Cognitive Change in Schizophrenia and Other Psychoses in the Decade Following the First Episode

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Abstract

Objective: Schizophrenia is associated with a large cognitive impairment that is widely believed to remain stable after illness onset. Yet, even to date, 10-year prospective studies of cognitive functioning following the first episode with good methodology are rare. We examined whether schizophrenia patients experience cognitive decline following the first episode, whether this decline is generalized or confined to individual neuropsychological functions, and whether decline is specific to schizophrenia.

Method: Participants were from a population-based, case-control study of patients with first-episode psychosis that were followed prospectively up to 10 years post first admission. A neuropsychological battery was administered at index presentation and at follow-up to patients with a diagnosis of schizophrenia (n=65), or other psychoses (n=41), as well as to healthy comparison subjects (n=103).

Results: The schizophrenia group exhibited declines in IQ and in measures of verbal knowledge, and memory, but not processing speed or executive functions. Processing speed and executive function impairments were already present at the first episode and remained stable thereafter. Magnitude of declines ranged between 0.28 and 0.66 standard deviations. Decline in measures of memory was not specific to schizophrenia and was also apparent in the group of patients with other psychoses. Healthy individuals with low IQ, on the other hand, showed no evidence of decline, suggesting that a decline is specific to psychosis.

Conclusions: Patients with schizophrenia and other psychoses experience cognitive decline after illness onset, but the magnitude of decline varies across cognitive functions. Distinct mechanisms consequent upon the illness and/or psychosocial factors may underlie impairments across different cognitive functions.
Introduction

Cognitive impairment is a core feature of schizophrenia (1, 2). Understanding the nature and course of this impairment may have important implications for our understanding of the pathophysiology of the disorder.

Research has shown that individuals diagnosed with schizophrenia experience cognitive decline from the premorbid to post-onset period. There is clear evidence for moderate cognitive deficits in children and adolescents who later develop schizophrenia, with meta-analyses showing an average premorbid deficit equal to 8 IQ points (0.5 Standard Deviation (SD)) (3, 4). Cognitive deficits in adults diagnosed with schizophrenia are more pronounced, with meta-analyses reporting a 14-point IQ deficit (0.90 SD) in first-episode schizophrenia patients (5) and 15- to 21-point IQ deficits (1.0 to 1.5 SD) in chronic schizophrenia patients (1, 6, 7). In line with cross-sectional evidence, longitudinal studies of cognitive change in schizophrenia from before to after illness onset have shown evidence for cognitive decline (8). Three population-based studies have reported cognitive declines ranging from 6 to 12 IQ points (0.4 – 0.8 SD) between childhood and adulthood in individuals later diagnosed with schizophrenia (8-10).

Despite evidence for cognitive decline from before to after illness onset, the course of cognitive decline in schizophrenia remains unclear. While it is widely believed that cognitive impairments stabilize after illness onset (11-13), at least until older adult life (12, 14), few longitudinal studies have examined cognitive change from illness onset through to a decade later (Table 1), and findings across studies and cognitive domains are mixed. Studies have reported a stabilization of the cognitive deficits, cognitive decline, as well as amelioration of cognitive functioning (Table 1 and ref # (15)).

Previous studies have been unable to comprehensively chart the course of cognitive deficits for several reasons. First, the majority of studies have used clinical samples, which may not be fully representative of the population of individuals with schizophrenia (8). Second, most studies followed participants for only 1 to 3 years from illness onset (Table 1). We previously reported a slow, gradual increase in premorbid cognitive deficits, with losses equal to between 0.5 and 1 IQ point per year (16). Studies with short follow-ups, therefore, may be underpowered to capture decline. Third, few studies have included comparison groups, and therefore have not considered the potential impact of normative age-associated changes in cognitive functioning, which is necessary to rigorously test for cognitive change. Since, brain maturation continues into the third decade of life (17), previous estimates of the magnitude of cognitive decline may be biased. Finally, few studies...
have examined the effect of medication on cognitive functioning, and yet recent findings suggest that antipsychotic medications may contribute to the severity of cognitive decline (18).

In a previous report on this population-based, case-control study, we provided evidence for an IQ deficit, as well as varying degrees of impairment across individual cognitive domains following the first psychiatric diagnosis of schizophrenia (19). Study participants have since been followed-up and underwent neuropsychological testing a second time. Using identical neuropsychological measures at first assessment and follow-up, we were able to directly examine change in IQ and in individual cognitive functions after the first episode. To provide an accurate estimate of cognitive change over time, we compared patients to the healthy comparison subjects in the study followed during the same period. We tested three hypotheses. First, we examined the “IQ decline” hypothesis to establish whether schizophrenia patients exhibit a static IQ deficit or IQ decline. Second, we tested the “generalized decline” hypothesis to determine whether decline occurs across multiple cognitive domains, namely verbal knowledge, memory, language, processing speed, executive function/working memory and visuospatial ability. Finally, we tested the “specificity” hypothesis to establish whether any cognitive decline is specific to schizophrenia or common to other psychoses by examining cognitive change in individuals with psychotic disorders other than schizophrenia.

Methods

AESOP Study

Data were derived from the Aetiology and Ethnicity in Schizophrenia and Other Psychoses (AESOP) study, a population-based, case-control study of first-episode psychosis. AESOP was approved by local research ethics committees and each participant gave written informed consent after receiving a complete description of the study. The study identified all first-episode psychosis cases (ICD-10: F10–F29 and F30–F33) aged 16 to 65 years presenting to specialist mental health services in tightly defined catchment areas of the United Kingdom (southeast London, Nottingham and Bristol) between September 1997 and August 2000. All potential cases making contact with psychiatric services (including adult community mental health teams, inpatient units, forensic services, learning disability services, adolescent mental health services, and drug and alcohol units) for the first time were screened. Exclusion criteria were previous contact with health services for psychosis, organic causes of psychotic symptoms, transient psychotic symptoms as the result of acute intoxication (as defined by ICD-10), and IQ<50. A random sample of control subjects with no past or present
psychotic disorder were recruited using a sampling method that matched cases and controls by area of residence. Hereafter, data collected at this phase of the AESOP study is referred to as ‘baseline’.

At baseline, detailed information was collected to enable patients to be traced, re-contacted and re-interviewed approximately 10 years later (‘follow-up’). At follow-up, patients currently in contact with mental health services were invited to participate through their clinical teams. Letters of invitation were sent to last known addresses of those not in contact with services. Non-responders were sent a second letter two to three weeks later. If patients were thought to have moved, contact was sought through their GP. Control subjects also provided contact details at baseline. Letters of invitation were sent and were followed-up with phone calls if no reply had been received within 2 weeks. If no reply had been received after 4 weeks, or where telephone numbers could not be obtained, in-person visits were made to the subject’s address. A detailed overview of the AESOP study design and methods, as well as the follow-up has been published elsewhere (20, 21).

Analytic Cohort

Derivation of the sample included in the present analysis is illustrated in Figure 1. The analytic cohort consisted of healthy comparison subjects and subjects who had a consensus ICD-10 diagnosis at last follow-up of schizophrenia (F20), bipolar disorder or mania (F30.2, F31.2, F31.5), depressive psychoses (F32.3, F33.3) or other psychotic disorders including persistent delusional disorders and psychosis NOS (F22, F23, F28, F29). Both case and comparison subjects were required to be native English speakers or to have migrated to the UK by age 11. The latter ensured that all participants had a good command of English, even as a non-native language, by verifying that participants had completed at least their secondary education in the UK. Thus, this minimized the effect of linguistic or cultural biases on cognitive performance in a multiethnic sample.

Figure 1. Derivation of first-episode psychosis patients and healthy comparison subjects from the Aetiology and Ethnicity in Schizophrenia and Other Psychoses (AESOP) Project Baseline and 10-Year Follow Up.

Neuropsychological assessment

At baseline and follow-up, participants underwent cognitive testing with a neuropsychological battery, which assessed general intellectual ability (IQ), as well as specific cognitive functions. Administration and scoring followed standard procedures. Full-scale IQ was estimated using the vocabulary, comprehension, digit symbol coding and block design subtests of the WAIS-R (22).
forms of the WAIS-R have been shown to produce accurate estimates of full-scale IQ (23, 24).

Specific functions were assessed using the following neuropsychological tests: Memory using the Rey Auditory-Verbal-Learning Test (RAVLT) trials 1 to 7 (learning, immediate and delayed verbal recall) (25), and the Visual Reproduction subtest of the Wechsler Memory Scale - Revised (WMS-R) (26); Verbal knowledge using the Vocabulary and Comprehension subtests of the WAIS-R (22); Processing speed using the WAIS-R digit symbol coding and the Trails-Making-Test Part A (27); Executive function/working memory using Trails-Making-Test - Part B (27), and Letter-Number Span (28); Language using Category (semantic) and Letter Fluency (categories: ‘body parts’; ‘fruits’; ‘animals’, letters: F; A; S) (29), and Visuospatial ability using the WAIS-R Block Design subtest.

Diagnostic Assessment

Clinical data were collected using the Schedules for Clinical Assessment in Neuropsychiatry (SCAN)(30). The SCAN incorporates the Present State Examination, Version 10, to elicit symptom-related data at time of presentation. Ratings on the SCAN are based on clinical interview, case note review, and information from informants (e.g. health professionals, close relatives). Researchers were trained on the SCAN with a World Health Organization-approved course and reliability was established prior to commencement of the study using independent ratings of videotaped interviews. Rater agreement was evaluated using Kappa statistics, which ranged from 1.0 for psychosis as a category to between 0.6 and 0.8 for individual diagnoses. ICD-10 diagnoses were determined using SCAN data through consensus meetings with one of the PIs and other team members. Symptom severity was classified based on the SCAN Symptom Severity Rating Scale 2 as: 0 = Absent, 1 = Mild, 2 = Moderate and 3 = Severe (21).

Covariates and medication information

Age was collected at baseline and follow-up. Sex, ethnicity, and level of education were collected at baseline. Treatment history with typical and/or atypical antipsychotic medication was ascertained for all patients from interview data and record review at follow-up.

Creating Norms for Neuropsychological Tests

A regression-based approach was used to create normative standards for the neuropsychological tests. Age at assessment, sex, ethnicity, and education were regressed on each of the neuropsychological measures in the healthy comparison sample at baseline and follow-up. Next, scores were adjusted on the basis of the regression results, and standard scores (i.e., z-scores) were
created. The same adjustment and standardization procedure were applied to the patient groups, using the normative standards from the healthy comparison group.

### Statistical analysis

Demographic and clinical characteristics of the baseline and follow-up cohorts were compared using summary statistics. For descriptive purposes, we compared patients with schizophrenia or other psychoses (including bipolar disorder, mania, depressive psychoses and other psychotic disorders) to the comparison group on normative-adjusted IQ and specific neuropsychological tests at baseline and follow-up using analysis of variance (ANOVA) models.

To examine the “IQ decline” “generalized decline” and “specificity” hypotheses, we compared the schizophrenia and other psychoses groups to the comparison group on change in normative-adjusted IQ and specific neuropsychological tests from baseline to follow-up. Change scores were calculated by subtracting follow-up test scores from baseline test scores, so that positive scores indicate cognitive amelioration and negative scores indicate cognitive decline. ANCOVA models with planned orthogonal comparisons of each psychosis group to the comparison group, adjusting for time from baseline assessment and baseline test score were used. Adjustment for baseline performance is common in studies on cognitive change (31, 32). For the “IQ decline” hypothesis, the significance level was set at \( p=0.05 \) (two-sided). For the “generalized decline” hypothesis, the significance level was set at a Bonferroni-corrected level of 0.0038 \((0.05/13)\). All analyses were conducted using IBM SPSS Statistics version 24.

### Results

Demographic characteristics of the baseline cohort and the cohort assessed at follow-up are presented in Table 1. Follow-up neuropsychological assessments were completed on 106 patients (63 males), and 103 comparison subjects (40 males). Average follow-up duration was 109.3 months (SD=29.5) for patients and 102.9 (SD=34.1) for comparisons. Overall, the patients and comparisons assessed at follow-up were similar to the respective patients and comparisons assessed at baseline in terms of demographic variables, suggesting that the cohort at follow-up was representative of the original cohort.

**Cognitive impairment in schizophrenia and other psychoses at baseline and follow-up**

As we have previously shown in the AESOP study cohort (19), patients with schizophrenia and
patients with other psychoses showed deficits in IQ and individual neuropsychological tests at baseline. Figure 2 illustrates that schizophrenia patients exhibited widespread, persistent, cognitive impairment, performing significantly worse than comparison subjects at both baseline and follow-up on 11 out of the 14 measures. Patients with other psychoses also showed widespread impairments, but these were generally of smaller magnitude than schizophrenia patients (Figure 2).

(Table 2 presents the non-adjusted performance in IQ and specific neuropsychological tests at baseline and follow-up)

Figure 2. Neuropsychological Performance Among Patients with Schizophrenia and Other Psychoses at Baseline and Follow-Up.

Effect sizes (expressed in standardized {z} scores) and 95% Confidence Intervals (95%CI) of difference from comparison subjects at baseline and follow up. Comparison subjects set to zero (dotted line). Effect sizes are adjusted for age, sex, ethnicity, and level of education. 95% CI that do not include zero indicate statistical significance level p<0.05. Trailmaking A=Trail Making Test, Part A; Trailmaking B=Trail Making Test, Part B.

Cognitive change in schizophrenia and other psychoses

Next, we compared cognitive change over time in each of the psychoses groups (schizophrenia and other psychoses) to cognitive change in controls to test the “IQ decline”, “generalized decline” and “specificity” hypotheses. Figure 3 presents effect sizes of the difference in the within group change from baseline to follow-up in IQ and individual neuropsychological tests between the psychoses groups and controls. Effect sizes of 0.20, 0.50, and 0.80 reflect small, medium, and large effects, respectively (33).

**IQ decline hypothesis**: IQ decline in the schizophrenia group was significantly larger than in controls, who showed no evidence of IQ decline. The IQ decline in the schizophrenia group compared to controls was of small magnitude (ES=-0.28, 95% Confidence Intervals: -0.47 to -0.09, p=0.003), but was not attenuated when adjusting for education, ethnicity, sex, age-at-baseline assessment, or duration of follow-up, suggesting that IQ decline could not be attributed to these variables.

**Generalized decline hypothesis**: Compared to controls, the schizophrenia group showed a larger cognitive decline across tests in the memory and verbal knowledge domains (Figure 3). In the memory domain, the schizophrenia group declines on verbal learning (p=0.001), immediate recall (p<0.00006), and delayed recall (p<0.00001) reached the Bonferroni-corrected level of significance. In the verbal knowledge domain, decline on vocabulary (p=0.003) reached the Bonferroni-corrected level of significance. Compared to controls, the schizophrenia group showed no significant cognitive
changes on Digit Symbol Coding and Trail-making-test Part A in the processing speed domain, Block Design in the visuospatial domain, and Trail-making-test Part B, Letter-Number Span, Letter Fluency and Category Fluency in the executive functions and working memory domain.

**Specificity hypothesis:** There was no evidence for IQ decline in the other psychoses group compared to controls (ES=-0.09, 95% Confidence Intervals: -0.30 to 0.11; p=0.37), (Figure 3). In terms of cognitive domains, like the schizophrenia group, the other psychoses group showed larger declines than controls across test in the memory domain, with verbal learning (p=0.001) reaching the Bonferroni-corrected level of significance. Like schizophrenia patients, the other psychoses group showed static deficits in tests of processing speed, executive functions and working memory, and visuospatial ability (Figure 3).

*Figure 3. Change in Neuropsychological Performance Among Patients with Schizophrenia and Other Psychoses*.

*a* - Presented are effect sizes and 95% Confidence Intervals of difference in change from baseline to follow up between the diagnostic group and comparison group. 95% Confidence Intervals that do not include zero indicate statistical significance level p<0.05. Effect sizes are adjusted for age, sex, ethnicity, level of education, time from baseline assessment and baseline test score. Trailmaking A=Trail Making Test, Part A; Trailmaking B=Trail Making Test, Part B.

* - Presents Bonferroni corrected level (p≤0.003)

**Medication**

We examined the potential moderating effect of antipsychotic medication on IQ decline in the schizophrenia group. There was no statistically significant difference in IQ decline (p=0.23) between patients with a history of treatment with typical antipsychotics only (45% of sample) and those with a history of treatment with both typical and atypical antipsychotics (55% of sample). Duration of antipsychotic medication (mean = 323±192 weeks) did not attenuate IQ decline in schizophrenia (F=7.30, p=0.008 vs. F=7.20, p=0.009 for ANCOVA models with vs. without duration of treatment as a covariate).

**Symptom severity**

Since illness severity might influence cognition, we also examined the association between baseline symptom severity and change in cognitive functioning, as well as change in symptom severity between baseline and follow up and change in cognitive functioning. Schizophrenia patients with severe symptoms at baseline showed statistically significantly greater cognitive decline than patients.
with mild or moderate symptoms across multiple tests in the memory domain (Figure 4). However, there was no association between change in symptom severity and change in cognitive functioning (Table 2 and Figure 1), and no evidence for a dose-response relationship across levels of severity (Figure 4). In the other psychoses group there was no evidence for an association between symptom severity, or change in symptom severity, and change in cognitive functioning (Figure 4, Figure 1).

**Figure 4.** Change in Neuropsychological Performance Among Patients With Schizophrenia and Other Psychoses in Relation to Symptom Severity at Baseline

- Presented are effect sizes and 95% Confidence Intervals of difference in change from baseline to follow up between the diagnostic group and comparison group as a function of symptom severity at baseline. 95% Confidence Intervals that do not include zero indicate statistical significance level p<0.05. Effect sizes are adjusted for age, sex, ethnicity, level of education, time from baseline assessment and baseline test score. Trailmaking A=Trail Making Test, Part A; Trailmaking B=Trail Making Test, Part B.

**Sensitivity analyses**

We also examined the potential impact of attrition by applying linear mixed models which permit varying numbers of measurements per person and time point, while adjusting for within-individual (i.e. between measures) variation. Similar results were obtained in models that included only cases and controls with data from both assessment time points, and in models that also included cases and controls with data from a single assessment, indicating results were not biased by attrition.

As a further comparison, we examined IQ change in controls with lower IQ (IQ<90 at baseline, equal to 1SD below the control group mean, N=17, 16.5% of sample). These individuals are of interest because, like schizophrenia patients, they also exhibit lower IQ, and yet they did not develop psychosis. In contrast to patients with schizophrenia, individuals with lower IQ did not show evidence of IQ decline, neither in absolute terms nor relative to controls without a cognitive impairment since mean IQ at baseline was 84.9, and at follow up was 89.8 (F=0.97, p=0.35).

**Discussion**

Using a population-based, case-control sample followed prospectively from the first psychotic episode we provide evidence for cognitive decline after illness onset in patients with schizophrenia. These findings advance knowledge in three important ways. First, the results lend support to the “IQ
As a group, schizophrenia patients showed IQ decline between baseline and follow up assessments, with an effect size of small magnitude (ES=0.28). This finding is in contrast with earlier studies reporting stabilization of cognitive deficits after the onset of psychosis (15). However, previous studies had important methodological limitations, including a short follow-up period of patients, and lack of a comparison group that is similarly followed-up. The finding of IQ decline is in line with findings from neuroimaging studies of greater age-associated brain volume loss (34), as well as deviated gyriﬁcation trajectories in schizophrenia patients in adulthood (35).

Moreover, reduction in cortical volume has been associated with IQ decline in schizophrenia patients (36).

Second, the current findings do not support the “generalized decline” hypothesis. Decline was not ubiquitous and varied across cognitive domains. The schizophrenia group exhibited declines in verbal knowledge and memory. In contrast, processing speed, executive functions and visuospatial ability did not decline. These contrasts can be generally viewed as reﬂecting differences between the impact of the illness on crystallized (verbal knowledge) vs. ﬂuid (processing speed, executive functions, visuospatial) abilities. Our ﬁndings of decreasing crystallized abilities and memory scores between baseline and follow-up is in line with previous evidence (37) and suggest that increasing deﬁcits in these domains may reﬂect actual loss of ability, rather than abnormal cognitive development (i.e. “lag”) (16). Alternatively, our ﬁndings may reﬂect diﬃculties with the maintenance and acquisition of new verbal knowledge due to substantial and increasing memory deﬁcits. While most cognitive abilities in the general population start to show stabilization or even decline in early adulthood, crystallized abilities may peak much later (38-40). In our study, measures of ﬂuid abilities showed a large deﬁcit already at the ﬁrst episode, which remained static thereafter. While previous longitudinal epidemiological studies have shown cognitive decline in schizophrenia from the premorbid period in childhood to the chronic stage in mid-adulthood (8-10), they were unable to determine when this decline occurred. Our ﬁndings suggest that most of the decline in ﬂuid abilities occurs before the ﬁrst episode, while crystallized abilities may continue to decline after onset.

Importantly, the decline in IQ after onset is likely to be due to the decline seen in crystallized abilities.

Third, the current ﬁndings do not support the “speciﬁcity” hypothesis since patients with schizophrenia, but also other psychoses, experienced cognitive decline. However, while patients with schizophrenia showed decline in IQ, memory and verbal knowledge, patients with other psychoses showed decline only in certain memory functions. Moreover, in line with previous reports (41, 42), the other psychoses group showed an overall impairment proﬁle that was qualitatively
similar, yet quantitatively smaller than the schizophrenia group. Thus, our findings suggest that
cognitive decline is not specific to schizophrenia, but also evident in other psychoses. However,
large, widespread, cognitive decline may still be specific to schizophrenia, since the other psychoses
group showed a smaller and less generalized cognitive decline. Interestingly, there was no evidence
of decline in a key comparison group, namely individuals with lower IQ who did not develop
psychosis. This group may in fact experience a different process of regression-to-the-mean.
The current findings should be viewed in the context of certain limitations. First, although we found
evidence for cognitive decline after illness onset, we could not fully map the course of deficits and
cognitive functions may vary in the timing of decline following the first episode. Second, group sizes
did not allow for an analysis of the heterogeneity of cognitive course and also limited our ability to
investigate more specific diagnostic sub-groups, such as bipolar/mania. Third, we ruled out two
explanations for the observed cognitive decline, namely, type or duration of antipsychotic
treatment. Unfortunately, we did not have information to examine other potential moderators of
cognitive decline, such as social isolation, smoking and illicit drug abuse, victimization, or physical
health problems such as obesity, diabetes and hypertensions. Moreover, despite the fact that we
adjusted for education in all our analyses, poor education in the schizophrenia group after the first
psychotic episode could still partly explain some of the group differences.
There is conflicting evidence regarding the relationship between change in symptoms and cognitive
functioning (43, 44). In our study, change in severity of psychosis was only minimally associated with
cognitive change. These results are consistent with cross-sectional findings of only a weak
association between positive symptoms and cognitive impairment (45). Longitudinal evidence also
suggests a minimal association between change in positive as well as negative symptoms, and
change in cognition (43, 44, 46). Interestingly, in our study, schizophrenia patients with severe
symptoms at baseline showed greater cognitive decline than patients with mild or moderate
symptoms. While this group was small (21% of overall group), the magnitude of decline in the
memory domain was large. Thus, this finding points to a potential subgroup of schizophrenia
patients that may greatly benefit from being specifically targeted for cognitive remediation.
Our findings have important implications for understanding the nature and course of cognitive
impairment in schizophrenia, as well as other psychoses. Integrating the current findings with those
of previous studies (16) suggests that cognitive dysfunction in schizophrenia may result from a
complex interplay between an early, static neuropathology (47, 48) and dynamic age-related
processes (49, 50). As such, cognitive functions that develop and peak relatively early in life, such as
processing speed and visuospatial abilities (39) may show aberrant development, resulting in slowed
growth prior to the onset of schizophrenia (16), but relative stabilization throughout the illness course. On the other hand, cognitive functions that continue to evolve through adult life, such as language (39), may show further deterioration throughout the course of schizophrenia. Finally, functions sensitive to age-related cognitive decline, such as memory, may begin to decline in middle adulthood before normative aging becomes apparent (40).

In conclusion, the present study demonstrates that while a substantial proportion of the cognitive impairment seen in adult patients with schizophrenia, as well as other psychoses, is present already at the first episode, these patients continue to experience cognitive decline after illness onset. However, the nature of this decline varies across neuropsychological functions. While large deficits in processing speed are already apparent at the first episode, deficits in verbal knowledge and memory continue to increase. These findings suggest that different pathophysiological mechanisms may underlie individual neuropsychological deficits seen in adult psychosis patients. Future research should determine which of these are consequent upon the illness itself, and which on the psychosocial factors patients experience. Finally, these findings highlight the importance of targeting early developmental stages in future studies of the causes of cognitive deficits associated with psychosis, as well as in cognitive remediation efforts.
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