

Hippocampal Volume Reduction in Schizophrenia as Assessed by Magnetic Resonance Imaging

A Meta-analytic Study

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Background: Although many quantitative magnetic resonance imaging studies have found significant volume reductions in the hippocampi of patients with schizophrenia compared with those of normal control subjects, others have not. Therefore, the issue of hippocampal volume differences associated with schizophrenia remains in question.

Methods: Two meta-analyses were conducted to reduce the potential effects of sampling error and methodological differences in data acquisition and analysis. Eighteen studies with a total patient number of 522 and a total control number of 426 met the initial selection criteria.

Results: Meta-analysis 1 yielded mean effect sizes of 0.37 ($P < .001$) for the left hippocampus and 0.39 ($P < .001$) for the right, corresponding to a bilateral reduction of 4%. Meta-analysis 2 indicated that the inclusion of the amygdala in the region of interest significantly in-

creased effect sizes across studies (effect size for the left hippocampus and amygdala, 0.67; for the right, 0.72), whereas variables such as illness duration, total slice width, magnet strength, the use of the intracranial volume as a covariate, measurement reliability, and study quality did not. No laterality differences were observed in these data.

Conclusions: Schizophrenia is associated with a bilateral volumetric reduction of the hippocampus and probably of the amygdala as well. These findings reinforce the importance of the medial temporal region in schizophrenia and are consistent with frequently reported memory deficits in these patients. Future quantitative magnetic resonance imaging studies evaluating the hippocampal volume should measure the hippocampus and amygdala separately and compare the volumetric reduction in these structures to that observed in other gray matter areas.

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THE HIPPOCAMPUS has long been considered an essential component of the medial temporal lobe memory system.^{1,2} The literature on both humans and animals has established that hippocampal lesions result in memory deficits.^{1,2} Patients with schizophrenia also have been shown to suffer from memory dysfunction, and several studies suggest that this selective impairment exceeds the global neuropsychological deficits reported.³⁻⁵ Specific deficits have been found in verbal^{3,5,9} and visual memory,⁵ including semantic memory¹⁰ and encoding,⁸ digit span,^{6,7} and design reproduction.^{6,7,11} Although there is some evidence to suggest that patients with schizophrenia are less significantly impaired on recognition tasks,^{3,6,11} this finding is somewhat equivocal.⁸

Recent studies have attempted to elucidate the anatomical correlates of memory impairment in schizophrenia as part of the general search for neuropathological correlates of the illness. Although some postmortem research has suggested that schizophrenia is associated with abnormal

hippocampal pyramidal cell density¹² and orientation,^{13,14} other studies^{15,16} have failed to support these findings. In addition, schizophrenia has been correlated with smaller neuron size in the CA1 region, subiculum, and layer II of the entorhinal cortex.¹⁷ Abnormalities also have been reported in the mossy fiber pathway,¹⁸ which arises from the axons of hippocampal granule cells and travels to the pyramidal cells of the CA3 region.¹⁹ Other postmortem evidence has suggested that schizophrenia is associated with reductions in overall volume of the hippocampus^{12,20,21} and parahippocampal gyrus.²²

Although postmortem techniques are useful in determining the neurohistological correlates of disease, their findings can be influenced by error variance associated with several variables, including the cause of death. Magnetic resonance imaging (MRI) allows for the in vivo volumetric measurement of various cerebral structures, and these techniques have been applied to the study of schizophrenia. Schizophrenia has been associated with an increased volume of the following brain components: total

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METHODS

STUDY SELECTION

Published reports that evaluated hippocampal volume differences between patients with schizophrenia and normal control subjects using MRI techniques were obtained through the use of computerized databases, including MEDLINE and PsychINFO, and by searching the bibliographies of relevant studies. This search identified 33 studies published between 1988 and 1997. To be included, published studies had to meet several initial criteria: means and SDs for hippocampal volumes had to be reported, a normal comparison group must have been evaluated, the region of interest (ROI) must have included the hippocampus alone or the hippocampal-amygdala complex, the mean patient age must have been between 25 and 40 years, and the study could not have used duplicate subjects from another included study. As many early studies were unable to distinguish between the hippocampus and the amygdala because of technical limitations, studies that measured the 2 structures together were not excluded. Four studies were excluded because of subject duplication,^{33,45,59,60} 3 because means and SDs were not reported,^{50,61,62} 6 because the hippocampal area was measured instead of the volume,^{49,53,63-66} 1 because control subjects were not included,⁶⁷ 1 because the parahippocampal gyrus was included in the ROI,⁶⁸ and 1 because it evaluated elderly patients (mean age, 78.1 years).⁶⁹ Therefore, 18 studies were ultimately included in this meta-analytic study, including data from our laboratory.⁷⁰ Demographic information on patients and normal control subjects is provided in **Table 1**, and scan-related information and mean hippocampal volumes are reported in **Table 2**.

GENERAL META-ANALYTIC METHODS

All analyses were conducted according to the methods of Hedges and Olkin⁷¹ as implemented by DStat,⁷² a commercially

available software program designed specifically for meta-analytic research. The DStat program computes both effect sizes and 95% confidence intervals. All analyses were conducted in several steps. First, Hedges g (an estimate of effect size) was computed for each study by subtracting the mean patient volume (m_p) from the mean control volume (m_c) and dividing by the pooled SD (s) according to the following formula⁷²:

$$(1) \quad g = \frac{m_c - m_p}{s}$$

Although Hedges g does provide an estimate of effect size, it is susceptible to bias introduced by a low sample size.^{72,73} Therefore, DStat was used to transform Hedges g into an unbiased measure of effect size, the Cohen d . Individual values of d were then combined across studies and weighted according to their variance (v) using the following formulas⁷¹:

$$(2) \quad v = \frac{n_1 + n_2}{n_1 n_2} + \frac{d^2}{2(n_1 + n_2)}$$

$$(3) \quad w_i = \frac{l}{v_i}$$

where w_i represents the individual weight for a given study. If more than 1 mean was reported (eg, separate volumes for men and women), a global mean for each hippocampus was computed using the weighted average of the individual means, a strategy that has been used previously in meta-analyses.⁵⁸

Laterality differences were evaluated by converting the effect size for each subset (left vs right) into the Fisher Z_r and computing the difference between the 2 hippocampi (eg, subtracting the Z_r associated with the left hippocampus from the Z_r associated with the right hippocampus) according to the following formulas⁷⁴:

$$(4) \quad r = \text{SQRT}[d^2/(d^2 + 4)]$$

Table 1. Demographic Information for 522 Patients With Schizophrenia and 426 Control Subjects*

Study	Diagnostic Criteria and Patient Type	No.	Age, y†	Illness Duration, y†	Age at Onset, y†	Male Sex, No. (%)
Torres et al, 1997 ⁴⁷	DSM-III-R, inpatients	20	25.6 (6.6)	4.3 (4.8)	NA	14 (70)
Becker et al, 1996 ⁴⁴	DSM-III-R, inpatients	20	27.0 (5.0)	4.25 (6.0)	22.0 (3.0)	20 (100)
Fukuzako et al, 1996 ³⁸	DSM-III-R, chronic type	18	37.8 (5.3)	13.9 (7.7)	23.9 (6.2)	18 (100)
Flashman et al, 1995 ⁷⁰	DSM-IV, inpatients	13	32.8 (9.2)	13.31 (8.04)	19.7 (3.7)	12 (92)
Flaum et al, 1995 ³⁹	DSM-III-R, inpatients	102	31.8 (8.6)	9.7 (8.1)	22.6 (6.4)	70 (69)
Rossi et al, 1994 ³¹	DSM-III, inpatients	19	33.42 (7.32)	11.57 (6.33)	21.84 (4.08)	19 (100)
Zipursky et al, 1994 ²⁶	DSM-III-R, chronic type inpatients	22	34.1 (5.5)	11.3 (6.1)	22.8 (4.3)	22 (100)
Bogerts et al, 1993 ⁴⁰	DSM-III-R, chronic type inpatients	19	27.6 (5.6)	7.8 (5.8)	NA	19 (100)
Kawasaki et al, 1993 ³²	DSM-III-R, inpatients, outpatients	20	28.5 (5.5)	6.72 (4.91)	21.8 (3.9)	20 (100)
McCarley et al, 1993 ⁴²	DSM-III-R, chronic type inpatients	15	38.0 (9.0)	15.8 (8.8)	NA	15 (100)
Breier et al, 1992 ³²	DSM-III-R, chronic type outpatients	44	35.8 (7.0)	14.0 (7.0)	NA	29 (66)
Hoff et al, 1992 ⁵⁴	DSM-III-R, first-episode	43	26.69 (7.03)	NA	NA	27 (63)
Swayze et al, 1992 ⁴⁸	DSM-III, inpatients	53	33.32 (NA)	NA	21.44 (4.44)	35 (66)
DeLisi et al, 1991 ⁵¹	DSM-III-R, chronic type inpatients	15	32.7 (7.2)	NA	24.53 (6.55)	9 (60)
Bogerts et al, 1990 ²⁸	DSM-III-R, first-episode inpatients	35	25.0 (NA)	1.4 (6.0)	NA	22 (63)
Dauphinais et al, 1990 ⁴⁶	DSM-III-R, outpatients	25	32.4 (6.0)	12.8 (5.8)	NA	13 (52)
Suddath et al, 1990 ²⁹	DSM-III-R, chronic type	15	32.4 (5.3)	10.5 (5.5)	22.1 (4.9)	8 (53)
Kelsoe et al, 1988 ⁵⁵	DSM-III, inpatients	24	29.0 (1.0)	8.4 (0.8)	21.0 (0.7)	19 (79)

*NA indicates not available; NIMH, National Institute of Mental Health.

†Data are given as mean (SD).

$$(5) \quad Z_r = 0.5\{\log_e[(1+r)/(1-r)]\}$$

$$(6) \quad q = Z_{r1} - Z_{r2}$$

Significance (*P*) for Cohen *q* was computed in the following way⁷⁵:

$$(7) \quad P = \frac{(Z_1 - Z_0)}{\text{SQRT}[1/(N - 3)]}$$

where $Z_1 = q = Z$ is associated with the experimental hypothesis ($D \neq 0$) and $Z_0 = Z$ is associated with the null hypothesis ($D = 0$). SQRT signifies square root; *D*, overall effect size.

Readers interested in further details of meta-analytic techniques are referred to the work of Hunter and Schmidt,⁷⁷ Hedges and Olkin,⁷¹ Rosenthal,⁷³ and Cooper and Hedges.⁷⁶

Meta-analysis 1

All 18 studies were included in the initial overall meta-analysis. During the initial analyses, however, it was determined that a significant outlier was present in the data set (**Figure 1**). Because the exclusion of outliers serves to increase the accuracy of the overall meta-analytic model through the reduction of extraneous variance,^{71,72} all further analyses were conducted excluding the outlier. An overall effect size (*D*) for each hippocampus was then obtained by computing the weighted mean across the remaining 17 studies.

Meta-analysis 2

A second meta-analysis was conducted to evaluate possible moderator variables (factors that could affect results between studies). Each study was coded on several theoretically relevant variables (mean illness duration, total slice width, magnet strength, the adjustment of hippocampal

volume for either hemispheric or intracranial volume, the inclusion of the amygdala in the ROI, and measurement reliability), and the studies were classified into subsets of each individual variable. As in previous meta-analyses,³⁷ studies were rated on several quality factors that were also treated as possible moderator variables. The first factor evaluated the completeness of the demographic information reported and the degree of patient-control matching. This dimension included patient-control matching by sex, age, handedness, height, weight, education level, and parental socioeconomic status. The reporting of demographic information, including illness duration, age at illness onset, diagnostic criteria, and the source of comparison subjects, was also considered. Studies were assigned 1 point for each item addressed, except in the case of patient-control matching by education level.

Because matching by education level may result in the selection of either high-functioning patients or low-functioning control subjects (or both),^{77,78} studies were assigned no points if such matching was used and 1 point if it was not. The second factor was related to overall scan quality and included magnet strength, slice plus gap width, measurement reliability, the adjustment of hippocampal volume for hemispheric or total intracranial volume, and the inclusion of the amygdala in the ROI. Scores on each of these factors were then transformed into percentages. The third factor measured overall study quality, which was created by averaging the percentages from the first 2 factors.

Potential differences in effect size between variable subsets were analyzed using the method of Hedges and Olkin⁷¹ as implemented by the categorical modeling feature in DStat. This procedure computes mean effect sizes and 95% confidence intervals for each variable subset and allows for the testing of the influence of each individual factor on the overall results. Because possible moderator variables were evaluated in a univariate manner, a Bonferroni correction was used, resulting in a critical *P* value of .005.

Subject Source	No.	Age, y†	Male Sex, No. (%)	Matched for					Parent Socioeconomic Status
				Hand	Height	Weight	Education Level		
Community	19	29.3 (8.9)	9 (47)	No	Yes	No	No	Yes	
NA	20	26.0 (5.0)	20 (100)	Yes	No	No	No	No	
Hospital staff and their relatives	18	38.1 (5.9)	18 (100)	Yes	Yes	Yes	No	No	
Hospital staff, community	12	33.3 (9.1)	11 (92)	Yes	No	No	No	Yes	
Community	87	30.4 (10.6)	45 (52)	Yes	No	No	No	Yes	
Hospital staff, community	14	31.00 (3.92)	14 (100)	Yes	Yes	No	Yes	No	
Community, military veterans	20	36.2 (7.0)	20 (100)	Yes	Yes	No	No	No	
Hospital staff, community	18	28.1 (5.3)	18 (100)	No	No	No	No	No	
Medical university graduates	10	30.0 (4.2)	10 (100)	Yes	Yes	Yes	No	No	
NA	14	38.0 (9.0)	14 (100)	Yes	Yes	Yes	No	Yes	
Community	29	34.4 (8.0)	20 (69)	Yes	Yes	Yes	No	No	
Hospital lobby	24	28.0 (6.7)	14 (58)	Yes	No	No	No	No	
Community	47	NA	28 (60)	Yes	Yes	Yes	Yes	No	
Minor neurological patients	20	28.7 (7.4)	12 (60)	Yes	Yes	No	No	No	
Community, hospital staff, other patients	25	28.2 (NA)	15 (60)	No	Yes	No	No	No	
NIMH personnel, community	20	33.3 (6.6)	10 (50)	No	Yes	No	No	No	
Monozygotic twins	15	32.4 (5.3)	8 (53)	Yes	Yes	Yes	No	Yes	
Hospital staff	14	31.0 (1.0)	10 (71)	Yes	Yes	No	No	No	

Table 2. Scan Information for 522 Patients With Schizophrenia Compared With 426 Control Subjects*

Study	Slice Width, mm	Gap Width, mm	Magnet Strength, T	Volume Adjusted for Total Volume?	Hippocampal-Amygdala Boundary	Mean Hippocampal Volume, cm ³ †				Corrected Effect Size‡ (Cohen d)		
						Reliability		Patients			Control Subjects	
						L	R	L	R	L	R	
Torres et al, 1997 ⁴⁷	1.5	0.0	1.5	No	Separate	0.94	0.94	2.78	3.11	2.78	2.85	L hip 0.00
Becker et al, 1996 ⁴⁴	4.0	0.0	1.5	Yes (by hemisphere)	Together	0.95	0.95	5.49	5.35	6.14	5.97	R hip -0.61 L hip 1.45 R hip 1.27
Fukuzako et al, 1996 ³⁸	1.0	0.0	1.5	Yes (ICV)	Separate	0.92	0.94	2.72	2.80	2.93	3.08	L hip 0.69
Flashman et al, 1995 ⁷⁰	1.5	0.0	1.5	Yes (by hemisphere)	Separate	0.94	0.94	3.40	3.30	3.70	3.76	R hip 0.68 L hip 0.60
Flaum et al, 1995 ³⁹	3.0	1.5	1.5	Yes (ICV)	Separate	0.53	0.53	2.59	2.53	2.67	2.64	L hip 0.21
Rossi et al, 1994 ³¹	5.0	1.0	0.25	No	Together	0.88	0.88	2.03	2.03	2.29	2.21	R hip 0.28 L hip 0.81 L amg/hip 1.06
Zipursky et al, 1994 ²⁶	3.0	0.0	1.5	No (head size/age corrected)	Separate	0.83	0.81	2.01	2.04	1.99	2.07	R amg/hip 0.71 L hip -0.28
Bogerts et al, 1993 ⁴⁰	3.1	0.0	1.0	No	Together	0.85	0.85	3.85	3.89	4.30	4.40	R hip 0.34 L hip 0.81
Kawasaki et al, 1993 ⁵²	5.0	0.0	1.5	No	Separate	0.79	0.79	2.02	2.17	1.94	2.05	R hip 1.14 L hip -0.27
McCarley et al, 1993 ⁴²	1.5	0.0	1.5	Yes (ICV)	Together	0.86	0.86	0.36§	0.38§	0.40§	0.41§	R hip -0.26 L amg/hip 0.80 R amg/hip 0.65
Breier et al, 1992 ³²	3.0	0.0	1.5	No	Separate	0.90	0.90	3.50	3.50	3.80	3.70	L hip 0.61 R hip 0.43
Hoff et al, 1992 ⁵⁴	5.0	2.0	1.5	Yes (total brain volume)	Separate	NA	NA	2.21	2.30	2.55	2.43	L hip 0.46 R hip 0.24
Swayze et al, 1992 ⁴⁸	10.0	0.0	0.5	No	Separate	0.90	0.90	1.45	1.42	1.44	1.47	R hip -0.02 L hip 0.11
DeLisi et al, 1991 ⁵¹	5.0	2.0	1.5	Yes (total brain volume)	Together	0.90	0.90	4.24	4.41	4.44	4.49	L hip 0.35 R hip 0.36
Bogerts et al, 1990 ²⁸	3.1	0.0	1.0	No	Together	0.85	0.85	4.49	4.54	4.70	4.81	L hip 0.36 R hip 0.49
Dauphinais et al, 1990 ⁴⁶	10.0	0.0	0.5	No (adj for age, ht, gender)	Together	0.93	0.93	3.91	3.62	4.20	4.08	L hip 0.31 R hip 0.61
Suddath et al, 1990 ²⁹	5.0	0.0	1.5	No	Separate	NA	NA	1.56	1.58	1.76	1.73	L hip 0.72 R hip 0.46
Kelsoe et al, 1988 ⁵⁵	10.0	0.0	0.5	No	Together	0.97	0.97	7.10	7.00	6.20	6.10	L amg/hip -1.64 R amg/hip -1.76

*ICV indicates intracranial volume; amg, amygdala; adj, adjusted; ht, height; and NA, not available.

†Data are given as mean (SD).

‡Corrected for sample size.

§Relative volume (%).

||Reliability information for the hippocampus was not reported. For other structures, reliability is 0.99.

cerebrospinal fluid,^{23,24} overall ventricular cerebrospinal fluid,^{24,25} lateral ventricles,²⁶⁻³⁰ the third ventricle,^{26,27,29,31} and the caudate.³² Decreases in volume have been found in the following structures: whole brain,²⁴ global gray matter,³³ thalamus,^{25,34} prefrontal cortex and white matter,^{24,32} temporal lobe gray matter,^{26,29} superior temporal gyrus,³⁵ amygdala,^{31,32} parahippocampal gyrus,^{35,36} and corpus callosum.³⁷ Magnetic resonance imaging volumetry has also been applied to the question of hippocampal volume differences in patients with schizophrenia, with inconclusive results. Many studies have reported substantial reductions of hippocampal volume in patients with schizophrenia when compared with normal control subjects,^{28,29,31,32,35,38-46} but others have not.^{26,47-55} In this context, the term “reduced” refers only to a smaller relative volume in patients compared with normal control subjects. Even less clear is the issue of laterality differences in hippocampal volume.⁵⁶ Considering that studies with negative findings are less likely to be published, definitive conclusions regarding hippocampal volume and laterality have remained elusive.

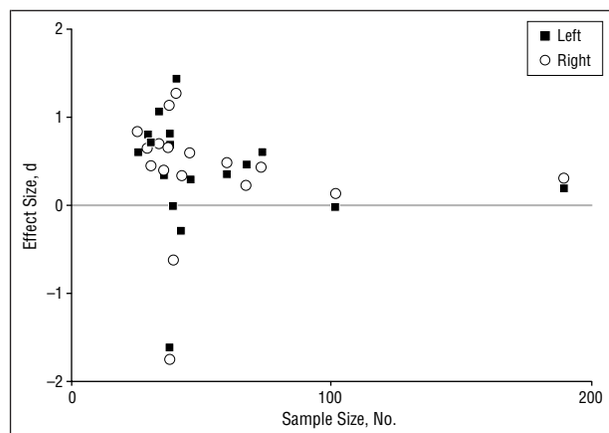


Figure 1. Funnel plot (sample size vs effect size) for the 18 studies included in the initial meta-analysis. Note the presence of an outlier in the data set (lower left).

Table 3. Results of Meta-analyses*

Variable (Subject N/Study N)	Effect Size (Cohen <i>d</i>)		95% Confidence Intervals		Model <i>P</i> †† (Between Variable Subsets)	
	Left	Right	Left	Right	Left	Right
Overall results (910/17)	0.37§	0.39§	0.25<δ<0.52	0.25<δ<0.52
Illness duration						
≤10 y (395/6)	0.34§	0.36§	0.13<δ<0.54	0.15<δ<0.56	.20	.13
>10 y (313/8)	0.52§	0.56§	0.29<δ<0.75	0.33<δ<0.79		
Missing data (202/3)	0.19	0.19	-0.09<δ<0.47	-0.09<δ<0.47		
Magnet strength, T						
1.5 (635/12)	0.38§	0.35§	0.22<δ<0.54	0.19<δ<0.51	.81	.45
<1.5 (275/5)	0.34	0.47§	0.10<δ<0.58	0.22<δ<0.71		
Total slice thickness, mm (slice + gap)						
≤3.1 (341/8)	0.42§	0.45§	0.20<δ<0.64	0.23<δ<0.67	.53	.46
>3.1 (569/9)	0.33§	0.35§	0.16<δ<0.50	0.18<δ<0.52		
Amygdala included in ROI?						
No (631/10)	0.24†	0.25†	0.08<δ<0.40	0.09<δ<0.41	.004	.002
Yes (279/7)	0.67§	0.72§	0.42<δ<0.91	0.47<δ<0.96		
Volume adjusted for ICV?						
Yes (494/8)	0.48§	0.46§	0.30<δ<0.67	0.28<δ<0.64	.07	.27
No (416/9)	0.23	0.30	0.03<δ<0.42	0.11<δ<0.50		
Reliability						
≥0.80 (594/13)	0.42§	0.46§	0.25<δ<0.59	0.30<δ<0.63	.18	.33
<0.80 (219/2)	0.15	0.23	-0.12<δ<0.42	-0.04<δ<0.50		
Missing data (97/2)	0.54§	0.31	0.12<δ<0.96	-0.10<δ<0.72		
Study quality						
Demographics/matching						
High (457/8)	0.41§	0.44§	0.22<δ<0.60	0.26<δ<0.63	.54	.42
Low (453/9)	0.32§	0.33§	0.13<δ<0.51	0.14<δ<0.52		
Scan quality						
High (433/7)	0.30†	0.33§	0.11<δ<0.50	0.14<δ<0.52	.39	.39
Low (477/10)	0.42§	0.44§	0.24<δ<0.61	0.26<δ<0.63		
Overall quality						
High (463/8)	0.33§	0.34§	0.15<δ<0.52	0.15<δ<0.52	.60	.44
Low (447/9)	0.40§	0.44§	0.21<δ<0.60	0.25<δ<0.64		

*ROI indicates region of interest; ICV, intracranial volume; and ellipses, data not applicable.

†Critical *P* = .005.

‡Model *P* refers to the probability that a given variable influenced results across studies for both left and right hippocampi.

§*P* < .001.

||*P* < .005.

There are a number of reasons why results may have been equivocal. For example, the reported differences in hippocampal volume in patients with schizophrenia may be due in large part to sampling error and/or methodological variations such as magnet field strength, slice thickness, and the adjustment of hippocampal volume for either hemispheric or total intracranial volume. One of the best methods for controlling the effects of such mediating variables is the use of meta-analytic techniques.⁵⁷ Meta-analysis provides a quantitative method for integrating research findings within a given body of literature so that a more definitive overall conclusion can be reached. Meta-analytic reviews have recently been conducted on MRI studies of ventricular enlargement,³⁰ corpus callosum size in patients with schizophrenia,³⁷ and ventricular size in patients with bipolar disorder.⁵⁸ Because these studies helped to clarify many unresolved issues, a meta-analytic review of the literature evaluating hippocampal volumes is likely to be a useful tool in addressing the question of brain structural differences in schizophrenia.

This meta-analytic study was conducted to test 2 major hypotheses. First, patients with schizophrenia were predicted to have significantly smaller hippocampal volumes

compared with normal control subjects. Second, we hypothesized that a moderator variable analysis would reveal important variables that could account for the divergence of results in the literature and thus inform future studies.

RESULTS

Results of both meta-analyses are provided in **Table 3**. The mean effect sizes and confidence intervals for left and right hippocampi are reported for each variable subset. Note that model *P* refers to the probability that a given variable affected results across studies for the left and right hippocampi. The results from the overall meta-analysis indicate that schizophrenia is significantly associated with bilaterally reduced hippocampal volume ($D_L=0.37$; $D_R=0.39$; $P < .001$ for both). As shown in **Figure 2**, these results correspond to a bilateral reduction of approximately 4%. The large number of significant mean effect sizes shown in Table 3 reflects the overall consistency of bilaterally reduced hippocampal volume. Similar effects were observed at most levels of the potential moderator variables. No laterality differences were seen in these data.

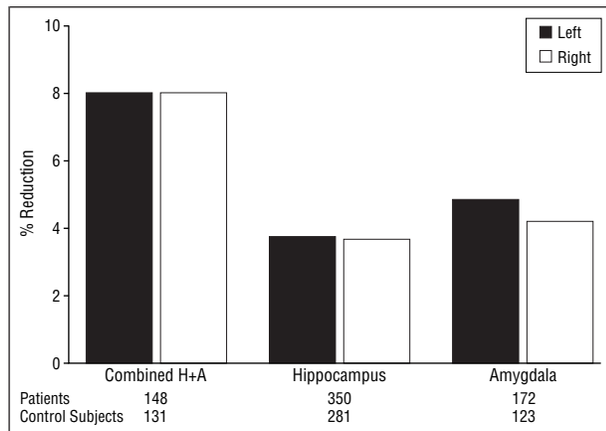


Figure 2. Percentage reduction in the hippocampal-amygdala complex (H+A) in patients with schizophrenia. With 1 exception, the studies measuring the amygdala alone also provided hippocampal measurements. The combined H+A studies do not overlap with the other groups. See the "Comment" section for details.

Only the inclusion of the amygdala in the ROI had a significant ($P < .005$) influence on effect sizes between studies, with those studies that measured both structures together having higher mean effect sizes (left, 0.67; right, 0.72), equaling a bilateral reduction of 8% (Figure 2). To further assess this potential relationship, studies^{32,48,52,54,68} that measured the amygdala separately were evaluated, 4 of which were included in the hippocampal analyses.^{33,48,52,54} Results of this analysis indicated a similar association between schizophrenia and volumetric reduction of the amygdala. The mean effect sizes were 0.29 for the left amygdala and 0.30 ($P < .05$ for both) for the right, equal to an approximately 4.5% reduction in volume (Figure 2). None of the other putative moderator variables had a significant effect on the results.

COMMENT

These meta-analyses produced several main results. First, there is a significant association between schizophrenia and bilateral volumetric reduction of the hippocampus. Second, the inclusion of the amygdala in the ROI significantly increased the magnitude of this relationship. No differences in volumetric reduction were observed between the left and right hippocampi. Finally, it appears that publication bias in favor of positive findings is unlikely to be a significant confounding variable in this literature because many of the studies evaluated in this meta-analysis reported results that were either nonsignificant or opposite the hypothesized direction (Figure 1 and Table 2).

These results have theoretical and methodological implications. Our finding of significant bilateral hippocampal volume reduction is consistent with the nature of the memory and cytoarchitectural abnormalities associated with schizophrenia. Patients with schizophrenia show specific memory deficits consistent with both left^{3-7,9,11} and right^{5-7,11} hippocampal disease. Schizophrenia also has been associated with reduced hippocampal pyramidal cell density, most notably in the left anterior CA3 and CA4 regions.¹²

The finding that the inclusion of the amygdala in the hippocampal ROI significantly increased the mean effect size suggests that volumetric reduction of the amygdala may

also be associated with schizophrenia. Note that in Figure 2 the reduction observed in those studies that measured the hippocampus and the amygdala together ("Combined H+A") was approximately 8%. Because most of the studies in the group in which the hippocampus and the amygdala were measured together had large slice widths (Table 2), however, the greater hippocampus-amygdala reduction observed in these studies could be due to partial volume effects or other measurement factors. Therefore, we recommend measuring the hippocampus and amygdala separately and in their entirety so that the volume of each structure can be assessed independently. The ability of modern scanners to delineate the alveus allows for the accurate separation of the amygdala from the anterior hippocampus.⁷⁹

Although neither the magnet strength nor the slice width had a significant effect on the quantitative volumetric measurement of the hippocampus in this sample, future MRI assessment of relatively small brain structures (ie, hippocampus) should be conducted using the best available scanning technology. For example, the total slice width has been shown to have a significant effect on the volumetric MRI measurement.⁸⁰ Using a calibrated phantom, Luft et al⁸⁰ demonstrated that slice widths greater than 3.0 mm lead to significantly increased error rates. Furthermore, it has been strongly recommended that MRI data be acquired or at least reformatted into a plane perpendicular to the long axis of the hippocampus, a process that helps correct for individual variations in head position.⁷⁹

Although the adjustment of the volume of small cerebral structures for hemispheric or total intracranial volume has been thought to facilitate the accuracy of their measurement,⁷⁹ the results of the moderator variable analysis failed to confirm this hypothesis. Therefore, it is unlikely that the observed differences in the hippocampal volume in patients with schizophrenia are due to a reduction in overall brain volume. Because people with larger heads and brains tend to have larger hippocampi,^{39,79} however, this correction appears warranted because it helps control for the confounding influence of overall brain volume.

Differences in interrater reliability, scan quality, demographic or subject matching quality, and overall quality did not influence the overall effect size in these meta-analyses. This may be due to a range restriction and the lack of variability in these dimensions, which were of consistently good quality across studies. These dimensions are clearly important. Large differences in interrater reliability can contribute to interstudy variation.⁷⁹ Also, the matching of patients and normal control subjects and the reporting of clinical and demographic information are necessary components of any quantitative MRI investigation involving clinical populations.

There are several limitations of this meta-analytic study. First, given the nature of the selection of studies that were included, there is no guarantee that many studies reporting either nonsignificant or countertheoretical results were not published. This problem is unlikely to have significantly influenced the overall conclusions of this study, however, because many of the analyzed studies did report negative results. Second, meta-analytic studies have been criticized for including studies of varying methods.⁵⁷ The inclusion of studies using different methodological techniques in this case revealed an important moderator variable that may account for some of the discrepancies in the published litera-

ture. Third, it was not possible to adequately compare the level of hippocampal volume reduction with that of other gray matter structures (except the amygdala) in these patients. Therefore, it is unclear whether our results are indicative of specific hippocampal volume reduction or a more general reduction in gray matter in patients with schizophrenia.* Fourth, the influence of gender and handedness on hippocampal volume reduction could not be evaluated because of the inclusion of mostly right-handed men in the available studies. Finally, the total number of studies was sufficient for univariate analyses but not for multivariate meta-analytic techniques assessing conjoint and interactive effects of moderator variables.

Future research on the cerebral morphometric correlates of schizophrenia should focus on several issues. First, longitudinal studies should be undertaken to assess the independent effects of patient age and illness duration on the volume of the hippocampus and other structures. Such an approach is important because neuroanatomical abnormalities may be present early in the disease^{25,50,51,67,82} and consequently interact with normal aging. Second, more female and left-handed subjects should be sampled to evaluate possible volumetric differences related to gender and handedness. Third, the relationship between the pattern of clinical symptoms and cognitive profile of patients with schizophrenia and the volume of implicated structures such as the parahippocampal gyrus, amygdala, prefrontal cortex, cingulate gyrus, and thalamus should be investigated. Fourth, concurrent volumetric and functional MRI studies should be undertaken to address the functional consequences and correlates of reduced hippocampal volume and differential role of the left and right hippocampus. Fifth, the volumetric reduction of the hippocampus vs that of other gray matter structures should be compared because, as noted earlier, such reductions have been associated with schizophrenia, including the first episode of the illness.^{50,51,67,82} Sixth, the relative involvement of the anterior and posterior hippocampus in the disease process of schizophrenia should be compared. Because these 2 subdivisions of the hippocampus have different afferent and efferent connections,⁶⁷ it is possible that they may be differentially affected in schizophrenia. Seventh, studies should report the anatomical boundaries used to separate the hippocampus and amygdala in enough detail to allow others to replicate them because differences in these boundaries have the potential to affect volumetry results.⁷⁹ Finally, future studies need to elucidate the underlying mechanism of reduced hippocampal and amygdalar volume in advancing an etiologic understanding of this illness. Recent neurobiological studies in other mammalian species have indicated that environmental factors such as enrichment⁸³ and psychosocial stress⁸⁴ can alter neurogenesis in the hippocampus. The possible developmental interactions among genetic and other neurobehavioral risks and environmental advantages and stressors should be a fruitful area of investigation in schizophrenia.

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REFERENCES

1. Squire LR, Zola-Morgan S. The medial temporal lobe memory system. *Science*. 1991;253:1380-1386.
2. Milner B. Disorders of learning and memory after temporal lobe lesions in man. *Clin Neurosurg*. 1972;19:421-446.
3. Saykin AJ, Gur RC, Gur RE, Mozley D, Mozley LH, Resnick SM, Kester DB, Stafiniak P. Neuropsychological function in schizophrenia: selective impairment in memory and learning. *Arch Gen Psychiatry*. 1991;48:618-624.
4. McKenna PJ, Tamlyn D, Lund CE, Mortimer AM, Hammond S, Baddeley AD. Amnesic syndrome in schizophrenia. *Psychol Med*. 1990;20:967-972.
5. Saykin AJ, Shtasel DL, Gur RE, Kester DB, Mozley LH, Stafiniak P, Gur RC. Neuropsychological deficits in neuroleptic naive patients with first-episode schizophrenia. *Arch Gen Psychiatry*. 1994;51:124-131.
6. Beatty WW, Jovic Z, Monson N, Staton RD. Memory and frontal lobe dysfunction in schizophrenia and schizoaffective disorder. *J Nerv Ment Dis*. 1993;181:448-453.
7. Goldberg TE, Torrey EF, Gold JM, Ragland JD, Bigelow LB, Weinberger DR. Learning and memory in monozygotic twins discordant for schizophrenia. *Psychol Med*. 1993;23:71-85.
8. Gold JM, Randolph C, Carpenter CJ, Goldberg TE, Weinberger DR. Forms of memory failure in schizophrenia. *J Abnorm Psychol*. 1992;101:487-494.
9. Sengel RA, Lovallo WR. Effects of cueing on immediate and recent memory in schizophrenics. *J Nerv Ment Dis*. 1983;171:426-430.
10. Tamlyn D, McKenna PJ, Mortimer AM, Lund CE, Hammond S, Baddeley AD. Memory impairment in schizophrenia: its extent, affiliations and neuropsychological character. *Psychol Med*. 1992;22:101-115.
11. Calev A, Korin Y, Kugelmass S, Lerer B. Performance of chronic schizophrenics on matched word and design recall tasks. *Biol Psychiatry*. 1987;22:699-709.
12. Jeste DV, Lohr JB. Hippocampal pathologic findings in schizophrenia: a morphometric study. *Arch Gen Psychiatry*. 1989;46:1019-1024.
13. Conrad AJ, Abebe T, Austin R, Forsythe S, Scheibel AB. Hippocampal pyramidal cell disarray in schizophrenia as a bilateral phenomenon. *Arch Gen Psychiatry*. 1991;48:413-417.
14. Kovelman JA, Scheibel AB. A neurohistological correlate of schizophrenia. *Biol Psychiatry*. 1984;19:1601-1621.
15. Altshuler LL, Conrad AJ, Kovelman JA, Scheibel AB. Hippocampal pyramidal cell orientation in schizophrenia: a controlled neurohistologic study of the Yakovlev collection. *Arch Gen Psychiatry*. 1987;44:1094-1098.
16. Christison GW, Casanova MF, Weinberger DR, Rawlings R, Kleinman JE. A quantitative investigation of hippocampal pyramidal cell size, shape, and variability of orientation in schizophrenia. *Arch Gen Psychiatry*. 1989;46:1027-1032.
17. Arnold SE, Franz BR, Gur RC, Gur RE, Shapiro RM, Moberg PJ, Trojanowski JQ. Smaller neuron size in schizophrenia in hippocampal subfields that mediate cortical-hippocampal interactions. *Am J Psychiatry*. 1995;152:738-748.
18. Goldsmith SK, Joyce JN. Alterations in hippocampal mossy fiber pathway in schizophrenia and Alzheimer's disease. *Biol Psychiatry*. 1995;37:122-126.
19. Kandel ER. Cellular mechanisms of learning and the biological basis of individuality. In: Kandel ER, Schwartz JH, Jessell TM, eds. *Principles of Neural Science*. 3rd ed. New York, NY: Elsevier Science; 1991:chap 65.
20. Bogerts B, Falkai P, Greve B, Schneider T, Pfeiffer U. The neuropathology of schizophrenia: past and present. *J Hirnforsch*. 1993;2:193-205.
21. Bogerts B, Meertz E, Bausch-Schonfeldt R. Basal ganglia and limbic system pathology in schizophrenia: a morphometric study of brain volume and shrinkage. *Arch Gen Psychiatry*. 1985;42:784-791.
22. Colter N, Battal S, Crow TJ, Johnstone EC, Brown R, Burton C. White matter reduction in the parahippocampal gyrus of patients with schizophrenia [letter]. *Arch Gen Psychiatry*. 1987;44:1023.
23. Gur RE, Mozley D, Resnick SM, Shtasel D, Kohn M, Zimmerman R, Herman G, Atlas S, Grossman R, Erwin R, Gur RC. Magnetic resonance imaging in schizophrenia. I: volumetric analysis of brain and cerebrospinal fluid. *Arch Gen Psychiatry*. 1991;48:407-412.
24. Andreasen NC, Flashman L, Flaum M, Arndt S, Swayze V II, O'Leary DS, Ehrhardt JC, Yuh WTC. Regional brain abnormalities in schizophrenia measured with magnetic resonance imaging. *JAMA*. 1994;272:1763-1769.
25. Corey-Bloom J, Jernigan T, Archibald S, Harris MJ, Jeste DV. Quantitative magnetic resonance imaging of the brain in late-life schizophrenia. *Am J Psychiatry*. 1995;152:447-449.
26. Zipursky RB, Marsh L, Lim KO, DeMent S, Shear PK, Sullivan EV, Murphy GM,

*References 24, 26, 29, 31-33, 35, 36, 56, 81.

- Csernansky JG, Pfefferbaum A. Volumetric MRI assessment of temporal lobe structures in schizophrenia. *Biol Psychiatry*. 1994;35:501-516.
27. Bornstein RA, Schwarzkopf SB, Olson SC, Nasrallah HA. Third-ventricle enlargement and neuropsychological deficit in schizophrenia. *Biol Psychiatry*. 1992;31:954-961.
 28. Bogerts B, Ashtari M, Degreef G, Alvir JMJ, Bilder RM, Lieberman JA. Reduced temporal limbic structure volumes on magnetic resonance images in first episode schizophrenia. *Psychiatry Res Neuroimaging*. 1990;35:1-13.
 29. Suddath RL, Christison GW, Torrey BF, Casanova MF, Weinberger DR. Anatomical abnormalities in the brain of monozygotic twins discordant for schizophrenia. *N Engl J Med*. 1990;322:789-794.
 30. Raz S, Raz N. Structural abnormalities in the major psychoses: a quantitative review of the evidence from computerized imaging. *Psychol Bull*. 1990;108:93-106.
 31. Rossi A, Stratta P, Mancini F, Gallucci M, Mattei P, Core L, Di Michelle V, Casacchia M. Magnetic resonance imaging findings of amygdala-anterior hippocampus shrinkage in male patients with schizophrenia. *Psychiatry Res*. 1994;52:43-53.
 32. Breier A, Buchanan RW, Elkashef A, Munson RC, Kirkpatrick B, Gellad F. Brain morphology and schizophrenia: a magnetic resonance imaging study of limbic, prefrontal cortex, and caudate structures. *Arch Gen Psychiatry*. 1992;49:921-926.
 33. Zipursky RB, Lim KO, Sullivan EV, Brown BW, Pfefferbaum A. Widespread cerebral gray matter volume deficits in schizophrenia. *Arch Gen Psychiatry*. 1992;49:195-205.
 34. Andreasen NC, Arndt S, Swayze V II, Cizadlo T, Flaum M, O'Leary D, Ehrhardt JC, Yuh WTC. Thalamic abnormalities in schizophrenia visualized through magnetic resonance imaging averaging. *Science*. 1994;266:294-298.
 35. Shenton ME, Kikinis R, Jolesz FA, Pollak SD, LeMay M, Wible CG, Hokama H, Martin J, Metcalf D, Coleman M, McCarley RW. Abnormalities of the left temporal lobe and thought disorder in schizophrenia: a quantitative magnetic resonance imaging study. *N Engl J Med*. 1992;327:604-612.
 36. Brown R, Colter N, Corsellis JA, Crow TJ, Frith CD, Jagoe R, Johnstone EC, Marsh L. Postmortem evidence of structural brain changes in schizophrenia: differences in brain weight, temporal horn area, and parahippocampal gyrus compared with affective disorder. *Arch Gen Psychiatry*. 1986;43:36-42.
 37. Woodruff PWR, McManus IC, David AS. Meta-analysis of corpus callosum size in schizophrenia. *J Neurol Neurosurg Psychiatry*. 1995;58:457-461.
 38. Fukuzako H, Fukuzako T, Hashiguchi T, Hokazono Y, Takeuchi K, Hirakawa K, Ueyama K, Takigawa M, Kajiya Y, Nakago M, Fujimoto T. Reduction in hippocampal formation volume is caused mainly by its shortening in chronic schizophrenia: assessment by MRI. *Biol Psychiatry*. 1996;39:938-945.
 39. Flaum M, Swayze VW II, O'Leary DS, Yuh WTC, Ehrhardt JC, Arndt SV, Andreasen NC. Effects of diagnosis, laterality, and gender on brain morphology in schizophrenia. *Am J Psychiatry*. 1995;152:704-714.
 40. Bogerts B, Lieberman JA, Ashtari M, Bilder RM, Degreef G, Lerner G, Johns C, Masiar S. Hippocampus-amygdala volumes and psychopathology in chronic schizophrenia. *Biol Psychiatry*. 1993;33:236-246.
 41. Buchanan RW, Breier A, Kirkpatrick B, Elkashef A, Munson RC, Gellad F, Carpenter WT Jr. Structural abnormalities in deficit and nondescript schizophrenia. *Am J Psychiatry*. 1993;150:59-65.
 42. McCarley RW, Shenton ME, O'Donnell BF, Faux SF, Kikinis R, Nestor PG, Jolesz FA. Auditory P300 abnormalities and left posterior superior temporal gyrus volume reduction in schizophrenia. *Arch Gen Psychiatry*. 1993;50:190-197.
 43. Bogerts B, Falkai P, Greve B. Evidence of reduced temporolimbic structure volumes in schizophrenia. *Arch Gen Psychiatry*. 1991;48:956-957. Letter.
 44. Becker T, Elmer K, Schneider F, Schneider M, Grodd W, Bartels M, Heckers S, Beckmann H. Confirmation of reduced temporal limbic structure volume on magnetic resonance imaging in male patients with schizophrenia. *Psychiatry Res*. 1996;67:135-143.
 45. Becker T, Elmer K, Mechela B, Schneider F, Taubert S, Schroth G, Grodd W, Bartels M, Beckmann H. MRI findings in medial temporal lobe structures in schizophrenia. *Eur Neuropsychopharmacol*. 1990;1:83-86.
 46. Dauphinais D, DeLisi LE, Crow TJ, Alexandropoulos K, Colter N, Tuma I, Gershon ES. Reduction in temporal lobe size in siblings with schizophrenia: a magnetic resonance imaging study. *Psychiatry Res Neuroimaging*. 1990;35:137-147.
 47. Torres IJ, Flashman LA, O'Leary DS, Swayze V, Andreasen NC. Lack of an association between delayed memory and hippocampal and temporal lobe size in patients with schizophrenia and healthy controls. *Biol Psychiatry*. 1997;42:1087-1096.
 48. Swayze VW II, Andreasen NC, Alliger RJ, Yuh WTC, Ehrhardt JC. Subcortical and temporal structures in affective disorder and schizophrenia: a magnetic resonance imaging study. *Biol Psychiatry*. 1992;31:221-240.
 49. Blackwood DHR, Young AH, McQueen JK, Martin MJ, Roxborough HM, Muir WJ, St Clair DM, Kean DM. Magnetic resonance imaging in schizophrenia: altered brain morphology associated with P300 abnormalities and eye tracking dysfunction. *Biol Psychiatry*. 1991;30:753-769.
 50. DeLisi LE, Tew W, Xie S, Hoff AL, Sakuma M, Kushner M, Lee G, Shedlack K, Smith AM, Grimson R. A prospective follow-up study of brain morphology and cognition in first-episode schizophrenic patients: preliminary findings. *Biol Psychiatry*. 1995;38:349-360.
 51. DeLisi LE, Hoff AL, Schwartz JE, Shields GW, Halthore SN, Gupta SM, Henn FA, Anand AK. Brain morphology in first-episode schizophrenic-like psychotic patients: a quantitative magnetic resonance imaging study. *Biol Psychiatry*. 1991;29:159-175.
 52. Kawasaki Y, Maeda Y, Urata K, Higashima M, Yamaguchi N, Suzuki M, Takashima T, Ide Y. A quantitative magnetic resonance imaging study of patients with schizophrenia. *Eur Arch Psychiatry Clin Neurosci*. 1993;242:268-272.
 53. Young AH, Blackwood DHR, Roxborough H, McQueen JK, Martin MJ, Kean M. A magnetic resonance imaging study of schizophrenia: brain structure and clinical symptoms. *Br J Psychiatry*. 1991;158:158-164.
 54. Hoff A, Riordan H, O'Donnell D, Stritzke P, Neale C, Boccio A, Anand AK, DeLisi LE. Anomalous lateral sulcus asymmetry and cognitive function in first-episode schizophrenia. *Schizophr Bull*. 1992;18:257-272.
 55. Kelsøe JR, Cadet JL, Pickar D, Weinberger DR. Quantitative neuroanatomy in schizophrenia: a controlled magnetic resonance imaging study. *Arch Gen Psychiatry*. 1988;45:533-541.
 56. Yurgelun-Todd DA, Kinney DK, Sherwood AR, Renshaw PF. Magnetic resonance in schizophrenia. *Semin Clin Neuropsychiatry*. 1996;1:4-19.
 57. Hunter JE, Schmidt FL. *Methods of Meta-Analysis: Correcting Error and Bias in Research Findings*. London, England: Sage Publications; 1990.
 58. Elkis H, Friedman L, Wise A, Meltzer HY. Meta-analyses of studies of ventricular enlargement and cortical sulcal prominence in mood disorders: comparisons with controls or patients with schizophrenia. *Arch Gen Psychiatry*. 1995;52:735-746.
 59. Nestor PG, Shenton ME, McCarley RW, Haimson J, Smith RS, O'Donnell B, Kimble M, Kikinis R, Jolesz FA. Neuropsychological correlates of MRI temporal lobe abnormalities in schizophrenia. *Am J Psychiatry*. 1993;150:1849-1855.
 60. Goldberg TE, Torrey EF, Berman KF, Weinberger DR. Relations between neuropsychological performance and brain morphological and physiological measures in monozygotic twins discordant for schizophrenia. *Psychiatry Res*. 1994;55:51-61.
 61. Weinberger DR, Berman KF, Suddath R, Torrey EF. Evidence of dysfunction of a prefrontal-limbic network in schizophrenia: a magnetic resonance imaging and regional cerebral blood flow study of discordant monozygotic twins. *Am J Psychiatry*. 1992;149:890-897.
 62. Weinberger DR, Berman KF, Torrey EF. Correlations between abnormal hippocampal morphology and prefrontal physiology in schizophrenia. *Clin Neuropharmacol*. 1992;15(suppl 1, pt A):393A-394A.
 63. Egan MF, Duncan CC, Suddath RL, Kirsh DG, Mirsky AF, Wyatt RJ. Event-related potential abnormalities correlate with structural brain alterations and clinical features in patients with chronic schizophrenia. *Schizophr Res*. 1994;11:259-271.
 64. Marsh L, Suddath RL, Higgins N, Weinberger DR. Medial temporal lobe structures in schizophrenia: relationship of size to duration of illness. *Schizophr Res*. 1994;11:225-238.
 65. Colombo C, Abbruzzese M, Livian S, Scotti G, Locatelli M, Bonfanti A, Scarone S. Memory functions and temporal-limbic morphology in schizophrenia. *Psychiatry Res*. 1993;50:45-56.
 66. DeLisi LE, Dauphinais ID, Gershon ES. Perinatal complications and reduced size of brain limbic structures in familial schizophrenia. *Schizophr Bull*. 1988;14:185-191.
 67. Bilder RM, Bogerts B, Ashtari M, Wu H, Alvir JM, Jody D, Reiter G, Bell L, Lieberman JA. Anterior hippocampal volume reductions predict frontal lobe dysfunction in first episode schizophrenia. *Schizophr Res*. 1995;17:47-58.
 68. Waldo MC, Cawthra E, Adler LE, Dubester S, Staunton M, Nagamoto H, Baker N, Madison A, Simon J, Scherzinger A, Drebing C, Gerhardt G, Freedman R. Auditory sensory gating, hippocampal volume, and catecholamine metabolism in schizophrenics and their siblings. *Schizophr Res*. 1994;12:93-106.
 69. Howard R, Mellers J, Petty R, Bonner D, Menon R, Almeida O, Graves M, Renshaw C, Levy R. Magnetic resonance imaging volumetric measurements of the superior temporal gyrus, hippocampus, parahippocampal gyrus, frontal and temporal lobes in late paraphemia. *Psychol Med*. 1995;25:495-503.
 70. Flashman LA, Saykin AJ, Carrol KE, Bono DB, McAllister TW, Weaver JB, Mamourian A, Kahn EM, Manschreck TC. Relation of hippocampal and neocortical volumetry to functional MRI memory activation in patients with schizophrenia. Paper presented at: 10th Annual Meeting of the Society for Research in Psychopathology; October 28, 1995; Iowa City, Iowa.
 71. Hedges LV, Olkin I. *Statistical Methods for Meta-analysis*. New York, NY: Academic Press Inc; 1985.
 72. Johnson BT. *DStat: Software for the Meta-analytic Review of Research Literatures*. Hillsdale, NJ: Lawrence Erlbaum Associates; 1989.
 73. Rosenthal R. *Meta-analytic Procedures for Social Research*. London, England: Sage Publications; 1986.
 74. Rosenthal R. Parametric measures of effect size. In: Cooper H, Hedges LV, eds. *The Handbook of Research Synthesis*. New York, NY: Russell Sage Foundation; 1994:231-244.
 75. Hays WL. *Statistics*. 5th ed. Fort Worth, Tex: Harcourt Brace College Publishers; 1991.
 76. Cooper H, Hedges LV, eds. *The Handbook of Research Synthesis*. New York, NY: Russell Sage Foundation; 1994.
 77. Resnick SM. Matching for education in studies of schizophrenia [letter]. *Arch Gen Psychiatry*. 1992;49:246.
 78. Goldberg TE, Torrey EF, Weinberger DR. Matching for education in studies of schizophrenia [letter]. *Arch Gen Psychiatry*. 1992;49:246.
 79. Jack CR Jr, Theodore WH, Cook M, McCarthy G. MRI-based hippocampal volumetrics: data acquisition, normal ranges, and optimal protocol. *Magn Reson Imaging*. 1995;13:1057-1064.
 80. Luft AR, Skalej M, Weite D, Kolb R, Klose U. Reliability and exactness of MRI-based volumetry: a phantom study. *J Magn Reson Imaging*. 1996;6:700-704.
 81. Harvey I, Ron MA, Boulay DU, Wicks D, Lewis SW, Murray RM. Reduction of cortical volume in schizophrenia on magnetic resonance imaging. *Psychol Med*. 1993;23:591-604.
 82. Lim KO, Harris D, Beal M, Hoff AL, Minn K, Csernansky JG, Faustman WO, Marsh L, Sullivan EV, Pfefferbaum A. Gray matter deficits in young onset schizophrenia are independent of age of onset. *Biol Psychiatry*. 1996;40:4-13.
 83. Kempermann G, Kuhn HG, Gage FH. More hippocampal neurons in adult mice living in an enriched environment. *Nature*. 1997;386:393-395.
 84. Gould E, McEwen BS, Tanapat P, Patima G, Liisa AM. Neurogenesis in the dentate gyrus of the adult tree shrew is regulated by psychosocial stress and NMDA receptor activation. *J Neurosci*. 1997;17:2492-2498.