The Effect of Neurogranin on Neural Correlates of Episodic Memory Encoding and Retrieval

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Neurogranin (NRGN) is the main postsynaptic protein regulating the availability of calmodulin-Ca(2+) in neurons. NRGN is expressed exclusively in the brain, particularly in dendritic spines and has been implicated in spatial learning and hippocampal plasticity. Genetic variation in rs12807809 in the NRGN gene has recently been confirmed to be associated with schizophrenia in a meta-analysis of genome-wide association studies: the T-allele was found to be genome-wide significantly associated with schizophrenia. Cognitive tests and personality questionnaires were administered in a large sample of healthy subjects. Brain activation was measured with functional magnetic resonance imaging (fMRI) during an episodic memory encoding and retrieval task in a subsample. All subjects were genotyped for NRGN rs12807809. There was no effect of genotype on personality or cognitive measures in the large sample. Haplozygote carriers of the T-allele showed better performance in the retrieval task during fMRI. After controlling for memory performance, differential brain activation was evident in the anterior cingulate cortex for the encoding and posterior cingulate regions during retrieval. We could demonstrate that rs12807809 of NRGN is associated with differential neural functioning in the anterior and posterior cingulate. These areas are involved in episodesic memory processes and have been implicated in the pathophysiology of schizophrenia in structural and functional imaging as well as postmortem studies.

Key words: NRGN/fMRI/memory/cingulate

Introduction

Neurogranin (NRGN) is the main postsynaptic protein regulating the availability of calmodulin-Ca(2+) in neurons, by binding to calmodulin in the absence of calcium. NRGN is expressed exclusively in the brain. It is abundantly expressed in brain regions involved in cognitive functioning and especially enriched in CA1 pyramidal neurons in the hippocampus and has been shown to play a role in long-term potentiation, spatial learning, and hippocampal plasticity. Furthermore, the NRGN gene has been implicated in schizophrenia and the T-allele of the single nucleotide polymorphism (SNP) rs12807809 (C/T) located upstream of the NRGN has recently been shown to be genome-wide significantly associated with schizophrenia in a meta-analysis of genome-wide association studies.

Although the exact etiology of schizophrenia still remains uncertain, abnormalities in brain structure and function along with a strong genetic component have consistently been implicated in the disorder. Furthermore, several cognitive domains are impaired, among which episodic memory appears to be one of the most severely affected. Recent functional magnetic resonance imaging (fMRI) studies on healthy participants document that variation in susceptibility genes for schizophrenia, among others NRG1, G72, DYNB1, and ZNF804A modulate the neural activation patterns associated with cognitive processing.

Among the cognitive domains impaired in schizophrenia, episodic memory deficits show high effect sizes (d = 0.74 in Heinrichs and Zakzanis), while other domains such as executive functions, working memory, and verbal fluency also show marked levels of impairment. Episodic memory processing has been linked to the hippocampus, cingulate, and frontal and temporal cortical regions.

Besides memory impairment in patients, relatives, and subjects at high risk, many studies have shown structural alterations in patients with schizophrenia in the medial temporal cortex/hippocampal formation. These alterations can also be found in subjects with an at risk
mental state. Further functional imaging has demonstrated dysactivation during episodic memory encoding (eg. ref. 21–23) and retrieval26 in the hippocampal cortex in patients. These alterations of medial temporal structures have also been demonstrated in relatives27–29 and high-risk subjects,30,31 which underlines the importance of these structures in the aetiology of schizophrenia. The cingulate cortex (especially the anterior part) has been shown to be hyperactivated in patients with schizophrenia compared with controls during both encoding and retrieval tasks.32–34 In patients, the anterior commissural line (AC) volume is decreased35 and the posterior cingulate cortex is smaller in both patients and their healthy relatives compared with controls.36

In animal models, NRGN has been shown to exert a profound influence on memory formation: It could be demonstrated that NRGN is involved in long-term potentiation memory formation and it enhances synaptic strength in the hippocampus.3,37–40 In addition, NRGN null mice exhibit anxiety-related behavior.41 During development in the rat brain, the hippocampus and the AC are among the first structures to express NRGN.42

Genetic risk variants for schizophrenia with high allele frequencies and limited effects are also present in a large proportion of the healthy population. It has been shown that investigating the impact of a risk variant in healthy individuals on objectively measurable phenotypes such as performance in fMRI-based neuropsychological paradigms constitutes a successful approach.5 Further functional imaging has been applied successfully in different samples previously.13 Investigating individuals on objectively measurable phenotypes such as that investigating the impact of a risk variant in healthy proportion of the healthy population. It has been shown healthy relatives compared with controls.36

In patients, the anterior commissural line (AC) volume is decreased35 and the posterior cingulate cortex is smaller in both patients and their healthy relatives compared with controls.36

Cognitive Tests and Personality Questionnaires

The following tests were administered in all subjects: A brief verbal IQ assessment,44 the d2 test for attention,45 the letter-number span,46 spatial span,47 the TMT-B,48 and semantic verbal fluency.49 In addition, all subjects completed the NEO-FFI50 and the brief version of the schizotypal personality questionnaire (SPQ-B)51 with the scales cognitive perceptual deficits, interpersonal deficits and disorganization.

Genotyping

Genomic DNA was extracted from ethylenediaminetetraacetic acid anticoagulated venous blood according to standard procedures.52 The SNP rs12807809 was genotyped on an Applied Biosystems 7900HT Fast Real-Time PCR System, using a TaqMan 5’ nuclease assay (TaqMan SNP Genotyping Assay ID C_32029000_20 Applied Biosystems). Genotyping accuracy was assessed by running 15% of the sample in duplicates. Reproducibility was 100%.

Encoding and Retrieval fMRI Paradigm

The paradigm consisted of an encoding and a retrieval task performed in different sessions. Both sessions were divided by a break of approximately 3 minutes. During this time, subjects stayed in the scanner.

Encoding Task. During the encoding phase, either single pictures of neutral faces (encoding condition) or the symbol “#” (baseline condition) were presented on a black background for 4000 ms in a pseudorandomized order using Presentation software package (Neurobehavioral Systems Inc, San Francisco, CA). Following this, stimuli were replaced by a blank screen for another 1000 ms completing one trial. During face encoding, participants were instructed to actively memorize each face for later recognition. In order to ensure continuous
attention to the task, participants had to indicate the sex of the displayed person via button press (LUMItouchTM; Lightwave Technologies, Richmond, BC, Canada). During low-level baseline, participants were enforced to press a button with the left index finger every time the symbol # appeared. There were 5 blocks of each condition with 6 responses in each block resulting in 30 faces (half male and half female) to be encoded. Similarly, during baseline, 30 button presses had to be accomplished. Each block lasted for 30 seconds. This task has been applied successfully in different samples previously by our group.25,53–55

Retrieval Session. After the encoding phase, a recognition phase of equal length and structure was administered. During the retrieval condition, 2 pictures of faces were presented simultaneously side by side, each trial comprising a previously presented face and a new face, randomly positioned at the left or right side. Subjects were requested to select the previously presented face and forced to make a choice by pressing the corresponding button with the left or right index finger. The baseline condition was the same as in the encoding phase.

**MRI Data Acquisition**

All MRI data were acquired on a 3-Tesla Tim Trio MR scanner (Siemens Medical Systems) at the Research Center Jülich. Functional images were collected with a T2* weighted echo planar imaging sequence sensitive to BOLD contrast (64 × 64 matrix, FOV 200 mm, in plane resolution 3.13 mm, 36 slices, slice thickness 3 mm, TR = 2.25 s, TE = 30 ms, flip angle 90°). Slices covered the whole brain and were positioned transaxially parallel to the anterior-posterior commissural line. One hundred and thirty-seven functional images were collected, and the initial 3 images excluded from further analysis in order to remove the influence of T1 stabilization effects.

**FMRI Data Analyses**

**Analysis of Behavioral Data.** Behavioral data (i.e., the number of correctly remembered faces in the retrieval session)
were analyzed using an independent t test with rs12807809 status (T/T vs C/T and C/C genotype) as grouping variable.

Analysis of fMRI Data. SPM5 (www.fil.ion.ucl.ac.uk/spm) standard routines and templates were used for analysis of fMRI data. The functional images were realigned, normalized (resulting voxel size $2 \times 2 \times 2$ mm$^3$), smoothed (8 mm isotropic Gaussian filter), and high-pass filtered (cut off period 120 s).

Statistical analysis was performed in a 2-level mixed-effects procedure. At the first level, the BOLD responses for the activation (encoding and retrieval, respectively) and the baseline condition were modeled by a boxcar function convolved with the canonical hemodynamic response function employed by SPM5. Parameter estimates ($\beta$-) and t statistic images were calculated for each subject. At the second level, the individual $\beta$-contrasts relating to activation differences between the activation and the baseline condition were entered into a t test design with rs12807809 status (T/T vs T/C and C/C status) as grouping variable. First, we calculated group activation maps related to the activation of memory encoding/retrieval (ie, encoding>baseline; retrieval>baseline). Activation maps were thresholded at $P < .05$, corrected for multiple comparisons (applying the family-wise error correction employed by SPM5). Second, we determined activation differences between the 2 rs12807809 genotype groups. Resulting first-level contrasts were entered in a second level, 2-sample t test. As groups differed with regard to gender distribution and memory performance (see results section and table 1), these 2 parameters were entered as covariates of no interest into the 2-sample t test.

In order to correct for multiple comparisons within a search volume, we applied a cluster extent threshold determined by Monte Carlo simulations. For a threshold at the voxel level at $P = .001$ and spatial properties as present in this study, 10,000 simulations resulted in an extent threshold of 26 resampled voxels. This procedure prevented a false positive rate above 5% due to multiple testing. The anatomical localization of activated brain regions was assessed both by the SPM anatomy toolbox and the Talairach atlas. Based on effect sizes from previous findings (eg, Kircher et al), a statistical power of 1$-\beta = .80$ resulted with the sample size in the fMRI task of the faces.

Cluster of At Least 26 Voxels (See Method Section) are Depicted

<table>
<thead>
<tr>
<th>Coordinates</th>
<th>Value Cluster Size (in voxels)</th>
<th>t</th>
</tr>
</thead>
<tbody>
<tr>
<td>L Anterior Cingulate Cortex 24</td>
<td>22 24 3.99 52</td>
<td>4.74</td>
</tr>
<tr>
<td>R Lingual Gyrus 18 24</td>
<td>26 64 4.22 46</td>
<td>4.28</td>
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<tr>
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<td>14 2 4.28 88</td>
<td>4.28</td>
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<td></td>
<td>14 2 4.28 88</td>
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</tbody>
</table>

Results

Cognitive Functioning and Personality Measures

Homzygous carriers of the T-allele did not differ from C-allele carriers in either of the administered tests or questionnaires with the sole exception of semantic verbal fluency. Homzygous T-allele carriers produced more words compared with the group of C-allele carriers ($31.8 \pm 9.2$ and $30.1 \pm 9.1$, respectively, $P = .046$). This difference did not withstand correction for multiple comparisons.

Genotype did not correlate with IQ, age, or years of education (all $P > .05$). During the fMRI task, there was an effect of NRGN genotype on the number of correctly recognized faces. Homzygous T-allele carriers showed a significantly better performance during retrieval ($P = .027$). These differences, along with a difference in gender distribution, were accounted for in the statistical analysis section. All results are given in table 1.

fMRI Results

During encoding, homzygous T-allele carriers exhibited stronger activations in the left lingual gyrus (Brodmann areas [BA] 19) and the anterior cingulate cortex (ACC, BA 24) compared with subjects with at least one C-allele ($P < .001$). There were no significant differences in the gender identification task: homzygous T-allele carriers correctly identified 98.14% (SD $= 2.3$) of the faces and C-allele carriers correctly identified 98.80% (SD $= 1.2$) of the faces.

The reversed contrast (T/C + C/C>T/T) did not yield any significant activations. Results are depicted in figure 1 and table 2.

During retrieval, homzygous T-allele carriers showed less deactivations compared to C-allele carriers in the left precentral gyrus, right cingulate gyrus (expanding to the right precentral gyrus), and the left insula ($P < .001$; see figure 2). Homzygous T-allele carriers showed better performance (correctly recognized faces) compared with the other group ($P = .027$; this was taken into account as a covariate of no interest, see above).

For both tasks, ROI analyses neither revealed significant activations within the hippocampus proper nor in the parahippocampal gyrus.

Discussion

This is the first report to describe the influence of a recently discovered genome-wide significant schizophrenia variant in NRGN on cognition and personality traits as well as neural correlates of episodic memory encoding and retrieval that was investigated in a large sample of healthy subjects. While no differences in personality dimensions were found in our study, homzygous T-allele carriers showed a trend toward higher performance in a semantic verbal fluency task. During fMRI scanning, homzygous T-allele carriers showed better performance with respect to correctly recognized faces. When controlling for performance and gender distribution, this group
C-allele carriers in either of the administered tests or homozygous carriers of the T-allele did not differ from healthy subjects. While no differences in personality dimensions were found in our study, homozygous T-allele carriers showed higher activations in the ACC and the lingual gyrus during encoding and less deactivation in the left insula, left precentral gyrus, and the cingulate gyrus during retrieval. It could previously be demonstrated that NRGN...

### Results

**Memory encoding task**

- **T/T vs T/C + C/C**
  - R Lingual Gyrus
    - BA: 18
    - Coordinates: $(x, y, z) = (24, 24, -94)$
    - $t$ Value: 4.28
    - Cluster Size: 88
  - L Anterior Cingulate Cortex
    - BA: 24
    - Coordinates: $(x, y, z) = (-4, 30, 12)$
    - $t$ Value: 3.85
    - Cluster Size: 35

- **C/C + C/T vs T/T**
  - No areas of differential activation

**Memory retrieval task**

- **T/T vs T/C + C/C**
  - L Precentral Gyrus
    - BA: 4
    - Coordinates: $(x, y, z) = (-14, -26, 64)$
    - $t$ Value: 4.22
    - Cluster Size: 46
  - R Cingulate Gyrus
    - BA: 24
    - Coordinates: $(x, y, z) = (20, -20, 46)$
    - $t$ Value: 4.14
    - Cluster Size: 161
  - L Insula
    - BA: 13
    - Coordinates: $(x, y, z) = (-34, -22, 24)$
    - $t$ Value: 3.99
    - Cluster Size: 52

- **C/C + C/T vs T/T**
  - No areas of differential activation

### Table 2

**Correlations of rs12807809 of NRGN Status With Neural Activations During Memory Encoding and Retrieval Processes; Only Clusters of At Least 26 Voxels (See Method Section) Are Depicted**

<table>
<thead>
<tr>
<th>Hemisphere</th>
<th>BA</th>
<th>Coordinates</th>
<th>$t$ Value</th>
<th>Cluster Size (in voxels)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Memory encoding task</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(T/T $&gt;$ T/C + C/C)</td>
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<td>$(x, y, z) = (24, 24, -94)$</td>
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<td>(C/C + C/T $&gt;$ T/T)</td>
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<tr>
<td>No areas of differential activation</td>
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<td></td>
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<tr>
<td>Memory retrieval task</td>
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<tr>
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<tr>
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**Note:** Coordinates are listed in MNI atlas space. BA is the Brodmann area nearest to the coordinate and should be considered approximate. NRGN, Neurogranin.
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is differentially expressed in the prefrontal cortex in patients with schizophrenia. A modulation of its function through genetic variation may therefore be a plausible explanation of our findings at the functional level.

Cognitive Functions and Personality
In the large sample, no differences in cognitive performance or any measures of personality could be detected after correction for multiple testing. This is in line with recent evidence from a study investigating a different German sample. However, there was a trend in differences in semantic verbal fluency. Importantly, it has to be noted that homozygous T-allele carriers showed a better performance than C-allele carriers. This result is somewhat surprising with regard to findings in schizophrenia where patients show impairments in semantic verbal fluency. However, there are several studies that also show these resulting patterns: in 2 studies investigating cognitive domains in healthy subjects, similar results have been demonstrated with respect to G72 and G73. In these studies, subjects with the risk diplotype showed a higher performance in working memory and attention than subjects without the risk diplotype. In a related fashion, Stefanis et al. found that healthy carriers of the minor allele in Dysbindin rs1018381 scored lower on the paranoid factor of the SPQ. Since these studies demonstrated that risk allele carriers also showed better results in a variety of variables, the present study may point to possible further investigations into semantic processing associated with NRG1 genotype variations.

fMRI Results
Encoding. The ACC has consistently been implied in control and decision-making processes (eg, Kennerley et al. as well as episodic memory (eg, Svoboda
et al.\(^4\) Particularly, the BOLD signal increase in this region has been shown to correlate with subsequent memory performance,\(^5\,6\) demonstrating its importance in encoding strategies. In schizophrenia, meta-analytical findings demonstrated higher ACC activations in patients as compared with healthy controls during episodic memory encoding.\(^3\) A similar activation pattern has also been reported for executive processing comparing patients with schizophrenia to healthy controls.\(^6\,6\) Meta-analyses additionally confirmed a volume reduction in the ACC in patients compared with controls (see also Introduction section).\(^6\,7\) Taken together, these findings implicate the ACC as an important structure in memory encoding as well as the pathophysiology of schizophrenia. Our T/T risk allele carriers exhibited stronger activation in the ACC during encoding new material. This may indicate that they must exert higher cognitive control in order to reach a similar performance level as the nonrisk carrier group.

The lingual gyrus (BA 19) is being activated when subjects process features of faces rather than the configuration of a face as a whole.\(^6\,8\) It could thus be argued that homozygous T-allele carriers may focus more on particular single facial components while encoding novel faces. This is a strategy that has been attributed to patients with schizophrenia.\(^6\,9,7\,0\)

**Retrieval.** The cingulate gyrus, especially the posterior part, has been implied as a part of a “core” network subserving episodic memory,\(^6\,4\) especially recognition of previously encoded material.\(^7\,1\) As such, it could be hypothesized that homozygous T-allele carriers rely more on this region in order to adequately perform memory processes.

Activations of the left insula in patients with schizophrenia during episodic memory processes are a common finding in fMRI studies (meta-analysis eg, Ragland et al.\(^3\)\(^3\)). In addition, left insular volume is decreased in schizophrenia compared with healthy subjects. These changes are already obvious in first-episode schizophrenia (meta-analysis in Ellison-Wright et al.\(^7\)\(^2\)).

Other studies also found higher activations in the left precentral gyrus in patients compared with controls during memory retrieval (eg, ref. \(^3\)\(^3,7\)\(^2,7\)\(^4\)). The differences in activation occurred while controlling for performance in the statistical analyses. It could be argued that homozygous T-allele carriers show less deactivation in the reported regions and thus compensating behavioral underperformance. Overrecruitment of task-related brain regions is a phenomenon found in the ageing brain\(^7\)\(^2\) and schizophrenia\(^7\)\(^6\) but has also been shown to be present in risk allele carriers of NRG1 in healthy subjects.\(^1\)\(^0\) Following this logic, one could hypothesize that fine tuning of cortical activations during memory performance is disturbed in homozygous T-allele carriers.

In contrast to our hypotheses, we could not detect an influence of NRG1 genotype on hippocampal activation in our ROI analyses, neither for encoding nor retrieval. The encoding paradigm was developed to particularly activate the hippocampus and we could previously detect activation differences in these areas between healthy subjects and patients with schizophrenia\(^5\)\(^5\) and Alzheimer’s Disease.\(^5\)\(^5\) It is unlikely that we missed differences in the current study. It seems that this common variant— even though associated with memory functions as evidenced by our fMRI task—exerts an influence on a widespread neural network that is implied in memory formation, such as the ACC and the insula, but not the hippocampal formation. Variations in other genes associated with schizophrenia have been shown to impact on hippocampal structure and functioning during memory processes.\(^7\)\(^7\) As such, it is suggested that some, but not all, risk polymorphisms involved in schizophrenia may impact on hippocampal formation processes. Consequently, different risk variants act on different neural networks, their prominent locus of expression depending on their relative function during brain development. Among other factors, this might explain the heterogeneous symptomatology and course of the disorder.

In sum, when correcting for multiple comparisons, the present findings demonstrate that rs12807809 does not seem to influence domains of cognition or personality dimensions in this sample of healthy subjects, which replicates the results of a recent study.\(^6\,6\) However, there was a trend toward differences in semantic verbal fluency performance and in episodic memory functions, which warrant further investigation. The main finding was that rs12807809 exerts a profound influence on the cortical activation/deactivation pattern evident during episodic memory processes. In this study, we could show that variation in rs12807809 does not necessarily influence memory processes orchestrated by the hippocampus proper but rather impacts on cortical regions that have been implicated in this function. Thus these findings suggest that rs12807809 has a pleiotropic effect on different brain networks. As this is the first study on NRG1 rs12807809 replications of our findings are warranted. Even though we studied a large sample of healthy subjects, it still remains the possibility that these are chance findings.

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