Longitudinal clinical and functional outcome in distinct cognitive subgroups of first-episode psychosis: a cluster analysis


Abstract

**Background.** Cognitive deficits may be characteristic for only a subgroup of first-episode psychosis (FEP) and the link with clinical and functional outcomes is less profound than previously thought. This study aimed to identify cognitive subgroups in a large sample of FEP using a clustering approach with healthy controls as a reference group, subsequently linking cognitive subgroups to clinical and functional outcomes.

**Methods.** 204 FEP patients were included. Hierarchical cluster analysis was performed using baseline brief assessment of cognition in schizophrenia (BACS). Cognitive subgroups were compared to 40 controls and linked to longitudinal clinical and functional outcomes (PANSS, GAF, self-reported WHODAS 2.0) up to 12-month follow-up.

**Results.** Three distinct cognitive clusters emerged: relative to controls, we found one cluster with preserved cognition (n = 76), one moderately impaired cluster (n = 74) and one severely impaired cluster (n = 54). Patients with severely impaired cognition had more severe clinical symptoms at baseline, 6- and 12-month follow-up as compared to patients with preserved cognition. General functioning (GAF) in the severely impaired cluster was significantly lower than in those with preserved cognition at baseline and showed trend-level effects at 6- and 12-month follow-up. No significant differences in self-reported functional outcome (WHODAS 2.0) were present.

**Conclusions.** Current results demonstrate the existence of three distinct cognitive subgroups, corresponding with clinical outcome at baseline, 6- and 12-month follow-up. Importantly, the cognitively preserved subgroup was larger than the severely impaired group. Early identification of discrete cognitive profiles can offer valuable information about the clinical outcome but may not be relevant in predicting self-reported functional outcomes.

Introduction

Despite relatively successful treatment of clinical symptoms after first-episode psychosis (FEP) (Kahn et al. 2018), many patients continue to experience ongoing functional impairment in day-to-day life (Henry et al. 2010; Lally et al. 2017). Large variability exists in the outcome of FEP with recovery rates ranging from 13.5% to 38% (Jääskeläinen et al. 2013; Lally et al. 2017). A significant minority of patients shows the excellent recovery, but a large proportion of patients continues to exhibit moderate or severe functional impairment (Jääskeläinen et al. 2013; Lally et al. 2017). Recovery rates appear to be stable 2 years after illness onset as demonstrated in a large meta-analysis (Lally et al. 2017), underscoring the importance of identifying factors that can predict outcome overtime in the early stages of disease onset. However, most longitudinal studies have examined predictors at the diagnostic group level and do not take the high heterogeneity between individual patients with the same diagnosis into account.
Also, the demonstrated that cognitive performance in a
ences in severity of cognitive dysfunction. Indeed, it has been
both functional and clinical outcomes may be related to differ-
outcome, leaving a significant proportion of the variance unex-
impairment may be less pronounced as previously thought.
predictive value of cognitive deficits in terms of functional
schizophrenia (Heinrichs, 2005). However, recent literature
have considered cognitive dysfunction to be the core feature of
Van Rheenen, Gurvich, Sumner, & Rossell, 2019; Moritz et al.
tial subset of patients that remains cognitively intact (Carruthers,
groups may exist within the FEP population, including a substan-
tial subset of patients that remains cognitively intact (Carruthers,
also, the predictive value of cognitive deficits in terms of functional
may be less pronounced as previously thought. Notably, a recent meta-analysis showed only small to medium
effect sizes for the association between cognition and functional
outcome, leaving a significant proportion of the variance unex-
plained (Halverson et al. 2019). It is plausible that variance in
both functional and clinical outcomes may be related to differ-
es in severity of cognitive dysfunction. Indeed, it has been
demonstrated that cognitive performance in a “neuropsychologi-
cally normal” range does not correlate well with aspects of every-
day functioning whereas more severe levels of cognitive
impairment do seem to be associated with functional outcomes
(Strassnig et al. 2018). This underscores the value of grouping
FEP patients into subtypes along the cognitive continuum, dem-
onstrating possible subgroups with distinct illness profiles.
An essential and relatively novel solution for determining
homogeneous subgroups is a data-driven clustering approach.
Defining subgroups based on baseline cognitive profile may pro-
vide crucial information regarding functional outcome and prog-
nosis. Such information is urgently needed, as the high
heterogeneity and lack of good predictors hamper clinicians in
providing optimal care for individual patients. Early identification
of risk factors associated with poor outcomes is highly valuable as
this would aid individually tailored interventions that may posi-
tively impact the long-term outcome.
The current study includes a large sample of FEP patients who
were 3–6 months in remission of their psychotic symptoms at
baseline, to identify homogeneous subgroups of cognition based
on a data-driven clustering approach. Factors that may influence
cognitive function, such as the distraction by unusual ideas and/or
hallucinations, long-term antipsychotic medication use or the
duration of illness, are limited in the current sample as all patients
were in a similar early stage of their illness. Emergent cognitive
subgroups were subsequently compared to healthy controls to assess
the level of cognitive (under)performance. Cognitive subgroups
were then evaluated according to clinical [Positive and Negative
Syndrome Scale (PANSS), Global Assessment of Functioning
(GAF)] and functional [WHO Disability Assessment Scale 2.0
(WHODAS2.0)] outcome at baseline and longitudinally at 6-
and 12-month follow-up. The clinician-rated GAF has been
widely used in clinical and research settings and has been
adopted as meaningful, however, the DSM-5 recommends a
new tool for the assessment of global functioning and impair-
ment, the WHODAS 2.0, a patient self-report assessment tool
that evaluates the patient’s ability to perform activities in six
domains of functioning (Gold, 2014). Based on a recent system-
atic review regarding cognitive subgrouping studies in schizo-
phrenia spectrum disorders, we expected to find three distinct
cognitive subtypes; a relatively intact cognitive subgroup, an
intermediate cognitive subgroup and a globally impaired sub-
group (Carruthers et al. 2019). We further hypothesized that
emergent cognitive subtypes are characterized by differences
in both clinical and functional outcomes at baseline and
follow-up.

Method
Participants
Data were used from the ongoing Handling Antipsychotic Medication: Long-term Evaluation of Targeted Treatment
(HAMLETT) study (Begemann et al. 2020). Patients were
recruited from outpatient settings in 24 healthcare centers
throughout the Netherlands. Written informed consent was
obtained from all participants and study procedures were per-
formed according to the Declaration of Helsinki (64th WMA
general assembly; October 2013). Ethics approval was obtained from
the research and ethics committee of the University Medical
Center Groningen, the Netherlands (protocol number: NL
62202.042.17, trial registration EudraCT number: 2017-002406-12).
Recruitment and study procedures are described in detail by
Begemann et al. (2020).
In short, the current study included data from 204 patients
aged between 16 and 60 years old with the first episode of
schizophrenia, schizoaffective disorder, schizophreniform dis-
order, brief psychotic disorder, delusional disorder, substance/medication-induced psychotic disorder, or those classified as
Unspecified Schizophrenia Spectrum and Other Psychotic Disorders (DSM-5, or as described in the International
Classification of Diseases-10). Diagnosis and duration of illness
were established by their treating psychiatrist and confirmed by
the Comprehensive Assessment of Symptoms and History
(CASH) (Andreasen, Flaum, & Arndt, 1992). At baseline, all
patients were 3–6 months in remission of their first psychotic epi-
sode and used antipsychotic medication. Symptomatic remission
is defined as “sustained improvement of psychotic symptoms to
the level that any remaining psychotic symptoms (such as hallu-
cinatory experiences, unusual thought content, conceptual disor-
ganization) are mild, which means (consistent with international
remission criteria) that they do not interfere with behavior and
daily functioning.”
Self-reports of current antipsychotic medication use (mg/day)
were converted into a chlorpromazine equivalent (CPZE, mg/day)
for each patient (Gardner, Murphy, O’Donnell, Centorrino, &
Baldessarini, 2010). The highest educational level achieved
(CASH) (Andreasen et al. 1992), was converted into the number
of years of education (YOE; see Online Supplementary Table S1).
Moreover, 40 healthy controls were included as a reference
group for cognitive functioning. Healthy controls did not have
any history of psychiatric illness and were aged between 19 and
45 years (Trial registration: ABR NL50657.041.14).

Procedures
Cognitive testing
Cognitive performance was assessed at baseline using the Dutch
version of the brief assessment of cognition in schizophrenia
https://doi.org/10.1017/S0033291721004153 Published online by Cambridge University Press
Performances of all participants on the subtests of the BACS were standardized by creating z-scores adjusted for gender and age using the norms of Keefe et al. (2004). A composite z-score was calculated by averaging all of the six standardized primary measures from the BACS. Participants missing more than 2 cognitive sub-scores were excluded from analysis (n = 2). For participants with ≤2 missing sub-scores, scores were replaced by the corresponding population mean for that specific domain (n = 8).

Clinical outcome
Clinical symptomatology was assessed by trained central study personnel using the Positive and Negative Symptom Scale (PANSS) at baseline, 6 months and 12-month follow-up (Kay, Fiszbein, & Opler, 1987).

In addition, clinical global functioning was evaluated by trained central study personnel at baseline, 6 months and 12-month follow-up using the GAF (Jones, Thornicroft, Coffey, & Dunn, 1995).

To ensure data quality, assessors are comprehensively trained and the central team of assessors have biannual meetings during which inter-rater reliability is assessed and protocol adherence is checked.

Self-reported functional outcome
Self-reported global functioning and disability were evaluated at baseline, 6 months and 12-month follow-up using the WHO Disability Assessment Schedule 2.0 (WHODAS 2.0). This questionnaire consists of 36 items covering six domains of functioning in everyday life: cognition (understanding and communicating), mobility (moving and getting around), self-care (hygiene, eating, and staying alone), getting along (interacting with other people), life activities (domestic responsibilities, leisure, work and school) and participation (joining in community activities). Participants respond to each item on a 5-point scale from 0 (No Difficulty) to 4 (Extreme Difficulty/Cannot Do). Overall scores range from 0 to 100 with higher scores indicating a greater level of self-reported disability (Üstün, 2010).

Statistical analyses
Statistical analyses were performed using Statistical Package for Social Sciences (SPSS) version 25 (IBM Corp. Released 2017. IBM SPSS Statistics for Windows, Version 25.0. Armonk, NY: IBM Corp.). Healthy controls and FEP patients were compared on demographic variables such as gender, age, years of education and cognitive performance using Pearson’s χ² (categorical variables) and One-way Analysis of Variance (ANOVA, continuous variables). Subsequently, cognitive patient clusters were compared on both clinical (PANSS, GAF) and functional (WHODAS 2.0) outcomes using ANOVA for baseline comparisons and ANCOVA for comparisons at 6- and 12-month follow-up. Post hoc comparisons were conducted for all significant ANOVA and ANCOVA effects, using Bonferroni correction for multiple comparisons.

Results
Demographics
A total of 204 patients and 40 healthy controls were included at baseline. Sociodemographic and clinical characteristics are presented in Online Supplementary Table S2. The group of patients consisted of 148 males (72.5%), the healthy controls included 32 males (80.0%). Patients were significantly older (M = 27.93, s.d. = 8.90) than healthy individuals (M = 24.48, s.d. = 4.98), (p = 0.018). Patients attained fewer years of education compared to healthy controls (p = 0.006) and patients scored significantly worse on both the BACS composite and all subtests (all p < 0.001, executive functioning p = 0.047). At 6- and 12-month follow-up, the sample consisted of 145 and 132 patients respectively.

Cluster solution
Hierarchical clustering (Ward’s method) and K-means optimization using BACS composite scores for the total sample of patients resulted in three distinct cognitive clusters (Table 1). Subgroups were subsequently compared to a group of healthy controls to assess the level of cognitive (under)performance.

One cluster could be described as a relatively preserved group (n = 76). The BACS composite score was not significantly different compared to healthy controls, yet these patients scored significantly lower on attention and processing speed compared to healthy controls (p = 0.008). An intermediate or moderately impaired cognitive cluster (n = 74) displaying reduced functioning on all cognitive domains compared to healthy controls (all p < 0.001), except for executive function (p = 0.730) was observed. Lastly, the severely impaired cognitive cluster (n = 54) showed significant impairments across all domains assessed relative to the controls, with working memory and motor speed showing the
most severe deficits (all $p < 0.001$). Results are demonstrated in Figs 1 and 2.

The relatively preserved cluster was significantly older than the healthy controls ($p = 0.011$), but no age differences were demonstrated between the three cognitive patient clusters. The moderately impaired cluster and severely impaired cluster had received significantly fewer years of education compared to both the healthy controls ($p = 0.007$ and $p < 0.001$, respectively) and the relatively preserved cluster ($p = 0.004$ and $p < 0.001$, respectively). Parental years of education attained showed an overall effect ($F(3) = 3.11, p = 0.027$) but no significant differences between clusters. Furthermore, chlorpromazine equivalents were not significantly different between clusters ($p = 0.107$).

Clinical outcome
Although all patients were in clinical remission at baseline, the subgroup of patients with severely impaired cognition had significantly higher symptom severity compared to the cognitively pre-

served subgroup, with higher scores on the PANSS total subscale ($p < 0.001$), as well as the positive ($p = 0.014$), negative ($p < 0.001$) and general subscales ($p = 0.014$). Results are demonstrated in Fig. 3.

After correcting for clinical symptoms at baseline, the patient groups with severely impaired and preserved cognitive performance showed significant differences on PANSS negative symptomatol-ogy at 6- and 12-month follow-up ($n = 145, p = 0.017; n = 132, p = 0.018$, respectively). Those with severely impaired and moder-

ately impaired cognitive performance differed on the PANSS negative subscale (6 months: $p = 0.010$; 12 months: $p = 0.010$). Thus, consistently across time points, the group with severely impaired cognition was characterized by more severe negative symptoms compared to the other clusters at baseline, 6- and 12-month follow-up.

Furthermore, the patient subgroup with severely impaired cog-

nition had lower clinical global functioning (total GAF score, Fig. 4) compared to patients with relatively preserved cognition, at baseline ($p = 0.001$), and trend-level effects were shown for 6-month follow-up ($n = 144; p = 0.094$) and 12-month follow-up ($n = 132; p = 0.052$), corrected for global functioning at baseline. In addition, lower clinical global functioning was shown in the subgroup with severely impaired cognition compared to the moder-

ately impaired cluster at baseline ($p = 0.045$) and 6-month follow-up ($p = 0.047$).

Functional outcome
Self-reported global functioning and disability were evaluated by the WHODAS 2.0. Although the clusters did not significantly dif-

fer across all time points, corrected for global functioning and dis-

ability at baseline (all $p > 0.05$), there was a gradual and stepwise increase in disability, with the relatively preserved cluster having lower disability scores compared to the moderately impaired and severely impaired cluster.

Discussion
To the best of our knowledge, this is the largest study investigating cognitive subgroups of FEP patients who all reached symptomatic remission after treatment in relation to longitudinal clinical and functional outcomes. We found three distinct cognitive subgroups in a sample of FEP, including one relatively large subgroup with preserved cognition (37.2%), one moderately impaired group (36.3%) and one severely impaired group (26.5%) as compared to healthy controls. Of note, the severely impaired group included only one-fourth of the sample. The cognitive subgroups were characterized by significant differences in clinical symptoms, with more severe clinical symptoms in the severely impaired cogni-

tive cluster compared to the relatively preserved cluster, at baseline (PANSS total and all subscales) and 6- and 12-month follow-up (PANSS negative subscale). In addition, evaluation of global functioning (GAF) was significantly higher in the relatively preserved cluster compared to the severely impaired cluster at baseline and showed trend-level effects at 6- and 12-month follow-up. No significant differences in self-reported measures of functional outcome (WHODAS 2.0) were found between the patient subgroups at baseline and follow-up, yet the same trend could be observed.

The current results provide support for cognitive heterogeneity in FEP, delineated by three cognitive subtypes. This is consistent with previous clustering studies reporting on three subgroups of cognition in both first episode and chronic samples of schizophrenia (Carruthers et al. 2019; Gilbert et al. 2014; Menkes, Armstrong, Blackford, Heckers, & Woodward, 2019; Sauvé, Malla, Joober, Brodeur, & Lepage, 2018; Uren et al. 2017; Wells et al. 2015). The relatively preserved subgroup did not perform worse on overall cognition compared to the healthy controls, confirming the existence of a subset of patients with relatively intact cognitive performance (Ammari et al. 2014; Carruthers et al. 2019; Menkes et al. 2019; Moritz et al. 2017; Uren et al. 2017). Although cognitive impairment has long been recognized as a core symptom of psychotic disorders, our results show that a significant proportion of patients (37.8%) perform in the same range as healthy controls. This underscores the importance of taking individual variability into account in both research and clinical practice. It should be noted that cognitive performance similar to that of healthy controls is not necessarily synonymous with cognitively unaffected. However, no differences in years of educa-

tion were observed between the relatively intact subgroup and healthy controls, and no decline relative to parents’ years of edu-

cation was observed, suggesting that cognitive functioning did not decline relative to a higher premorbid level (Keefe, Eesley, & Poe, 2005). We further showed that both the moderately and severely impaired subgroups had attained significantly fewer years of edu-

cation compared to the relatively preserved subgroup. The moder-

ately impaired subgroup showed global cognitive impairment compared to the healthy controls, including all subdomains except for executive function. Findings regarding the intermediate cluster show global impairments of cognitive performance rather than domain-specific deficits. This is in line with previous studies performed in both first episode and chronic schizophrenia samples, which identified an intermediate cluster with overall moderate cog-

nitive impairment (Lewandowski, Sperry, Cohen, & Öngür, 2014; Uren et al. 2017; Van Rheenen et al. 2016). The severely impaired subgroup (25.5%) showed pronounced cognitive impairments that were not restricted to specific domains, with more severe perform-

ance deficits compared to the other cognitive subgroups. The exist-

ence of a severely impaired cognitive subgroup has been previously demonstrated (Lewandowski et al. 2014; Uren et al. 2017; Van Rheenen et al. 2016). However, the percentage of individuals show-

ing severely impaired cognition in this study is lower than the 44% reported in a large recent systematic review (Carruthers et al. 2019). Remarkably, executive function was relatively spared across all sub-

groups of cognition, although previous FEP studies demonstrated
Table 1. Mean (s.a.) baseline demographic and cognitive characteristics for FEP cognitive clusters and healthy controls

<table>
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<th>Test statistic</th>
<th>df</th>
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<th>Post hoc analyses*</th>
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### Male, n (%)
- Healthy controls (n = 40): 32 (80.0%)
- Preserved cognition (n = 76): 47 (61.8%)
- Moderately impaired cognition (n = 74): 60 (81.1%)
- Severely impaired cognition (n = 54): 41 (75.9%)

### Age
- Healthy controls (n = 40): 24.48 (4.98)
- Preserved cognition (n = 76): 29.62 (10.26)
- Moderately impaired cognition (n = 74): 26.54 (7.26)
- Severely impaired cognition (n = 54): 27.46 (8.65)

### Years of education
- Healthy controls (n = 40): 14.95 (1.95)
- Preserved cognition (n = 76): 14.75 (1.93)
- Moderately impaired cognition (n = 74): 13.42 (2.15)
- Severely impaired cognition (n = 54): 12.84 (3.28)

### Years of education parents
- Healthy controls (n = 40): 13.63 (2.20)
- Preserved cognition (n = 76): 13.61 (2.68)
- Moderately impaired cognition (n = 74): 12.52 (3.87)
- Severely impaired cognition (n = 54): 12.15 (3.06)

### Chlorpromazine equivalent
- Healthy controls (n = 40): N.A.
- Preserved cognition (n = 76): 210.11 (127.24)
- Moderately impaired cognition (n = 74): 258.62 (139.43)
- Severely impaired cognition (n = 54): 249.88 (145.86)

### BACS, Z-score
- Composite score: 0.13 (1.15)
- Verbal memory: 0.43 (1.01)
- Working memory: 0.07 (1.05)
- Motor speed: −0.15 (0.96)
- Verbal fluency: 0.11 (1.05)
- Attention & Processing speed: −0.23 (1.19)
- Executive function: 0.24 (0.87)

### FEP, first-episode psychosis; BACS, brief assessment of cognition in schizophrenia; df, degrees of freedom.
* a HC significantly different from the relatively preserved cluster; b HC significantly different from the moderately impaired cluster; c HC significantly different from the severely impaired cluster; d relatively preserved cluster significantly different from moderately impaired cluster; e relatively preserved cluster significantly different from severely impaired cluster; f moderately impaired cluster significantly different from a severely impaired cluster.
reduced executive function compared to healthy controls (Kravariti et al. 2009). Attention and speed of processing showed most severe impairments across all subgroups, which is in line with previous studies performed in FEP (Kravariti et al. 2009; Leeson et al. 2010; Weinberg et al. 2016).

Our finding of more severe clinical symptoms, specifically negative symptoms in the group with severely impaired cognition is in line with previous research demonstrating an association between cognitive function and negative symptoms in both FEP (Engen et al. 2019; Reser, Allott, Killackey, Farhall, & Cotton, 2015; Uren et al. 2017) and chronic schizophrenia (Lewandowski et al. 2014; Weinberg et al. 2016; Wells et al. 2015). However, the relationship between cognitive function and negative symptoms seems complex. Severe negative symptoms such as lack of motivation or decreased effort may impact cognitive performance but similarly, cognitive impairment could affect the manifestation of negative symptoms as more preserved cognitive function may be essential for the ability to plan, initiate, motivate and carry out daily activities (Beck, Himelstein, Bredemeier, Silverstein, & Grant, 2018; Fervaha et al. 2014; Fortgang, Srihari, & Cannon, 2020; Jurado & Rosselli, 2007; Lindgren et al. 2020). More longitudinal studies are required to gain more insight into the relationship between cognitive function and negative symptoms in FEP.

In the subgroup of individuals with severely impaired cognition, we found lower objectively evaluated global functioning (GAF) when compared to the relatively preserved subgroup. These findings are substantiated by other studies suggesting that global functioning is related to cognitive cluster membership (Gilbert et al. 2014; Lewandowski et al. 2014; Uren et al. 2017; Wells et al. 2015). Moreover, studies investigating cognitive subtypes in both psychotic patients and unaffected siblings showed that patients with cognitively impaired siblings reflect a poorer course of the disease. This suggests that cognitive impairment may indeed be predictive for the course of illness (Burger et al. 2021; Quee et al. 2014). However, no significant differences between cognitive subgroups could be demonstrated on self-reported measures of functional outcome (WHODAS 2.0). This is remarkable, as both the GAF and the WHODAS 2.0 assess measures of outcome. It is plausible that not all types of cognition are associated with the evaluation of functional outcomes. It has been suggested that not global cognition but specifically social cognition plays a critical role in outcome regarding everyday functioning. A recent study by Kim et al. (2021) demonstrated significant correlations between the WHODAS 2.0 and social cognition, such as communication and learning abilities (Kim et al. 2021). Similarly, Tan, Rossell, and Lee (2020) demonstrated that mostly verbal-linguistic cognitive skills such as semantics and language are associated with subjective measures of functioning and well-being, as those have a direct effect on community functioning (Tan et al. 2020). Indeed, medium to large associations between social cognition and community functioning have been reported in a meta-analysis (Fett et al. 2011), whereas only small to moderate associations have been reported between nonsocial cognition and functional outcome (Halverson et al. 2019). This suggests that interventions targeting social cognition may improve functional
outcomes more than neurocognitive interventions. Another explanation for the lack of differences in WHODAS 2.0 between the cognitive subgroups may be the lack of awareness of functioning and disability in patients as the accuracy of assessing daily functioning in patients with schizophrenia is under debate (Jongs et al. 2020). An overestimation of functioning by the patient may be affected by disease-related factors such as negative symptomatology and lack of insight (Jongs et al. 2020; Sabbag et al. 2012). This indicates that despite symptoms or restrictions in clinician observed functioning, patients may be satisfied with their lives and consider their level of functioning high. Thus, our findings suggest that daily functioning from a patient’s perspective is not necessarily synonymous with the clinician’s interpretation of recovery and may be related to a different set of predictors. Finally, the WHODAS 2.0 includes domains of daily functioning that are hardly affected in the current FEP sample and only minimally associated with cognitive function, such as mobility (getting around, standing up, walking a long-distance) and self-care (getting dressed, washing, eating) and hence do not differentiate between the groups (Chen et al. 2018).

Strengths, limitations and future directions

A strength of the current study is its large sample size and longitudinal design, evaluating both clinical and functional outcomes over a 12-month follow-up period. The participants were included shortly after diagnosis and had all achieved symptomatic remission before the baseline measurement. Therefore, factors that may influence cognitive function, such as long-term antipsychotic medication use or duration of illness, are being limited. In addition to the assessment of clinical outcome (PANSS and GAF) by trained central raters, we extensively measured functioning and disability with the self-reported WHODAS 2.0 questionnaire, which is recommended for the assessment of functioning in the DSM-5 (Gold, 2014). We also note that our study comes with some limitations. First, antipsychotic medication use was not stable for all participants throughout follow-up as some participants may have tapered off their antipsychotic medication gradually. However, the process of medication discontinuation also occurs in the general population of first-episode patients. Furthermore, although we did not include a cumulative dose of antipsychotic medication as a factor, all participants were in a similar early stage of the illness (3 to 6 months in remission of their first psychotic episode) at the time of inclusion and we found that current chlorpromazine equivalents were not significantly different between clusters. Moreover, although cognitive performance at baseline was not affected by psychotic symptoms as solely patients in symptomatic remission were included, generalizability to wider FEP populations may be limited within this study. Finally, cluster analyses come with the limitation that the
Fig. 3. PANSS mean scores illustrated for FEP cognitive clusters at baseline, 6-month follow-up and 12-month follow-up comparisons at 6- and 12-month follow-up were corrected for clinical symptoms at baseline. * illustrates \( p < 0.05 \); Error bars represent standard deviations. PANSS, Positive and Negative Syndrome Scale; FEP, first-episode psychosis.

Fig. 4. GAF mean scores illustrated for FEP cognitive clusters at baseline, 6-month follow-up and 12-month follow-up GAF, Global Assessment of Functioning; FEP, first-episode psychosis.
determination of the number of clusters may be arbitrary as it depends on the methods used. However, we followed the recommended guidelines for reporting on cluster analysis (Carruthers et al. 2019).

Conclusion

The results of the present study provide strong support for high heterogeneity in cognition among FEP patients who reach symptomatic remission. Besides finding a moderately impaired and severely impaired subgroup, we also show that a significant subset of patients have relatively preserved cognitive function. This underscores the importance of taking individual variability into account. In addition, we found that FEP patients with severe cognitive impairment have poor clinical outcomes compared to those with relatively preserved cognitive function. These findings suggest that grouping patients in subtypes along the cognitive continuum may offer crucial information about illness profiles and clinical prognosis. In conclusion, early identification of distinct cognitive profiles in FEP and corresponding longitudinal differences in clinical profile has clear implications for prognosis and personalized treatment of psychotic disorders. However, self-reported measures of functional outcome seem to have different sets of predictors in FEP and more longitudinal studies are required to further assess determinants of functional outcome.

Supplementary material. The supplementary material for this article can be found at https://doi.org/10.1017/S0033291721004153.

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Conflicts of interest. The authors report no potential conflicts of interest.

Ethical standards. The authors assert that all procedures contributing to this work comply with the ethical standards of the relevant national and institutional committees on human experimentation and with the Helsinki Declaration of 1975, as revised in 2008.

References


