Attenuated niacin-induced skin flush response in individuals with clinical high risk for psychosis

Ranpiao Gan, Yanyan Wei, Guisen Wu, Jiahui Zeng, Yegang Hu, Lihua Xu, Xiaochen Tang, Xiaohua Liu, Haichun Liu, Tao Chen, Jijun Wang, Tianhong Zhang

ABSTRACT

Background Impaired sensitivity of the skin flush response to niacin is one of the most replicated findings in patients with schizophrenia. However, prior studies have usually focused on postonset psychosis, and little is known about the clinical high-risk (CHR) phase of niacin sensitivity in psychosis.

Aims To profile and compare the niacin flush response among CHR individuals (converters and non-converters), patients with first-episode schizophrenia (FES) and healthy controls (HCs).

Methods Sensitivity to four concentrations (0.1–0.0001 M) of aqueous methylnicotinate was tested in 105 CHR individuals, 57 patients with FES and 52 HCs. CHR individuals were further grouped as converters and non-converters according to the 2-year follow-up outcomes. Skin flush response scores were rated on a 4-point scale.

Results Of the 105 CHR individuals, 21 individuals were lost during the study, leaving 84 CHR individuals; 16 (19.0%) converted to full psychosis at 2 years of follow-up. Flush response scores identified in the CHR samples were characterised as modest degree levels, intermediate between those of HC individuals and patients with FES. The flush responses in the CHR group mimicked the responses observed in the FES group at higher concentrations (0.01 M, 0.1 M and longer time points (15 min, 20 min); however, these became comparable with the responses in the HC group at the shorter time points and at lower concentrations. The converters exhibited lower mean flush response scores than the non-converters.

Conclusions Attenuated niacin-induced flushing emerged during the early phase of psychosis. New devices should be developed and verified for objective quantification of skin responses in the CHR population.

INTRODUCTION

While there is obvious heterogeneity in the clinical manifestations and mechanisms in patients with psychosis,1 biological markers for the early identification of psychosis remain undiscovered.2 In particular, trait markers with endophenotypic characteristics3 4 that can be applied in the premorbid phase to increase power by predicting the outcome of conversion to psychosis are desirable. Blunted skin flush response to niacin has been widely reported in patients with psychosis,3–7 and nonpsychotic first-degree relatives.8 9

Attenuated niacin response in patients with psychosis has been consistently reported in previous studies,6 10 and such abnormalities are heritable traits within psychosis-affected families.11 Other indirect evidence has shown that diminished flush responses are stable across time and depend on the stage of illness.12 These findings imply that the attenuated niacin flush response may be a useful trait for the early identification of psychosis.
Although there are many hypotheses regarding the pathogenetic mechanism of biochemical metabolism-related factors in psychosis, the mechanism of the attenuated skin flush response to niacin has been studied in-depth and found to be associated with disordered signalling in the phospholipase A2 (PLA2)-arachidonic acid (AA)-prostaglandin cascade. Niacin interacts with a specific G-protein-coupled receptor, leading to the active release of AA, which is mediated by PLA2 located at cell membranes. AA interacts with cyclo-oxygenase (COX), leading to its conversion to vasodilator prostaglandins (figure 1). Thus, there is a subgroup of patients with psychosis characterised by an attenuated niacin flush response linked to the disordered GPR109A-COX-prostaglandin pathway, which can be used as a biomarker for predicting a subtype of psychosis from the premorbid phase.

With rapid progress being made around the world in the identification of individuals at clinical high risk (CHR) for psychosis, there is now hope that psychosis can be treated to postpone and prevent the conversion to a first psychotic episode. Among individuals with CHR, about 20% are at risk of conversion within 2 years; efforts are currently underway to refine risk identification strategies to increase their predictive power. While there have been promising results from blunted niacin-induced skin flush responses in psychosis, whether this biomarker has early warning value in the prodromal phase of psychosis, such as in the CHR population, remains unknown. In 2017, Langbein et al. compared the visual ratings of niacin sensitivity between CHR adolescents and healthy controls (HCs) and first-episode schizophrenia (FES) subjects. They found that the niacin sensitivity of the entire CHR group was significantly increased compared with the HC group, whereas there was no difference between converters and non-converters after 1-year follow-up.

In this study, we employed a larger CHR cohort (105 CHR individuals with a 2-year follow-up) from the Shanghai At-Risk for Psychosis-extended Program (SHARP-extended), a disease control population with FES, and an HC population. We used a 4-point scale measurement to compare the niacin flush responses in different stages of psychosis, with the aim of identifying the biomarker properties of niacin flush responses, namely, the differential response in the FES, CHR and HC groups. The identification of such CHR subgroups will hopefully lead to the development of a method for the precise prediction and personalised prevention of psychosis in the future.

METHODS
Sample and procedures
This was an observational study of 105 individuals who were confirmed to have CHR through face-to-face interviews. A total of 57 patients with FES were included as the disease control group and 52 subjects were used as HCs. The CHR individuals were from the SHARP-extended...
Figure 2  Flowchart showing study subjects selection. CHR, clinical high risk; FES, first-episode schizophrenia; HC, healthy controls.

98 patients with FES referred from Shanghai Mental Health Center between 2016-2017

169 individuals with CHR for psychosis referred from Shanghai Mental Health Center between 2016-2017

52 HCs recruited from local Shanghai community

41 excluded:
• 28 refused consent
• 13 did not meet inclusion criteria

64 excluded:
• 14 refused consent
• 50 did not meet inclusion criteria

57 FES patients enrolled

105 CHR individuals enrolled

52 HCs enrolled

21 CHR individuals lost in 2-yr follow-up:
• 11 declined for lack of time
• 4 unwilling to re-assess
• 6 unable to be reached

84 CHR subjects completed 2-yr follow-up

16 converted to psychosis

68 did not convert to psychosis

cohort; their samples were recruited between 2016 and 2021 at the Shanghai Mental Health Center (SMHC) in China. All participants agreed to participate in the study. Subjects younger than 18 years of age had their consent forms signed by their parents, and the youths gave informed assent. The patients fulfilled at least one of the prodromal syndrome criteria: (1) Brief intermittent psychotic syndrome, (2) Attenuated positive symptom syndrome, or (3) Genetic risk and deterioration syndrome. The inclusion criteria were as follows: (1) Age younger than 45 years old; (2) Individuals younger than 18 years of age had to be accompanied by either a parent or legal guardian; (3) Capacity to provide informed consent or assent if under 18 years; (4) Must have completed at least 6 years of primary education; and (5) Psychotropically naïve. The exclusion criteria were: (1) Severe somatic diseases, such as pneumonia, cancer or heart failure; (2) Intellectual disability; or (3) History of drug (such as methamphetamine) abuse or dependence. Zhang et al provided further details regarding the SHARP methodology.

The research procedure was independent of the routine clinical treatment procedures at the SMHC. In the present study, of the 105 CHR individuals who completed the baseline assessment, 21 individuals were lost, and 84 individuals completed both the baseline assessment and the 2-year follow-up (figure 2). Recruited CHR individuals were followed up every 6 months until the end of 24 months and reassessed by telephone or face-to-face interview every 6 months using the Structured Interview for Prodromal Syndromes (SIPS). The patients with FES met the Diagnostic and Statistical Manual of Mental Disorders-IV (DSM-IV) diagnostic criteria for schizophrenia through the Structured Clinical Interview for DSM-IV Axis I Disorders (SCID-I) interview and had their first psychotic episode in the past year. Fifty-two HC individuals with a negative family history of mental disorders were recruited from the local community in Shanghai and matched to the FES group according to age, sex and education.

Clinical measurement and outcome criteria

The SIPS was used to identify individuals with CHR. It consists of 19 items that assess four symptom domains: positive symptoms (scales P1–P5: P1 unusual thought content; P2 suspiciousness; P3 grandiosity; P4 perceptual abnormalities; and P5 disorganised communication), negative symptoms (scales N1–N6: N1 social anhedonia; N2 avolition; N3 expression of emotion; N4 experience of emotions and self; N5 ideational richness; and N6 occupational functioning), disorganised symptoms (scales D1–D4: D1 odd behaviour or appearance; D2 bizarre thinking; D3 trouble with focus and attention; and D4 impaired personal hygiene), and general symptoms (scales G1–G4: G1 sleep disturbance; G2 dysphoric mood; G3 motor disturbances; and G4 impaired tolerance to normal stress). During the SIPS interview, the global assessment of function (GAF) was used to measure the participants’ global psychological, social, and occupational functioning. Decreases in GAF scores were used to assess functional deterioration (ie, scores compared with the scores 12 months prior) during the SIPS interview.

Conversion to psychosis was the major outcome of this study. Of the remaining 84 CHR individuals, 16 (19.0%)...
Table 1  Demographic and clinical variables, comparison among FES, CHR and HC groups

|                          | FES          | CHR          | HC           | Comparison |
|--------------------------|--------------|--------------|--------------|------------|
| **Cases, n**             | 57           | 105          | 52           | –          |
| **Demographic variables**|              |              |              | F=31.5 (<0.001) | χ²=1.303 (0.521) | F=41.2 (<0.001) | F=1.6 (0.206) | F=2.4 (0.093) |
| Age (year), mean (SD)    | 24.4 (8.5)   | 18.3 (4.8)   | 27.3 (9.2)   |            |
| Male, n (%)              | 30 (52.6)    | 46 (43.8)    | 26 (50.0)    |            |
| Education (year), mean (SD) | 11.8 (3.2)   | 10.3 (2.8)   | 16.2 (5.7)   |            |
| Height (cm), mean (SD)   | 168.1 (7.2)  | 167.1 (8.0)  | 167.0 (7.4)  |            |
| Weight (kg), mean (SD)   | 59.5 (9.9)   | 58.1 (11.4)  | 61.3 (9.7)   |            |
| BMI (kg/m²)              | 21.0 (3.3)   | 20.7 (3.4)   | 21.9 (2.6)   |            |
| **PANSS variables**      |              |              |              |            |
| PANSS_P                  | 22.8 (5.5)   | –            | –            |            |
| PANSS_N                  | 16.9 (8.1)   | –            | –            |            |
| PANSS_G                  | 44.2 (6.9)   | –            | –            |            |
| PANSS_TOTAL              | 83.8 (15.1)  | –            | –            |            |
| **SIPS variables**       |              |              |              |            |
| SIPS_P                   | –            | 10.5 (3.4)   | –            |            |
| SIPS_N                   | –            | 12.1 (5.7)   | –            |            |
| SIPS_D                   | –            | 6.5 (2.8)    | –            |            |
| SIPS_G                   | –            | 9.6 (2.8)    | –            |            |
| SIPS_TOTAL               | –            | 38.0 (10.1)  | –            |            |

BMI, body mass index; CHR, clinical high risk; FES, first-episode schizophrenia; HC, healthy control; PANSS, Positive and Negative Syndrome Scale; PANSS_G, scores of general psychopathology scale in PANSS; PANSS_N, scores of negative symptoms scale in PANSS; PANSS_P, scores of positive symptoms scale in PANSS; PANSS_TOTAL, total scores of PANSS; SD, standard deviation; SIPS, Structured Interview for Prodromal Syndromes; SIPS-D, scores of disorganisation symptoms scale in SIPS; SIPS_G, scores of general symptoms scale in SIPS; SIPS_N, scores of negative symptoms scale in SIPS; SIPS_P, scores of positive symptoms scale in SIPS; SIPS_TOTAL, total scores of SIPS.

converted to full psychosis at 2 years of follow-up. Conversion to psychosis was defined using the presence of psychotic symptoms in the SIPS criteria. The conversion was defined as the development of at least one psychotic-level symptom (with a rating of ‘6’ on the SIPS positive symptoms scale) with either sufficient frequency or duration, or occurring at least an hour a day on average over days a week for at least 16 hours.

Measurement of the niacin response

Before the niacin skin patch test, participants were screened for the following criteria: no major systemic illness (especially heart disease, allergic skin illnesses and asthma), and no use of anti-inflammatory drugs (eg, aspirin, non-steroidal anti-inflammatory drugs and steroids) within 7 days before the test. The temperature may affect the skin’s response to niacin; therefore, the temperature in the test room was maintained at 25 °C. A round filter paper patch was used to apply niacin in the form of AMN. Different concentrations (0.1 M, 0.01 M, 0.001 M and 0.0001 M) of AMN solution were prepared on the same day of the test. To better set the reference distance, a sticky ruler was attached to the inner side of the participant’s forearm. Four wet paper patches from each of the four AMN solutions were applied to four neighbouring sites on the forearm skin for 1 min and then removed. The skin flush response was photographed from a fixed vertical view at 5 min, 10 min, 15 min and 20 min after patch removal. The skin flush response was rated as follows: 0, no erythema; 1, incomplete erythema; 2, complete erythema within the defined area of the patch; and 3, erythema plus oedema beyond the definite area of the patch. The scoring of skin flush response was performed first by a research assistant and then by a senior researcher. Any inconsistent scores were discussed and evaluated together. Both groups were blinded to the grouping information. The niacin-induced flushing response scores consisted of 16 score values obtained at different AMN concentrations (0.0001 M, 0.001 M, 0.01 M, 0.1 M) and at different time points (5 min, 10 min, 15 min and 20 min).

Statistical analysis

Demographic and baseline clinical features are presented separately. Quantitative variables are expressed as mean (SD), while qualitative variables are presented as frequencies (%). The two groups were compared using χ² tests for comparisons of categorical variables, rank-sum tests for comparisons of individual SIPS item scores and independent t-tests for comparisons of continuous variables.
dependent variables included the 16 raw scores obtained at different AMN concentrations (0.0001 M, 0.001 M, 0.01 M and 0.1 M) and at different time points (5 min, 10 min, 15 min and 20 min). CHR individuals were further grouped as converters and non-converters according to the 2-year follow-up outcomes. Multivariate analysis of variance (MANOVA) was conducted with age, education and group as the independent variables and the 16 raw scores in the skin flush response as the dependent variables for group comparisons and to assess covariate effects. Considering gender imbalance, the flush scores were compared using the MANOVA test for converters and non-converters. The level of statistical significance was set at a two-tailed p value of 0.05.

RESULTS
Sample characteristics
Sex, height and weight were not significantly different among the FES, CHR and HC groups. The FES and HC groups were much older than the CHR group. The mean SIPS scores for the CHR and Positive and Negative Syndrome Scale (PANSS) measures for the FES group are listed in table 1.

Comparison of the niacin response in the FES, CHR and HC groups
The MANOVA showed that the covariates of age and education had no impact on our effort to compare 16 raw scores in the skin flush response among three groups. The mean (SD) values of the niacin response for the FES, CHR and HC groups are shown in figure 3. Overall, many scores in the flush response were significantly different among the groups. The patients with FES exhibited lower mean flush response scores than the HC and CHR groups. Flush response scores identified in the CHR samples were characterised by a modest degree level, intermediate between those of HC individuals and patients with FES. The flush responses in the CHR group mimicked the responses observed in the FES group at the higher concentrations (0.01 M, 0.1 M) and longer time periods (15 min, 20 min); however, these became comparable with the responses in the HC group in the shorter time periods and lower concentrations.

Table 2 highlights the significant differences between converters and non-converters in the baseline demographic variables. Male subjects, individuals with lower baseline or current GAF scores, and individuals with higher total SIPS scores were more likely to convert to psychosis during the follow-up. Comparisons of the baseline demographic, clinical and niacin-induced flushing response variables between those who quit and those who completed follow-up can be found in the online supplemental table S1. No significant differences in demographic and clinical characteristics between these two groups were found. However, significant differences were observed for the niacin-induced flushing response between the two groups. The CHR individuals in the former showed lower mean flush response than those subjects who completed the follow-up.

The mean scores of the flush responses for converters and non-converters are shown in figure 4. Overall, the converters exhibited lower mean flush response scores than the non-converters. The flush responses in the converter group differed from those observed in the non-converter group at the highest concentration (0.1 M); however, they became comparable at shorter time points and at lower concentrations.

DISCUSSION
Main findings
The first aim of this study was to investigate niacin sensitivity in CHR individuals compared with patients with FES, who met the criteria for schizophrenia for the first time, and with HCs. We found that niacin sensitivity of our CHR sample was attenuated, but to a lesser extent than in FES subjects. At higher concentrations or longer time points, the flush responses in the CHR group were
close to the responses observed in the FES group, while at shorter time points or at lower concentrations, the flush responses became comparable with the responses in the HC group. Second, we investigated whether baseline niacin sensitivity differed between those who progressed towards psychosis (converters) versus those who did not (non-converters). We found that niacin sensitivity was more attenuated in the CHR converters than in the non-converters, especially at the highest concentration. To the best of our knowledge, this is the first attempt to investigate the niacin-induced flush response in a relatively large-scale CHR cohort in a Chinese clinical setting.

Compared with the FES and HC groups, our data showed that attenuated flush responses were identified in the CHR samples; these were characterised by a modest level of severity that was intermediate between HC individuals and patients with FES. Specifically, at a shorter time point (10 min), the flush responses in the CHR group were close to those in the HC group, while at longer durations and greater concentrations, differences between the CHR and FES groups were not observed, which implied that attenuated flush responses might already be present prior to the onset of full psychosis. Compared with HCs, patients with FES showed blunted flush responses, which was consistent with previous studies. Furthermore, the results in the CHR group in the current study replicate previous findings by Langbein et al in a different CHR sample. A novel finding is that niacin-induced flush responses appear to be attenuated in the CHR group, with similar alterations to the HC group at the lower concentrations and shorter time points but similar alterations to the FES group at higher concentrations and longer time points. This could indicate that the application of the niacin test in risk groups needs to be tailored to improve the accuracy of identification.

In the context of CHR follow-up research, the association of a biological marker such as attenuated flush responses and conversion outcomes is particularly valuable, as it helps to assign a specific pathological mechanism (and its extent) to refine risk identification strategies to increase predictive power for the CHR population. We identified several flush response scores that were significantly different between converters and non-converters, with converters showing a lower response to niacin than non-converters and more similar responses to those in patients with FES. Although the sample size of

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Table 2  Baseline demographic and SIPS variables, comparison between converters and non-converters.

| Variables                          | Lost | Converters | Non-converters | Comparison |
|------------------------------------|------|------------|----------------|------------|
|                                    | Cases, n |           |                | T/Z/χ²/κ² | P value |
| Demographic variables             | 21   | 16         | 68             |           |         |
| Age (year), mean (SD)              | 17.7 (3.8) | 19.4 (5.5) | 18.2 (4.9)     | t=0.886   | 0.378   |
| Male, n (%)                        | 9 (42.9) | 11 (68.8)  | 26 (38.2)      |           |         |
| Female, n (%)                      | 12 (57.1) | 5 (31.3)   | 42 (61.8)      |           |         |
| Education (year), mean (SD)        | 9.5 (1.9) | 10.6 (2.4) | 10.5 (3.1)     | t=0.022   | 0.982   |
| Height (cm), mean (SD)             | 170.0 (9.8) | 168.9 (7.9) | 165.8 (7.2)     | t=1.517   | 0.133   |
| Weight (kg) (mean (SD))            | 61.3 (14.6) | 58.5 (14.3) | 56.9 (9.3)     | t=0.553   | 0.582   |
| BMI (kg/m²)                        | 21.1 (3.9) | 20.4 (4.2) | 20.7 (3.0)     | t=0.361   | 0.719   |
| If antipsychotic taken (n %)       | –    | 13 (81.3)  | 42 (61.8)      | χ²=2.176  | 0.140   |
| Dose                               | –    | 6.4 (2.7)  | 7.8 (4.6)      | t=1.101   | 0.276   |
| SIPS variables                     |      |            |                |           |         |
| SIPS_P (median, mean (SD))         | 10.10 (3.1) | 11.11 (3.6) | 10.10 (3.6)     | Z=1.040   | 0.302   |
| SIPS_N (median, mean (SD))         | 12,12.4 (6.5) | 13,13.7 (4.9) | 11,11.6 (5.6)     | Z=1.356   | 0.179   |
| SIPS_D (median, mean (SD))         | 6,6.4 (3.0) | 7,7.3 (2.4) | 7,6.4 (2.8)     | Z=1.267   | 0.209   |
| SIPS_G (median, mean (SD))         | 9,9.2 (2.9) | 10,10.2 (2.0) | 10,9.5 (2.5)     | Z=1.011   | 0.302   |
| SIPS_TOTAL (median, mean (SD))     | 40,39.2 (12.5) | 42,42.3 (6.8) | 35,36.2 (9.7)     | Z=2.333   | 0.023   |

Dose of antipsychotics was measured using an olanzapine-equivalent dose of antipsychotics. BMI, body mass index; GAF, Global Assessment of Functioning; GAF drop, GAF baseline score that drops from highest in the past year; SD, standard deviation; SIPS, the Structured Interview for Prodromal Syndromes; SIPS_D, scores of disorganisation symptoms scale in SIPS; SIPS_G, scores of general symptoms scale in SIPS; SIPS_N, scores of negative symptoms scale in SIPS; SIPS_P, scores of positive symptoms scale in SIPS; SIPS_TOTAL, total scores of SIPS; t/Z/χ², t for independent t test, Z for Mann-Whitney U test (non-parametrical test), χ² for χ test.
converters was small, the distinctiveness of their baseline niacin-induced responses was established, which is highly valuable in clinical practice for predicting psychosis since only 20% of CHR individuals would convert to psychosis in the near future. The results of the differences between converters and non-converters in the flush scores in niacin skin were presented. P values were calculated by the MANOVA test. AMN, aqueous methylnicotinate; CHR, clinical high risk; MANOVA, multivariate analysis of variance.

**Implications**

The current study provides evidence that blunted flush responses to niacin appear in the early stage of psychosis and are associated with the disease progression. This could represent a psychosis subgroup with a similar underlying pathophysiology. The association of membrane polyunsaturated fatty acids (PUFAs) with psychosis, and with niacin sensitivity (especially in AA PUFAs) has been shown previously. Together with our findings, we hypothesise that there is indeed a subgroup of early psychosis with blunted flush response phenotypes and an underlying pathophysiology of abnormal fatty acid spectrum in neural cell membranes that hampers neuron activity and signal transduction. Therefore, these patients may benefit from the implementation of more specific strategies (eg, supplementation of eicosapentaenoic acid) for early prevention or treatment.

**CONCLUSION**

In summary, SHARP-extended CHR individuals showed attenuated niacin-induced flush responses characterised and unreliable; thus, this method is far from being quantifiable and useful for accurate clinical applications in early psychosis. The laser Doppler flowmeter for the niacin test was introduced by Messamore in 2003; it measures microcirculation in a non-invasive way, which is a useful method to quantify the extent of skin flush responses. The cutaneous blood flow response to increasing concentrations of topical methylnicotinate was measured to derive the log_{10}(EC_{50}) and maximal blood flow values from the dose-response curve of each subject. Another method, denoted optical reflection spectroscopy, introduced by Smesny et al, enables the objective assessment of colour changes and can be used to quantify skin responses. Recently, by applying artificial intelligence technologies for image recognition, our team developed a niacin test device with easy access and high credibility for quantifying skin responses. Details of this method can be found in a companion article by Chen et al.

**Limitations**

This study has several limitations. First, clinical symptoms were not assessed in the HC subjects, whereas the PANSS was applied to the CHR group. Therefore, whether relevant variables from these measures could be confounders when analysing niacin responses remains unknown. Second, the niacin tests were performed only once. The evidence would have been strengthened if we had tested them multiple times. Third, the sample size of the converter group was small; therefore, the present findings from comparisons between the converter and non-converter groups are insufficient and require further study. Finally, the SHARP-extended CHR cohort was surveyed naturally. There were 61 CHR individuals treated with antipsychotics for at least 2 weeks during the follow-up period. Therefore, it remains unknown whether antipsychotics can affect conversion outcomes.
by a modest level of severity that was intermediate between those of HCs and patients with FES. The niacin flush response is more blunted in CHR converters than in non-converters. Future research should develop and validate new devices for the objective quantification of skin responses. Previous reviews have shown that an attenuated niacin response is associated with abnormalities in membrane fatty acid composition and activation of neuroinflammatory processes.\textsuperscript{t3}2 39 There are potential implications for the rational, early and precise identification and intervention for psychosis.

**Author affiliations**
1 Shanghai Intelligent Psychological Evaluation and Intervention Engineering Technology Research Center, Shanghai Key Laboratory of Psychotic Disorders, Shanghai Mental Health Center, Shanghai Jiao Tong University School of Medicine, Shanghai, China
2 Department of Automation, Shanghai Jiao Tong University, Shanghai, China
3 Big Data Research Lab, University of Waterloo, Waterloo, Ontario, Canada
4 Center for Excellence in Brain Science and Intelligence Technology (CEBSIT), Chinese Academy of Science, Shanghai, China
5 Brain Science and Technology Research Center, Shanghai Jiao Tong University, Shanghai, China

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**Data availability statement** Data are available upon reasonable request. The data that support the findings of this study are available from the corresponding author upon reasonable request.

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**ORCID iDs**
Ranpiao Gan http://orcid.org/0000-0002-6920-1447
Yanyan Wei http://orcid.org/0000-0002-6218-1954
Lihua Xu http://orcid.org/0000-0002-2237-9336
Yao Chen http://orcid.org/0000-0003-2987-6711
Tianhong Zhang http://orcid.org/0000-0002-5379-7119

**REFERENCES**
1 Jablensky A. Subtyping schizophrenia: implications for genetic research. *Mol Psychiatry* 2006;11:815–36.
2 Taylor JH, Calkins ME, Gur RE. Markers of psychosis risk in the general population. *Biol Psychiatry* 2020;88:337–48.
3 Messamore E. The niacin response biomarker as a schizophrenia endophenotype: a status update. *Prostaglandins Leukot Essent Fatty Acids* 2018;136:95–7.
4 Gottseman M. The endophenotype concept in psychiatry: etymology and strategic intentions. *Am J Psychiatry* 2003;160:636–45.
5 Messamore E. Relationship between the niacin skin flush response and essential fatty acids in schizophrenia. *Prostaglandins Leukot Essent Fatty Acids* 2003;69:413–9.
6 Yao JK, Dougherty GG, Gautier CH, et al. Prevalence and specificity of the abnormal niacin response: a potential endophenotype marker in schizophrenia. *Schizophrenia Bull* 2016;42:369–76.
7 Sun L, Yang X, Jiang J, et al. Identification of the niacin-blunted subgroup of schizophrenia patients from mood disorders and healthy individuals in Chinese population. *Schizophrenia Bull* 2018;44:896–907.
8 Lien Y-J, Huang S-S, Liu C-M, et al. A genome-wide quantitative linkage scan of niacin skin flush response in families with schizophrenia. *Schizophrenia Bull* 2013;39:68–76.
9 Chang S-S, Liu C-M, Lin S-H, et al. Impaired flush response to niacin skin patch among schizophrenia patients and their nonpsychotic relatives: the effect of genetic loading. *Schizophrenia Bull* 2009;35:213–21.
10 Hu Y, Xu L, Gan R, et al. A potential objective marker in first-episode schizophrenia based on abnormal niacin response. *Schizophrenia Res* 2021;101(6):s084. [Epub ahead of print: 26 Jun 2021].
11 Waldmo MC. Co-distribution of sensory gating and impaired niacin flush response in the parents of schizophrenics. *Schizophrenia Res* 1999;40:49–53.
12 Tavares H, Yacobian J, Talib LL, et al. Increased phospholipase A2 activity in schizophrenia with absent response to niacin. *Schizophrenia Res* 2003;61:1–6.
13 Shen H, Chen M, Cui D. Biological mechanism study of medication and its application in mental disorders. *Gen Psychiatr* 2020;33:e100214.
14 Messamore E, Hoffman WF, Yao JK. Niacin sensitivity and the arachidonic acid pathway in schizophrenia. *Schizophrenia Res* 2010;122:248–56.
15 Tang Y, Zhou L, Gunnet JW, et al. Enhancement of arachidonic acid signaling pathway by nicotinic acid receptor HM74A. *Biochem Biophys Res Commun* 2006;345:29–37.
16 Murakami M, Kudo I. Recent advances in molecular biology and physiology of the prostaglandin E2-biosynthetic pathway. *Prog Lipid Res* 2004;43:3–35.
17 Ansarey SH. Inflammation and JNK’s role in niacin-GRPR109A diminished flushed effect in microglial and neuronal cells with relevance to schizophrenia. *Front Psychiatry* 2021;12:771144.
18 Zhang T, Li H, Woodberry KA, et al. Prodromal psychosis diagnosis in a counseling center population in China: an epidemiological and clinical study. *Schizophrenia Res* 2014;152:391–9.
19 Zhang T, Xu L, Tang Y, et al. Prediction of psychosis in prodrome: development and validation of a simple, personalized risk calculator. *Psychol Med* 2019;49:1990–8.
20 Zhang TH, Li JH, Woodberry KA, et al. Two-year follow-up of a Chinese sample at clinical high risk for psychosis: timeline of symptoms, help-seeking and conversion. *Epidemiol Psychiatr Sci* 2017;26:287–98.
21 Nadalin S, Jonovska S, Šendula Jengi V, et al. An association between niacin skin flush response and plasma triglyceride levels in individuals with schizophrenia. *Prostaglandins Leukot Essent Fatty Acids* 2020;155:102084.
22 Langbein K, Schmidt U, Schack S, et al. State marker properties of niacin skin sensitivity in ultra-high risk groups for psychosis.
Ranpiao Gan is a medicine postgraduate at Shanghai Jiao Tong University School of Medicine. She graduated from Chongqing Medical University in 2018 with a bachelor's degree in medicine. Since 2019, she has been engaged in the assessment of clinical high-risk for psychosis and the detection of niacin at the Shanghai Mental Health Center. Her main interest is the early identification and intervention of clinical high-risk for psychosis.

- an optical reflection spectroscopy study. Schizophr Res 2018;192:377–84.
23 Berger GE, Smesny S, Schäfer MR, et al. Niacin skin sensitivity is increased in adolescents at ultra-high risk for psychosis. PLoS One 2016;11:e0148429.
24 Barako T, Li C, Yeung A, Massachusetts General Hospital and the Shanghai Mental Health Center: the past, present and future of a psychiatric research partnership. Gen Psychiatr 2019;32:e100157.
25 Zhang T, Li H, Tang Y, et al. Validating the predictive accuracy of the NAPLS-2 psychosis risk calculator in a clinical high-risk sample from the SHARP (Shanghai At Risk for Psychosis) program. Am J Psychiatry 2018;175:906–8.
26 Zhang T, Xu L, Li H, et al. Calculating individualized risk components using a mobile app-based risk calculator for clinical high risk of psychosis: findings from Shanghai At Risk for Psychosis (SHARP) program. Psychiatr Med 2021;51:653–60.
27 Zhang T, Xu L, Tang X, et al. Real-world effectiveness of antipsychotic treatment in psychosis prevention in a 3-year cohort of 517 individuals at clinical high risk from the SHARP (Shanghai At Risk for Psychosis). Aust N Z J Psychiatry 2020;54:696–706.
28 Miller TJ, McGlashan TH, Rosen JL, et al. Prodromal assessment with the structured interview for prodromal syndromes and the scale of prodromal symptoms: predictive validity, interrater reliability, and training to reliability. Schizophr Bull 2003;29:703–15.
29 Miller TJ, McGlashan TH, Rosen JL, et al. Prospective diagnosis of the initial prodrome for schizophrenia based on the structured interview for prodromal syndromes: preliminary evidence of interrater reliability and predictive validity. Am J Psychiatry 2002;159:863–5.
30 Miller TJ, Cicchetti D, Markovich PJ, et al. The SIPs screen: a brief self-report screen to detect the schizophrenia prodrome. Schizophr Res 2004;70:78.
31 Ward PE, Sutherland J, Glen EM, et al. Niacin skin flush in schizophrenia: a preliminary report. Schizophr Res 1998;29:269–74.
32 Nadalin S, Buretić-Tomljanović A, Rubesa G, et al. Niacin skin flush test: a research tool for studying schizophrenia. Psychiatr Danub 2010;22:14–27.
33 Smesny S, Berger G, Rosburg T, et al. Potential use of the topical niacin skin test in early psychosis—a combined approach using optical reflection spectroscopy and a descriptive rating scale. J Psychiatr Res 2003;37:237–47.
34 Sethom MM, Fares S, Bouaziz N, et al. Polyunsaturated fatty acids deficits are associated with psychotic state and negative symptoms in patients with schizophrenia. Prostaglandins Leukot Essent Fatty Acids 2010;83:131–6.
35 Laugharne JD, Mellor JE, Peet M. Fatty acids and schizophrenia. Lipids 1996;31 Suppl:S163–5.
36 Glen AI, Cooper SJ, Rybakowski J, et al. Membrane fatty acids, niacin flushing and clinical parameters. Prostaglandins Leukot Essent Fatty Acids 1996;55:9–15.
37 Emsley R, Myburgh C, Oosthuizen P, et al. Randomized, placebo-controlled study of ethyl-eicosapentaenoic acid as supplemental treatment in schizophrenia. Am J Psychiatry 2002;159:1596–8.
38 Jamilian H, Solhi H, Jamilian M. Randomized, placebo-controlled clinical trial of omega-3 as supplemental treatment in schizophrenia. Glob J Health Sci 2014;6:103–8.
39 Horrobin DF. The membrane phospholipid hypothesis as a biochemical basis for the neurodevelopmental concept of schizophrenia. Schizophr Res 1998;30:193–208.