Learning and Generalization in Schizophrenia: Effects of Disease and Antipsychotic Drug Treatment

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Background: Schizophrenia involves alterations in hippocampal function. The implications of these alterations for memory function in the illness remain poorly understood. Furthermore, it remains unknown how memory is impacted by drug treatments for schizophrenia. The goal of this study was to delineate specific memory processes that are disrupted in schizophrenia and explore how they are affected by medication. We specifically focus on memory generalization—the ability to flexibly generalize memories in novel situations.

Methods: Individuals with schizophrenia (n = 56) and healthy control subjects (n = 20) were tested on a computerized memory generalization paradigm. Participants first engaged in trial-by-error associative learning. They were then asked to generalize what they learned by responding to novel stimulus combinations. Individuals with schizophrenia were tested on or off antipsychotic medication, using a between-subject design in order to eliminate concerns about learning-set effects.

Results: Individuals with schizophrenia were selectively impaired in their ability to generalize knowledge, despite having intact learning and memory accuracy. This impairment was found only in individuals tested off medication. Individuals tested on medication generalized almost as well as healthy control subjects. This between-group difference was selective to memory generalization.

Conclusions: These findings suggest that individuals with schizophrenia have a selective alteration in the ability to flexibly generalize past experience toward novel learning environments. This alteration is unaccompanied by global memory impairments. Additionally, the results indicate a robust generalization difference on the basis of medication status. These results suggest that hippocampal abnormalities in schizophrenia might be alleviated with antipsychotic medication, with important implications for understanding adaptive memory-guided behavior.

Key Words: Antipsychotic drugs, hippocampus, human, memory, schizophrenia

Considerable evidence suggests that individuals with schizophrenia display subtle but consistent memory impairments (1,2). However, individuals with schizophrenia are not impaired on all forms of memory (1,3). Furthermore, the effect of treatment on memory impairments in schizophrenia remains poorly understood at both the cognitive and neurobiological levels. Thus, a central challenge is to characterize the specificity of the memory impairments in schizophrenia, their relation to specific neural systems, and their modulation by antipsychotic drugs (APDs).

One aspect of memory function that might be particularly vulnerable in schizophrenia is the ability to flexibly generalize memories from past events when encountering novel situations in the future. Understanding how memories might guide behavior in novel situations provides insight into a fundamental aspect of adaptive human behavior. Emerging evidence suggests that, in healthy individuals, generalization of memories depends critically on the hippocampus (4–6), a brain region widely known to support the formation of accurate declarative memories (7–9). Here, we examine the hypothesis that individuals with schizophrenia are specifically impaired at memory-based generalization, and we explore how generalization is impacted by treatment with APDs.

There are a number of reasons to expect that memory generalization might be impaired in schizophrenia and might be impacted by APDs. Extensive evidence suggests that schizophrenia is associated with abnormal hippocampal function. Reductions in hippocampal volume (10,11), increases in hippocampal perfusion (e.g., regional cerebral blood flow) (12,13), and reductions in task-associated activations in the hippocampus (14,15) are found in schizophrenia, along with impaired performance on hippocampal-dependent memory tasks (1,2). A particularly strong link has been found between hippocampal function and memory generalization in schizophrenia with a transitive inference paradigm (1,2). Transitive inference is a form of hippocampal-dependent generalization in which learned information is used to later guide logical inferences. Individuals with schizophrenia (tested on medication) are impaired at transitive inference, and their impairment is related directly to hippocampal dysfunction (16). However, it is not known how memory-based generalization might be impacted by APDs or how these findings relate to other forms of generalization that do not rely on logical inference.

Evidence suggesting that generalization might be impacted by APDs comes from recent reports that generalization might also depend on dopaminergic mechanisms in the midbrain (6). For example, in one study, healthy participants engaged in a novel two-phase learning and generalization task while being scanned with functional magnetic resonance imaging (fMRI). In this paradigm, the first phase involved feedback-based learning of a series of associations. In the second phase, subjects were probed to generalize what they learned in response to novel stimulus combinations. Here, generalization does not depend on logical inference but on associative memory processes that take place during learning (6). The fMRI data revealed that in this context
the ability to generalize involved correlated activation in the hippocampus and in midbrain dopaminergic regions (ventral tegmental area [VTA]) during learning, suggesting a cooperative hippocampal-midbrain interaction that supports generalization. The precise anatomical underpinnings of this interaction remain to be determined. However, VTA dopamine neurons are known to project directly to the hippocampus (17–19), where dopamine modulates hippocampal plasticity (20,21), suggesting a likely mechanism by which midbrain dopaminergic signals might modulate hippocampal representations.

These findings in healthy individuals raise questions regarding the effects of APDs on memory-based generalization. Antipsychotic drugs are known to antagonize dopamine (as well as other monoamine receptors) (22). Thus, one possibility is that APDs might worsen generalization performance. However, other evidence demonstrates that dopamine antagonism generally reduces symptoms in schizophrenia (23,24), raising the possibility that APDs might improve generalization of memories. Indeed, one early report indicated that APDs improve verbal memory performance on a recent memory task. However, because that study did not specifically examine generalization, it remains unknown how or whether APD treatment impacts generalization in schizophrenia. Understanding the effects of APDs on memory and generalization is important from both a clinical and basic science perspective, because such understanding might provide insight into possible mechanisms underlying memory-based generalization in the illness.

The goal of the present study was to address two main open questions. First, we sought to examine the effect of schizophrenia on a memory generalization paradigm that does not depend on logical inference. On the basis of prior reports (25), we predicted that individuals with schizophrenia would be impaired at generalization. Second, we sought to determine the effects of APDs on generalization in schizophrenia to distinguish the effects of disease from the effects of medication on this fundamental cognitive process. To that end, we tested generalization in individuals with schizophrenia who were either taking or not taking their medication. Given that the effect of APDs on hippocampal function is largely unknown, we explored whether APDs would impair, facilitate, or have no effect on generalization performance. Importantly, the generalization paradigm used here allows for separate assessments of: 1) learning, 2) memory retention, and 3) memory-guided generalization. Therefore, to the extent that APDs impact generalization, this paradigm allows us to determine whether APDs specifically affect generalization or whether they have a broad impact on global memory function.

We solicited informed consent, validated potential diagnosis and group placement, and subsequently tested all normal volunteers and most schizophrenic volunteers at the UT SW Research Clinic; volunteers were tested with the behavioral task within 2 weeks of consenting. Additional information regarding patient recruitment, medication status, and exclusion criteria are found in Supplement 1.

All patient volunteers received a thorough diagnostic workup with the Structured Clinical Interview for Diagnosis (SCID) and met DSM-IV criteria for schizophrenia or schizoaffective disorder. Consensus diagnoses were based on all available psychiatric information and made by at least two experienced research psychiatrists. Ratings for psychosis (i.e., Positive and Negative Symptom Scale [PANSS]) were conducted by three trained research coordinators whose inter-rater reliability was $r = .84$, intraclass correlation. Informed consent was obtained for all participants in accordance with procedures approved by the UT SW Institutional Review Board.

Ages of the volunteers ranged from 18 to 59 years. Control volunteers (NV) averaged 39.9 years of age (SD = 11.8 years) and were not different from either SV-ON (mean = 40.4; SD = 9.1; $p = .85$) or SV-OFF (mean = 36.4; SD = 11.9, $p = .40$); nor were there differences in age between the SV-ON and the SV-OFF ($p = .18$). The ages for disease onset for all SV volunteers were between 21 and 27 years of age. The education level of the control group was 14.0 years (SD = 1.8), which was not significantly different from the SV-ON (mean = 13.7; SD = 2.7, $p = .55$). The education level of SV-OFF was 11.9 years (SD = 2.1), which was lower than the level of the NV ($p = .003$) and, to a lesser extent, the SV-ON ($p = .04$). Of the 20 control subjects, 19 were administered the Wechsler Test of Adult Reading (WTAR) in predicting IQ; 36 of the 40 SV-ON and 13 of the 16 SV-OFF were also administered the WTAR for predicting premorbid IQ; this measurement yielded significant differences between NV (mean = 107; SD = 8.4) and both SV-ON (mean = 98; SD = 12.2, $p = .001$) and SV-OFF (mean = 96; SD = 10.9, $p = .001$) but no significant differences in predicted premorbid IQ between the SV-ON and SV-OFF ($p = .65$).

Patient assessments included the PANSS for symptom severity (26), the Repeatable Battery for the Assessment of Neuropsychological Status (RBANS) (27), and the Birchwood Social Functioning Scale (SFS) (28). Demographic and clinical characteristics of the participants are presented in Table 1. Consistent with their medication status, the SV-OFF scored significantly higher than the SV-ON with respect to the total PANSS score ($t(35) = 2.57, p = .014$) and on PANSS-general score ($t(35) = 3.10, p = .004$). The groups did not differ on other measures.

### Methods and Materials

#### Subjects

The effects of APDs on generalization were examined with a between-subject design to avoid effects of order and learning-set which are common in studies of learning and memory. Results are reported from 56 volunteers with schizophrenia and 20 healthy control volunteers at the University of Texas Southwestern (UTSW) Schizophrenia Research Clinic.

Patients were tested either on antipsychotic medication (SV-ON; $n = 20$) or off antipsychotic medication (SV-OFF; $n = 16$). All volunteers were recruited from the Dallas metropolitan area; the healthy control subjects through advertising and the schizophrenic volunteers through advertising and directly at a community mental health clinic.

#### Task Stimuli and Procedures

We used an “acquired equivalence” task, previously shown to be sensitive and specific to hippocampal function (25,29,30). As shown in Figure 1, the task consisted of two phases. Subjects first engage in associative learning (Learning Phase) and then are tested on generalization (Test Phase) (25,29,30). During the Learning phase, participants use trial-by-error feedback to learn associations between stimuli (cartoon faces and colored fish). Although each person–fish association is learned individually, there is partial overlap between them, such that two different people are associated with the same fish ($P_1$–$F_1$, $P_2$–$F_1$). This overlap provides participants with the opportunity to form an associative link between the two people ($P_1$ and $P_2$), even though they have not been experienced together. Importantly, a
Table 1. Demographic and Neuropsychological Measures for All Participants

<table>
<thead>
<tr>
<th></th>
<th>NV (n = 20)</th>
<th>SV-ON (n = 40)</th>
<th>SV-OFF (n = 16)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male/female, n</td>
<td>5/15</td>
<td>26/14</td>
<td>12/4</td>
</tr>
<tr>
<td>Age (yrs)</td>
<td>39.9 (11.8)</td>
<td>40.4 (9.1)</td>
<td>36.4 (11.9)</td>
</tr>
<tr>
<td>Education (yrs)</td>
<td>14.0 (1.84)</td>
<td>13.7 (2.7)</td>
<td>11.9 (2.9)</td>
</tr>
<tr>
<td>Chlorpromazine Equivalent (mg)</td>
<td>—</td>
<td>483.5 (404.4)</td>
<td>—</td>
</tr>
<tr>
<td>PANSS</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>—</td>
<td>81.07 (10.8)</td>
<td>92.43 (8.8)</td>
</tr>
<tr>
<td>Positive</td>
<td>—</td>
<td>21.30 (4.3)</td>
<td>23.3 (5.8)</td>
</tr>
<tr>
<td>Negative</td>
<td>—</td>
<td>19.20 (4.7)</td>
<td>22.3 (4.1)</td>
</tr>
<tr>
<td>General</td>
<td>—</td>
<td>40.63 (5.7)</td>
<td>47.8 (4.8)</td>
</tr>
<tr>
<td>RBANS</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>—</td>
<td>80.05 (17.8)</td>
<td>73.9 (17.9)</td>
</tr>
<tr>
<td>Immediate memory</td>
<td>—</td>
<td>88.84 (24.0)</td>
<td>80.5 (21.9)</td>
</tr>
<tr>
<td>Visuospatial/</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>constructional</td>
<td>—</td>
<td>79.07 (18.5)</td>
<td>75.2 (14.9)</td>
</tr>
<tr>
<td>Language</td>
<td>—</td>
<td>88.12 (12.0)</td>
<td>89.8 (10.7)</td>
</tr>
<tr>
<td>Attention</td>
<td>—</td>
<td>81.42 (18.5)</td>
<td>75.6 (22.6)</td>
</tr>
<tr>
<td>Delayed memory</td>
<td>—</td>
<td>81.91 (22.3)</td>
<td>74.0 (20.7)</td>
</tr>
<tr>
<td>SFS</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>156.6 (16.3)</td>
<td>118.9 (27.9)</td>
<td>113.9 (24.6)</td>
</tr>
<tr>
<td>Social withdrawal</td>
<td>13.1 (1.8)</td>
<td>9.52 (2.6)</td>
<td>8.27 (2.72)</td>
</tr>
<tr>
<td>Interpersonal</td>
<td>—</td>
<td>8.7 (4.6)</td>
<td>6.57 (1.7)</td>
</tr>
<tr>
<td>functioning</td>
<td>—</td>
<td>6.36 (2.16)</td>
<td>—</td>
</tr>
<tr>
<td>Independence</td>
<td>—</td>
<td>34.9 (3.3)</td>
<td>28.8 (6.7)</td>
</tr>
<tr>
<td>(performance)</td>
<td>—</td>
<td>20.6 (6.6)</td>
<td>—</td>
</tr>
<tr>
<td>Independence</td>
<td>—</td>
<td>38.5 (1.1)</td>
<td>32.0 (5.7)</td>
</tr>
<tr>
<td>(competence)</td>
<td>—</td>
<td>31.7 (4.35)</td>
<td>—</td>
</tr>
<tr>
<td>Recreational</td>
<td>24.6 (6.0)</td>
<td>18.1 (7.6)</td>
<td>12.5 (6.9)</td>
</tr>
<tr>
<td>activities</td>
<td>—</td>
<td>28.8 (10.4)</td>
<td>18.2 (8.5)</td>
</tr>
<tr>
<td>Pro-social</td>
<td>9.8 (5.5)</td>
<td>5.19 (3.8)</td>
<td>2.0 (2.2)</td>
</tr>
<tr>
<td>activities</td>
<td>—</td>
<td>96.0 (10.9)</td>
<td>—</td>
</tr>
<tr>
<td>Employment/</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>occupation</td>
<td>—</td>
<td>107.2 (8.38)</td>
<td>97.6 (12.21)</td>
</tr>
<tr>
<td>WTAR predicted IQ</td>
<td></td>
<td>84.05 (10.0)</td>
<td>—</td>
</tr>
</tbody>
</table>

NV, control volunteers; SV-ON, on antipsychotic medication; SV-OFF, off antipsychotic medication; PANSS, Positive and Negative Symptom Scale; RBANS, Repeatable Battery for the Assessment of Neuropsychological Status; SFS, Birchwood Social Functioning Scale; WTAR, Wechsler Test of Adult Reading.

Results

Among the schizophrenia participants, there were two who made more than 200 errors during the very first stage of learning (>3 SDs from the mean of the group, one receiving and one not receiving medication). These outliers were removed from all further analyses.

Learning

Figure 2 shows performance of all remaining subjects across the three stages of the Learning phase. A repeated-measures analysis of variance (ANOVA), with number of errors as the dependent variable and learning stage (1–3) and group (NV, SV-OFF, SV-ON) as independent variables, revealed a trend for a difference in learning between the groups $[F(2,71) = 2.56, p = .08]$, no difference across the learning stages $[F(2,142) = 1.55, p = .21]$, and a trend for a group × stage interaction $[F(4,142) = 2.17, p = .08]$. To further examine differences between the groups, we conducted a more liberal test with a separate ANOVA for each learning stage. This revealed a significant difference between the groups only during the first stage of learning $[F(2,71) = 3.35, p = .04]$, reflecting significantly better performance among the NV than among either of the schizophrenia groups [NV vs. SV-ON, $t(57) = 2.27, p < .05$; NV vs. SV-OFF, $t(55) = 2.23, p < .05$], whereas the SV-ON and the SV-OFF did not differ $[t(52) = 1.07, p = .29]$. By contrast, the effect of group was not significant during either the second $[F(2,71) = 2.44, p = .04]$ or third $[F(2,71) = .56, p = .57]$ stage of learning, revealing that the groups had reached a comparable level of learning before test. Note that the NV showed a trend $[t(19) = 1.8; p = .08]$ toward an increase in number of errors between the second and third learning phase. The different phases of learning differ in the extent to which they test learning for each of the three different trial types (P1–F1, P2–F2, P3–F3, respectively). Thus, this trend is likely due to the increased memory load and increased conflict in the third stage, where subjects must learn a new outcome for an already well-learned stimulus.

Test

Figure 3 shows performance of the three groups during the Test phase. A repeated measures ANOVA revealed a main effect of group $[F(1,71) = 5.53, p < .01]$, a main effect of trial type (“generalized” vs. “trained”), $[F(1,71) = 36.15, p < .001]$, and a group × trial type interaction $[F(2,71) = 4.676, p < .01]$. Post hoc analyses indicated that the interaction was due to significantly worse generalization among the SV-OFF relative to the NV ($p < .01$), whereas the groups did not differ on the “trained” trials (all $p > .50$). No differences were found between the SV-ON and NV on either “trained” or “generalized” trials (all $p$ values > .15).

Consistent with these results, one-sample $t$ tests confirmed that generalization performance was significantly above chance for both the NV and SV-ON groups [NV, $t(19) = 10.47, p < .001$; SV-ON, $t(38) = 4.35, p < .001$] but not for the SV-OFF group ($t < 1.0$).

Relation Between Learning and Generalization

We found no significant relationship between learning rate (number of errors during the Learning phase) and generalization performance. This was the case for all three of the learning stages and when we examined this relationship across all participants (Stage 1: $r = .13$; Stage 2: $r = .08$; Stage 3: $r = .17$; all $p$ values > .20) and for each group separately (NV, Stage 1: $r = .36$; Stage 2: $r = .03$; Stage 3: $r = .18$; SV-OFF, Stage 1: $r = .13$; Stage 2: $r =
Stage 3: \( r = .15 \); Stage 3: \( r = .09 \); SV–ON, Stage 1: \( r = .10 \); Stage 2: \( r = .005 \); Stage 5: \( r = .20 \); all \( p \) values > .13.

**Relation Between Medication Dose and Performance**

We examined the relation between medication dose and performance for individuals in the SV–ON group. Dose was calculated according to two different methods (as described in [31,32]). We found no correlation between dose of medication and performance on either the Learning or the Test phase, for either calculation (Learning, Stage 1: \( r = .10 \); Stage 2: \( r = .16 \); Stage 3: \( r = .22 \); Test, trained: \( r = .01 \) generalized; \( r = .13 \); all \( p \) values > .16).

**Relation of Performance to Demographic and Neuropsychological Measures**

We examined the relation between task performance and all demographic and neuropsychological measures.

**Learning.** Across all subjects, the number of errors during the first stage of learning was correlated with education (\( r = .33 \)), WTAR-IQ (\( r = .41 \)), and with SFS scores (\( r = .41 \) (Bonferroni corrected for multiple comparisons; all \( p \) values < .01). Performance during the second and third stages of learning was not correlated with any demographic measures (all \( r \) values < .20, all \( p \) values > .13).

A separate examination of neuropsychological measures among the patient groups revealed that performance on the first and second stage of learning correlated with RBANS total scores (Stage 1: \( r = .46 \); Stage 2: \( r = .51 \); \( p \) values < .005) and with RBANS delayed memory scores (Stage 1: \( r = .45 \); Stage 2: \( r = .55 \); \( p \) values < .005). No other measures correlated with learning (all \( r \) values < .35, nonsignificant when corrected for multiple comparisons).

**Figure 1.** Sample events and task structure for the learning and generalization paradigm. A single equivalence set is shown here. In the task, participants were trained simultaneously on two sets (four people, four fish). A fish that was correct for one person was incorrect for another person, so the task could not be learned by simple stimulus–response associations.

**Figure 2.** Learning performance across the three shaping stages for healthy control subjects (NV), individuals with schizophrenia tested off-medication (SV–OFF), and individuals with schizophrenia tested on-medication (SV–ON). Both patient groups made more errors in the very first learning stage; however, after this initial phase, the groups did not differ in their ability to learn the associations.

**Figure 3.** Test phase performance for NV, SV–OFF, and SV–ON individuals. All groups performed equally well on the test of previously trained associations. However, the groups differed in their ability to generalize what they learned. Schizophrenic subjects off medication did not generalize what they learned; by contrast, schizophrenic subjects tested on medication generalized well and did not differ significantly from healthy control subjects. Abbreviations as in Figure 2.
**Test.** Across all subjects, performance during Test (trained and generalization trials) was not correlated with any demographic measures (all r values < .20). Given the difference in education levels between the SV–OFF and the NV groups, it is especially important to note that education and generalization were not even weakly correlated ($r = .08; p = .47$).

To further verify that differences in education between SV–OFF and SV–ON do not account for the differences in generalization performance, we conducted a separate analysis of Learning and Test phase performance while equating for education levels in the groups. To this end, we excluded those subjects from the SV–ON and NV groups who had the highest level of education ($\geq$ 16 years), resulting in two new subgroups that did not differ in education from the SV–OFF group (SV–ON: $n = 28$, mean years education $= 12.36, SD = 1.73$; NV: $n = 13$, mean years education $= 12.9, SD = 1.32$; SV–OFF vs. SV–ON, $p = .20$; SV–OFF vs. NV, $p = .49$).

Analyses of task performance among these education-matched groups replicated the findings described in the preceding text. Specifically, the SV–OFF did not differ from either the SV–ON or the NV in performance on the trained trials ($p > .40$). Critically, however, the SV–OFF demonstrated significantly lower generalization performance compared with the NV ($p < .005$) and the SV–ON ($p < .05$).

We also examined the group differences found during Stage 1 of learning among these education-matched groups. This analysis revealed that, among the education-matched groups, there were no significant differences in errors during Stage 1 of learning ($F(2,55) = 2.25; p = .12$).

Finally, when separately examining the neuropsychological measures limited to the education-matched patient groups, performance on the trained trials was correlated with the RBANS-Total Scale (TOT) scores ($r = .44, p < .005$) but not with any other measures. Generalization performance was not significantly correlated with any neuropsychological measure, when correcting for multiple comparisons. With a more liberal, uncorrected threshold, the only measure that showed a relationship with generalization performance was the RBANS-Language Scale (LANG) ($r = .31; p < .05$; all other measures, $p$ values > .40).

**Discussion**

This study sought to examine the effect of schizophrenia and APDs on memory generalization. We found that individuals with schizophrenia are impaired at generalization, an impairment that was significant only among those schizophrenic volunteers who were tested off APDs. In contrast, schizophrenic individuals treated with APDs did not perform significantly worse than healthy control subjects. The effect of both disease and medication was selective to generalization: the on- and off-medication groups showed no impairments on retention of what they learned, and importantly, both groups performed similarly during learning, suggesting that the lack of differences in retention is not explained due to ceiling effects.

**Memory Generalization in Schizophrenia**

The paradigm used here has previously been shown to be specifically sensitive to hippocampal function (25,29,30). The present findings thus complement previously reported data demonstrating impaired hippocampal-dependent generalization in schizophrenia (4,33,34) and extend these prior observations by demonstrating that generalization is sensitive to modulation by antipsychotic medication. This raises important questions regarding the cause of generalization deficits in schizophrenia as well as the mechanism by which antipsychotic medication might impact generalization. In particular, recent fMRI data in healthy individuals demonstrate that hippocampal–midbrain interactions contribute to generalization. Taken together with the present findings, this raises the intriguing possibility that there might be an intrinsic alteration in hippocampal function in schizophrenia that is remediated by APDs. This idea is also consistent with recent theories suggesting that disrupted hippocampal–midbrain function might be a key feature of schizophrenia (35). It is currently unknown as to precisely what effect APDs have on hippocampal function. Future studies exploring the interaction between hippocampal and midbrain dopamine regions in schizophrenia and the impact of APDs on hippocampal function will provide leverage on these speculative possibilities.

**Effect of APDs on Memory and the Hippocampus**

It is a prevailing assumption that antipsychotic medication does not have an action on any aspect of hippocampal-dependent memory in schizophrenia (36). Antipsychotic drugs reduce psychotic symptoms; however, reports regarding their effect on memory and other aspects of cognition have not been documented (37). Here, we demonstrate for the first time that medication might positively impact generalization in a paradigm previously shown to depend on hippocampal function (25,29,30).

This key finding must be understood in light of what is known about APD action in the medial temporal lobe. Antipsychotic drugs block D2 receptors that are located anteriorly in hippocampus and in highest concentrations in dentate gyrus, CA1, and subiculum. Additionally, APDs have been shown to alter long-term potentiation in hippocampal subfields, through D1 dopamine receptor actions (38,39), modulating activity-dependent plasticity (21). Activity within the perforant pathway is modulated by dopamine, affecting the integration of perforant pathway and Schäfer collateral transmission into CA1 (40). Antipsychotic drugs might also have indirect effects on hippocampal transmission by altering activity within the cortical-subcortical long-track pathways. Precisely what part of the pharmacology of dopamine in hippocampus is involved in the actions we report here remains unknown.

The present contrast between SV–ON and SV–OFF shows improved memory-based generalization in individuals tested on APDs, which has not been shown previously. Generalization performance was impaired in the SV–OFF group and was significantly better although still not normal in the SV–ON group. This latter observation supports a speculative therapeutic action of APDs on memory-based generalization in schizophrenia.

**Selective Effects of Schizophrenia and APDs on Memory Generalization**

The present findings indicate that APDs had a selective effect on the ability to generalize learned associations. By contrast, there were no differences between medication groups on feedback-based associative learning or on memory for previously learned associations. Thus, our data suggest that schizophrenia—and APDs—might selectively impact only particular aspects of memory function. The selectivity of this effect might have important implications for understanding the nature and cause of cognitive symptoms in schizophrenia.

Additionally, the present data add to a growing number of studies suggesting that associative learning and generalization might tap into dissociable processes (25,30,41). First, we found that medication selectively impacted generalization but not
learning. Second, we found that performance in the early stages of learning correlated with a number of demographic and neuropsychological measures, whereas generalization did not. This last finding is particularly interesting given that, among volunteers with schizophrenia, both the on- and off-medication subjects made more errors than healthy control subjects on the earliest learning stage. Taken together, these findings suggest that early learning in this task varies on the basis of IQ, age, education, and/or general attentional-learning-set abilities—all abilities that do not seem to contribute directly to generalization processes.

In contrast to generalization, the “learning phase” of the paradigm used here is thought to depend on striatal function based on fMRI and patient studies (25,30). There has been one previous report that medication dose in schizophrenia correlated with number of errors in the learning phase of a task similar to the one used here (33). However, we found neither an effect of medication on learning performance nor a correlation with medication dose, consistent with a more recent report (34). This difference might be related to the overall higher APD doses tested in that study (33), compared with the present one. Future studies are necessary to gain a deeper understanding of how medication dose might differentially impact different aspects of memory function.

Limitations and Future Directions

A potential limitation in interpreting the present findings is that the on- versus off-medication effects were compared between individuals. Because cognitive tasks (particularly learning and memory tasks) are sensitive to learning-set effects, between-subjects designs—as used here—are best suited for this kind of study. However, this design necessitates a consideration of possible differences between the groups other than medication status. In particular, it is important to consider that the groups might differ not only in treatment but in disease state as well. Nonetheless, there are several indications that the effects we found are not due to global differences between the groups. First, the effects are selective to generalization rather than global cognitive differences. Second, the difference between the groups does not seem to be related to any other demographic or neuropsychological measure. Although the off-medication group had on average lower education levels, this does not seem to relate to differences in generalization. Over all participants, education did not correlate with generalization performance. This does not seem to be due to lack of sensitivity in the education measure, because education did correlate with number of errors in the first learning stage. Furthermore, the difference in education levels between the groups was driven primarily by a number of high-education subjects in the on-medication and the control groups. When removing these high-education subjects from the analyses, the effects of medication on generalization performance remain significant.

Conclusions

Although schizophrenia is a complex, multifaceted disorder, the present findings show that cognitive dysfunctions in memory are selective, not global. Therefore, schizophrenia can provide a relatively selective model for certain aspects of cognitive disruption. In particular, the disease and its treatment with APDs might be specifically vulnerable to memory processes that depend on the hippocampus and that might be impacted by dopamine modulation. Taken together with recent functional imaging data, the current results suggest that the hippocampus might be powerfully modulated by dopaminergic mechanisms (either through a direct effect, a systems action, or both) that enable the acquisition of associative memories that support later generalization across experiences, functions that are impaired in schizophrenia.

This work was supported by the National Alliance for Research on Schizophrenia and Depression, National Institute of Mental Health (5R01–MH083039 to ADW; 1 R34 MH75863, SMRI 05–RC–001 to CT, and 5F32MH72135 to DS). The authors are grateful to R. Alison Adcock and Ed Smith for insightful discussion.

The authors report no biomedical financial interests or potential conflicts of interest.

Supplementary material cited in this article is available online.


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