| Current\* antipsychotics  | 14 (26.4%)     | 24 (55.8%)           | 21 (36.8%)        | 33 (68.8%)           |
| Current\* antidepressants | 13 (24.5%)     | 12 (28.6%)           | 16 (28.1%)        | 11 (22.9%)           |
| Current\* mood stabilizers| 2 (3.8%)       | 5 (11.6%)            | 4 (7.0%)          | 4 (8.3%)             |
| Current\* benzodiazepines | 1 (1.9%)       | 6 (14.0%)            | 7 (12.3%)         | 4 (8.3%)             |
| Diagnose of current\* abuse | 0 (0.0%)   | 0 (0.0%)             | 1 (1.9%)          | 0 (0.0%)             |
| Diagnose of current\* dependency | 0 (0.0%) | 0 (0.0%)            | 1 (1.9%)          | 0 (0.0%)             |
| Clinical outcomes         |                    |                       |                    |                       |
| CAARMS composite          | 51.9 (11.9)    | 38.1 (16.7)          | 49.7 (16.5)       | 33.2 (15.3)          |
| SANS                      | 1.6 (0.7)      | 1.4 (0.9)            | 1.5 (0.8)         | 1.1 (1.6)            |
| BPRS                      | 42.9 (7.6)     | 37.8 (12.6)          | 41.0 (10.7)       | 38.0 (10.8)          |
| MADRS                     | 16.3 (7.4)     | 12.0 (7.5)           | 15.0 (7.0)        | 13.1 (7.2)           |
| Remission from UHR-status | 7 (16.3%)      | 5 (9.3%)             | 2 (4.1%)          | 8 (17.8%)            |
| Conversion to psychosis   | 5 (9.3%)       | 7 (13.3%)            | 7 (13.3%)         | 7 (13.3%)            |
| Drop-out                  | 11 (20.4%)     | 9 (15.8%)            |                    |                       |
| Cognitive functions       |                    |                       |                    |                       |
| IQ                        | 102.8 (12.7)   | 104.5 (12.4)         |                    |                       |
| BACS                      |                    |                       |                    |                       |
| List-Learning             | 51.7 (8.9)     | 53.2 (8.8)           | 50.4 (7.8)        | 53.1 (9.4)           |
| Digit sequencing          | 21.0 (3.9)     | 21.1 (4.1)           | 20.3 (4.4)        | 21.1 (4.2)           |
| Fluency                   | 58.2 (15.2)    | 59.2 (18.6)          | 57.4 (11.7)       | 59.9 (12.3)          |
| Symbol coding             | 57.3 (11.3)    | 62.2 (11.7)          | 59.0 (11.5)       | 62.3 (12.7)          |
| CANTAB                    |                    |                       |                    |                       |
| SOC                       | 9.8 (1.7)      | 10.8 (2.1)           | 10.1 (1.8)        | 10.3 (1.6)           |
| IED                       | 18.6 (16.0)    | 20.1 (34.8)          | 19.2 (16.0)       | 11.4 (10.0)          |

The clinical characteristics and cognitive functions for UHR individuals at baseline and follow-up are shown.

*Atypical antipsychotics in low dose: aripiprazole, amisulpride, olanzapine, paliperidone, quetiapine, risperidone.

†Current, the last month.

APS, attenuated psychotic symptoms; BACS, brief assessment of cognition in schizophrenia battery; BLIPS, brief limited intermittent psychotic symptoms; CAARMS, comprehensive assessment of at-risk mental state; CANTAB< Cambridge neuropsychological test automated battery; CR, cognitive remediation; IED, intra-extradimensional set shifting test, total errors, adjusted; IQ, intelligence quotient; N, count; SD, standard deviation; SOC, stockings of Cambridge, Problems solved in minimum moves; UHR, ultra-high risk.

FIGURE 2 | Boxplots illustrating the development of mean whole brain fractional anisotropy at baseline and 26 weeks follow-up for UHR individuals allocated to cognitive remediation versus treatment as usual are displayed. Error bars indicate the 95% confidence intervals. Note that the y-axis has been altered to enhance visual display. CR, cognitive remediation; TAU, treatment as usual; UHR, ultra-high risk.
processes may be associated with WM-measures such as FA. Various candidate mechanisms of activity-dependent WM-reorganization has been suggested, such as long-term potentiation (55), growth of astrocytes and oligodendrocyte precursor cells (56), myelination (57), and reorganization of axons and fiber-bundles (58). Our results stress the perspective of investigating duration and intensity as prerequisites of neuroplasticity-based interventions embedded in a broader clinical setting. Moreover, future research specifying the potential active treatment ingredients beyond duration and intensity, such as choice of remediation strategies (i.e. integrative versus simple programs, or compensatory versus restorative remediation approaches) may support the clinical tailoring of treatment strategies. The clinical perspectives includes enhancement of motivation and attendance (21, 59), as well as the question of whether this comprehensive treatment may only be viable for the subgroup of UHR individuals who can be engaged sufficiently to practice the skills in the recommended dosage.

Exploratory analyses revealed mental flexibility to be correlated with a change of left medial lemniscus AD in the cognitive remediation group, but not in the TAU-group. Although our observation is marginal and should be interpreted very cautiously, medial lemniscus is part of the cerebello-thalamocortical connections [CTC (60)], which previously has been associated with symptom course in UHR individuals (61, 62). Interestingly, our observation would be consistent with a recent study in patients with schizophrenia, reporting that WM organization in medial lemniscus predicted training induced improvements in executive functioning.

### TABLE 3 | White matter characteristics at baseline and follow-up in individuals at ultra-high risk for psychosis.

|                      | FA          | AD Mean (SD) x10^{-2} | RD Mean (SD) x10^{-2} | MD Mean (SD) x10^{-2} |
|----------------------|-------------|-----------------------|-----------------------|-----------------------|
|                      | CR          | TAU                   | CR                    | TAU                   | CR          | TAU       |
| **Primary outcomes** |             |                       |                       |                       |             |           |
| Whole Brain          |             |                       |                       |                       |             |           |
| BL                   | 0.578 (0.0139) | 0.5805 (0.0167)      | 0.1360 (0.0224)       | 0.1361 (0.0039)       | 0.0462 (0.0019) | 0.0459 (0.0025) |
| FU                   | 0.579 (0.0160) | 0.5814 (0.0160)      | 0.1354 (0.0021)       | 0.1357 (0.0028)       | 0.0459 (0.0022) | 0.0456 (0.0023) |
| Interaction Time*Group | P=0.81       | F=0.06               | P=0.85               | F=0.04               | P=0.28      | F=1.18    |
| **Secondary outcomes** |             |                       |                       |                       |             |           |
| Fornix               |             |                       |                       |                       |             |           |
| BL                   | 0.5656 (0.0743) | 0.5546 (0.0625)    | 0.1985 (0.0196)       | 0.2003 (0.0148)       | 0.0797 (0.0252) | 0.0829 (0.0018) |
| FU                   | 0.5594 (0.0837) | 0.5543 (0.0649)    | 0.1999 (0.0211)       | 0.2022 (0.0166)       | 0.0814 (0.0209) | 0.0829 (0.0019) |
| Interaction Time*Group | P=0.23       | F=0.28               | P=0.64               | F=0.44               | P=0.59      | F=0.53    |
| Right medial lemniscus |             |                       |                       |                       |             |           |
| BL                   | 0.6313 (0.0384) | 0.6336 (0.0249)    | 0.1399 (0.0068)       | 0.1395 (0.0062)       | 0.0440 (0.0048) | 0.0435 (0.0031) |
| FU                   | 0.6302 (0.0346) | 0.6411 (0.0265)    | 0.1386 (0.0059)       | 0.1389 (0.0065)       | 0.0437 (0.0039) | 0.0424 (0.0031) |
| Interaction Time*Group | P=0.76       | F=0.28               | P=0.33               | F=1.11               | P=0.69      | F=0.62    |
| Left medial lemniscus |             |                       |                       |                       |             |           |
| BL                   | 0.6179 (0.0386) | 0.6253 (0.0334)    | 0.1396 (0.0064)       | 0.1387 (0.0067)       | 0.0456 (0.0045) | 0.0443 (0.0040) |
| FU                   | 0.6266 (0.0318) | 0.6341 (0.0327)    | 0.1399 (0.0053)       | 0.1367 (0.0072)       | 0.0447 (0.0041) | 0.0428 (0.0036) |
| Interaction Time*Group | P=0.37       | F=0.99               | P=0.02               | F=3.74               | P=0.14      | F=2.03    |
| Left superior cerebellar peduncle |             |                       |                       |                       |             |           |
| BL                   | 0.6422 (0.0319) | 0.6492 (0.0373)    | 0.1618 (0.0071)       | 0.1613 (0.0074)       | 0.0498 (0.0046) | 0.0488 (0.0059) |
| FU                   | 0.6403 (0.0311) | 0.6472 (0.0342)    | 0.1602 (0.0058)       | 0.1593 (0.0075)       | 0.0495 (0.0048) | 0.0486 (0.0058) |
| Interaction Time*Group | P=0.40       | F=0.92               | P=0.18               | F=1.74               | P=0.17      | F=2.20    |
| Left uncinate fasciculus |             |                       |                       |                       |             |           |
| BL                   | 0.5034 (0.0412) | 0.5027 (0.0303)    | 0.1277 (0.0066)       | 0.1277 (0.0066)       | 0.0541 (0.0037) | 0.0543 (0.0031) |
| FU                   | 0.5017 (0.0431) | 0.5009 (0.0369)    | 0.1272 (0.0054)       | 0.1277 (0.0067)       | 0.0541 (0.0022) | 0.0544 (0.0029) |
| Interaction Time*Group | P=0.77       | F=0.27               | P=0.94               | F=0.06               | P=0.92      | F=0.74    |

Mean and standard deviation (SD) of DWI data at baseline and 26-weeks follow-up for UHR individuals, comparing the cognitive remediation intervention-group (CR) to treatment as usual (TAU), and the interaction effect between timepoint*group are shown. The model comparing CR versus TAU are covared for age, gender, IQ, antipsychotic medication, and movement in scanner. Group*Time interaction significant before correction for multiple comparisons are marked in bold.

AD, axial diffusivity; CR, cognitive remediation; DWI, diffusion weighted imaging; FA, fractional anisotropy; MD, mean diffusivity; RD, radial diffusivity; SD, standard deviation; BL, Baseline; FU, Follow-up; TAU, treatment as usual; UHR, ultra-high risk.
CTC-connections may be an important determinant for training induced neurocognitive plasticity (63). We speculate whether this correlation may indicate a subtle protective effect of cognitive remediation on loss of WM-integrity. The protective approach of intact cognitive functions have been discussed as an alternative to focusing on restoration of impaired functions by cognitive remediation (64). Our observations lend support to further investigations of strategies for cognitive remediation, i.e. specifically aimed at preserving targeted cognitive functions while potentially preventing WM-deterioration in early psychosis, versus the integrative program combining various remediation strategies as applied in the current trial.

A strength of this study is that it was undertaken as part of a high quality randomized, assessor-blinded clinical trial. Limiting the study is the moderate attrition-rate, along with the low number of mean neurocognitive training hours, which impede reaching firm conclusions about the effect of cognitive remediation delivered per the trial-protocol. Furthermore, sample-size was calculated on different outcomes from the main trial, which is a weakness in our study-design. However, this study presents a large sample-size compared to previous studies examining neuroplastic effects of cognitive interventions.

CONCLUSION

Cognitive remediation comprising 12 h of neurocognitive training on average did not improve global or regional WM organization in UHR individuals. Further investigations of the duration and intensity of neuroplasticity-based cognitive training as necessary prerequisites of WM-changes are warranted.

DATA AVAILABILITY STATEMENT

The data sets presented in this article are not readily available because: Not allowed by the Danish Data Protection Agency.

Requests to access the datasets should be directed to TK, tina.dam.kristensen@regionh.dk.

ETHICS STATEMENT

The studies involving human participants were reviewed and approved by The study protocol was approved by the Committee on Health Research Ethics of the Capital Region Denmark (study: H-6-2013-015) and the Danish Data Protection Agency (RHP-2014-009-02670). The patients/participants provided their written informed consent to participate in this study.

AUTHOR CONTRIBUTIONS

MN, LG, and BF conceived and designed the trial protocol. MN, BG, LG, CH, TK, and CW raised the funding. JJ and BF supervised the data collection. BE, RM, and CP supervised the study. JR processed MRI-data. TK conducted the analyses supervised by CH. TK wrote the initial draft. All authors contributed to the article and approved the submitted version.

FUNDING

The study was funded by The Danish Council for Independent Research (DFF-4004-00314); TrygFoundation (108119); the Mental Health Services in the Capital Region of Denmark; the research fund of the Capital Region of Denmark; the Lundbeck Foundation Center for Clinical Intervention and Neuropsychiatric Schizophrenia Research, CINS (R155-2013-16337). CP was supported by a NHMRC Senior Principal Research Fellowship (1105825) and by a grant from Lundbeck Foundation (R246-2016-3237).
ACKNOWLEDGMENTS

We thank the participants and clinical staff for their participation. In particular, we are grateful that Lise Mariegaard as the senior psychologist delivered the FOCUS-intervention with excellence.

REFERENCES

1. Fusar-Poli P, Bonoldi I, Yung AR, Borgwardt S, Kempton MJ, Valsaggia L, et al. Predicting Psychosis. Arch Gen Psychiatry (2012) 69(3):220–9. doi: 10.1001/archgenpsychiatry.2011.1472
2. Bora E, Lin A, Wood SF, Yung AR, McGorry PD, Panetis C. Cognitive deficits in youth with familial and clinical high risk to psychosis: A systematic review and meta-analysis. Acta Psychiatr Scand (2014) 130(1):1–15. doi: 10.1111/acps.12261
3. Fusar-Poli P, Deste G, Smieskova R, Barlati S, Yung AR, Howes O, et al. Cognitive Functioning in Prodromal Psychosis. Arch Gen Psychiatry (2012) 69(6):562–71. doi: 10.1001/archgenpsychiatry.2011.1592
4. De Herdt A, Wampers M, Vancampfort D, De Hert M, Vanhees L, Demunter H, et al. Neurocognition in clinical high risk young adults who did or did not convert to a first schizophrenic psychosis: A meta-analysis. Schizophr Res (2013) 149(1–3):48–55. doi: 10.1016/j.schres.2013.06.017
5. Mam-lam-Fook C, Danset-Alexandre C, Pedron L, Amado I, Gaillard R, Krebs M-O. Neuropsychology of subjects with ultra-high risk (UHR) of psychosis: A critical analysis of the literature. Encephale (2017) 43(3):241–53. doi: 10.1016/j.enep.2017.02.001
6. de Paula ALD, Hallak JEC, Machado-de-Sousa JP. Cognition at risk-at-risk mental states for psychosis. Neurosci Biobehav Rev (2015) 57:199–208. doi: 10.1016/j.neubiorev.2015.09.006
7. Vijayakumar N, Bartholomeusz C, Whitford T, Hermens DF, Nelson B, Rice S, et al. White matter integrity in individuals at ultra-high risk for psychosis: a systematic review and discussion of therole of polyunsaturated fatty acids. BMC Psychiatry (2016) 16(1):1. doi: 10.1186/s12888-016-0932-4
8. Le Bihan D, Iima M. Diffusion magnetic resonance imaging: What water tells us about biological tissues. PLoS Biol (2015) 13(7):1–13. doi: 10.1371/journal.pbio.1002203
9. Alexander AL, Lee JE, Lazar M, Field AS. Diffusion tensor imaging of the brain. Neurotherapeutics (2007) 4(3):316–29. doi: 10.1016/j.neurt.2007.05.011
10. Kristensen TD, Mandl RCW, Raghava JM, Jessen K, Jepsen JR, Fagerlund B, et al. Widespread higher fractional anisotropy associates to better cognitive function in individuals at ultra-high risk for psychosis. Hum Brain Mapp (2019) 40(18):5185–201. doi: 10.1002/hbm.24765
11. Krakauer K, Ebdrup BH, Glenthoj BY, Raghava JM, Nordentoft M, et al. White matter microstructure in individuals at ultra-high risk for psychosis: Associations to level of functioning and clinical symptoms. Psychol Med (2017) 47(15):2689–707. doi: 10.1017/S0033291717001210
12. Peters BD, Dingemans PM, Dekker N, Blaas J, Akkerman E, van Amelsvoort T, et al. White matter connectivity and psychosis in ultra-high-risk subjects: A meta-analysis of non-psychiatric and psychiatric studies. Acta Psychiatr Scand (2013) 138(10):954–62. doi: 10.1111/j.1600-0447.2013.12393
13. Penades R, Vallespi LG, Franck N, Dekerle M. Neuroimaging Studies of White Matter Organization: A systematic review of non-psychiatric and psychiatric studies. Neurosci Biobehav Rev (2018) 88(August 2017):84–97. doi: 10.1016/j.neubiorev.2018.03.013
14. Penades R, Pujol N, Catalan R, Massana G, Rametti G, Garcia-Rizo C, et al. Brain effects of cognitive remediation therapy in schizophrenia: A structural and functional neuroimaging study. Biol Psychiatry (2013) 73(10):1015–23. doi: 10.1016/j.biopsych.2013.01.017
15. Keshavan MS, Högarty GE. Brain maturational processes and delayed onset in schizophrenia. Dev Psychopathol (1999) 11(3):525–43. doi: 10.1017/S0954579499002199
16. Wu Y, Huddy V, Cellard C, McGurk SR, Czobor P. A meta-analysis of cognitive remediation for schizophrenia: Methodology and effect sizes. Am J Psychiatry (2011) 168(5):472–85. doi: 10.1176/appi.ajp.2010.10060855
17. Mogami T. Cognitive remediation for schizophrenia with focus on NEAR. Front Psychiatry (2018) 8:304[IAN]. doi: 10.3389/fpsyg.2017.00304
18. McGurk SR, Twwamey EW, Sitzer DJ, GM, Mueser KT. A Meta-Analysis of Cognitive Remediation in Schizophrenia. Am J Psychiatry (2007) 164(12):1791–802. doi: 10.1176/appi.ajp.2007.07060906.A
19. Blehaut JB, Hjorthøj C, Kristensen TD, Davidson CA, Nordentoft M. The effect of cognitive remediation in individuals at ultra-high risk for psychosis: a systematic review. NPJ Schizophrenia (2017) 3(1):20. doi: 10.1038/s41537-017-0021-9
20. Loewy R, Fisher M, Schlosser DA, Biagianti B, Stuart B, Mathalon DH, et al. Intensive Auditory Cognitive Training Improves Verbal Memory in Adolescents and Young Adults at Clinical High Risk for Psychosis. Schizophr Bull (2016) 42 Suppl 1:S18–26. doi: 10.1093/schbul/bsw009.
21. Choi J, Corcoran CM, Fiszdon JM, Stevens M, Javitt DC, Deasy M, et al. Pupilometer-Based Neurofeedback Cognitive Training to Improve Processing Speed and Social Functioning in Individuals at Clinical High Risk for Psychosis. Psychatri Rehabil J (2016) 40(1):33–42. doi: 10.1037/prj0000027
22. Glenthøj LB, Fagerlund B, Jepsen JR, Fagerlund B, Jepsen JM, Nordentoft M, et al. Cognitive remediation plus standard treatment versus standard treatment alone for individuals at ultra-high risk of developing psychosis: results of the FOCUS randomised clinical trial. Schizophrenia Res (2020) 19:2(September):35–4. doi: 10.1016/wps.26760
23. Yung AR, Yuen HP, McGorry PD, Phillips LJ, Kelly D, Dell’Olio M, et al. Mapping the onset of psychosis: The Comprehensive Assessment of At-Risk Mental States. Aust N Z J Psychiatry (2005) 39(11–12):964–71. doi: 10.1111/ j.1440-1614.2005.0174X.x
24. First MB, Gibbon M. The Structured Clinical Interview for DSM-IV Axis I Disorders (SCID-I) and the Structured Clinical Interview for DSM-IV Axis II Disorders (SCID-II). In: Comprehensive Handbook of Psychological Assessment, Vol. 2: Personality Assessment. Hoboken, NJ, US: John Wiley & Sons Inc (2004). p. 134–43.
25. Spitzer RL, Williams JBW, Gibbon M, First MB. User’s Guide for the Structured Clinical Interview for DSM-III-R: SCID. Arlington, VA, US: American Psychiatric Association (1990).
26. Ventura J, Nuechterlein KH, Subotnik KL, Gutfried D, Gilbert E. A symptom dimensions in recent-onset schizophrenia and mania: A principal components analysis of the 24-item Brief Psychiatric Rating Scale. Psychiatry Res (2000) 97(2–3):129–35. doi: 10.1016/S0165-1781(00)00228-6
27. Andreasen NC. Scale for the Assessment of Negative Symptoms. Iowa City, Iowa: Dept. of Psychiatry, College of Medicine, the University of Iowa (1984).
33. Montgomery S a., Asberg M. A new depression scale designed to be sensitive to change. *Br J Psychiatry* (1979) 134(4):382–9. doi: 10.1192/bjp.134.4.382

34. Wechsler D. *Wais-3 Administration and Scoring Manual*. San Antonio TX: The Psychological Corporation (1997). Available at: http://www.amazon.com/Wais-3-Administration-Scoring-Wechsler/dp/0158981030.

35. Keefe RSE, Harvey PD, Goldberg TE, Gold JM, Walker TM, Kennel C, et al. Norms and standardization of the Brief Assessment of Cognition in Schizophrenia (BACS). *Schizophr Res* (2008) 102(1-3):108–15. doi: 10.1016/j.schres.2008.03.024

36. Sahakian BJ, Owen a. M. Computerized assessment in neuropsychiatry using CANTAB: discussion paper. *R Soc Med* (1992) 85(July):399–402. doi: 10.1177/014107689208500711

37. Anderson JLR, Skare S, Ashburner J. How to correct susceptibility distortions in spin-echo echo-planar images: Application to diffusion tensor imaging. *Neuroimage* (2003) 20(2):870–88. doi: 10.1016/S1053-8119(03)00336-7

38. Jenkinson M, Beckmann CF, Behrens TEJ, Woolrich MW, Smith SM. Fsl. *Neuroimage* (2012) 62(2):782–90. doi: 10.1016/j.neuroimage.2011.09.015

39. Smith SM, Jenkinson M, Johansen-Berg H, Rueckert D, Nichols TE, Mackay CE, et al. Tract-based spatial statistics: Voxelwise analysis of multi-subject diffusion data. *Neuroimage* (2006) 31(4):1487–505. doi: 10.1016/j.neuroimage.2006.02.024

40. Mori S, Van Zijl P. Human white matter atlas. *Am J Psychiatry* (2007) 164(July):73390. doi: 10.1176/appi.ajp.164.7.1005

41. Roalf DR, Quarzme M, Elliott MA, Satterwhite TD, Vandenak SN, Ruparel K, et al. The Impact of Quality Assurance Assessment on Diffusion Tensor Imaging Outcomes in a Large-Scale Population-Based Cohort. *Neuroimage* (2016) 125:903–19. doi: 10.1016/j.neuroimage.2015.10.068

42. Eack SM. Cognitive Enhancement Therapy for Early-Course Schizophrenia: Effects of a. *Psychiatr Serv* (2009) 60:1468–76. doi: 10.1176/ps.2009.60.11.1468

43. Medalia A, Freidlich B. The Neuropsychological Educational Approach to Cognitive Remediation (NEAR) Model: Practice Principles and Outcome Studies. *Am J Psychiatry Rehabil* (2008) 11(2):123–43. doi: 10.1080/1548769081963660

44. Roberts DL, David P, Combs DR. *Cognitive Remediation (NEAR) Model: Practice Principles and Outcome Study*. All grants are the property of the Mental Health Services in the Capital Region of Denmark, and other foundations. Her group has also received a grant from the Lundbeck Foundation based on international review and partially financed by the Lundbeck Foundation Centre of Excellence for Clinical Intervention and Neuropsychiatric Research. The Psychological Corporation (1997). Available at: http://www.amazon.com/ Corsini’s Encyclopedia of Psychology/Print/1-59385-233-4

45. Penn DL, Roberts DL, Combs D, Sterne A. Best practices: The development of the Social Cognition and Interaction Training program for schizophrenia spectrum disorders. *Psychiatr Serv* (2007) 58(4):449–51. doi: 10.1176/appi.ps.58.4.449

46. Lu K, Mehrotra DV. Speci-...
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Kristensen, TD; Ebdrup, BH; Hjorthoj, C; Mandl, RCW; Raghava, JM; Jepsen, JRM; Fagerlund, B; Glenthoj, LB; Wenneberg, C; Krakauer, K; Pantelis, C; Glenthoj, BY; Nordentoft, M

Title:
No Effects of Cognitive Remediation on Cerebral White Matter in Individuals at Ultra-High Risk for Psychosis-A Randomized Clinical Trial

Date:
2020-08-28

Citation:
Kristensen, T. D., Ebdrup, B. H., Hjorthoj, C., Mandl, R. C. W., Raghava, J. M., Jepsen, J. R. M., Fagerlund, B., Glenthoj, L. B., Wenneberg, C., Krakauer, K., Pantelis, C., Glenthoj, B. Y. & Nordentoft, M. (2020). No Effects of Cognitive Remediation on Cerebral White Matter in Individuals at Ultra-High Risk for Psychosis-A Randomized Clinical Trial. FRONTIERS IN PSYCHIATRY, 11, https://doi.org/10.3389/fpsyt.2020.00873.

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