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**ORIGINAL PAPER**

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3 **Neurocognitive functioning in the early stages of bipolar**  
 4 **disorder: visual backward masking performance in high risk**  
 5 **subjects**

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■ **Abstract** *Introduction* Cognitive deficits, including deficits in early information processing, are associated with remitted bipolar disorder. The temporal relationship between these deficits and the clinical course is not known. The current study investigated whether or not deficits in early information processing were present before the onset and/or during the early stages of bipolar disorder. *Methods* Unaffected and remitted high risk offspring of well-characterized bipolar parents completed a visual backward masking task. For comparison we included a cohort of unaffected offspring of well parents and a clinically referred group of remitted bipolar patients. *Results* There was no evidence of a deficit in early information processing in well high risk subjects. As expected, the referred patient group had the highest error rates. After excluding the patients, interaction effect showed that the affected remitted high risk subjects performed differently in terms of error rates

than unaffected high risk and control subjects. There were no significant differences in response times across study groups. Exploratory analyses revealed an association between a lifetime history of psychosis and increased errors on the task. *Conclusions* There was no evidence of a vulnerability in early information processing in offspring at risk for bipolar disorder. However, there were emergent changes in performance in the affected remitted high risk group. Psychosis appears to be an important clinical correlate associated with cognitive deficits. Mapping of the early course of bipolar disorder and associated changes in cognition has important implications for establishing critical periods for intervention.

■ **Key words** bipolar disorder · early stages · neurocognitive functioning · high risk · visual backward masking

**Introduction**

There is a substantial agreement in findings across a number of studies that even during periods of remission, adults diagnosed with bipolar disorder show deficits in neurocognitive functioning [26, 27]. This has in part been attributed to a burden of illness effect, likely representing a composite of neurobiological changes associated with repeated episodes and/or residual symptoms, complications of the disorder including substance use and perhaps to the effects of medications [20, 25, 30]. It is not clear how early in the course of illness neurocognitive deficits arise or if there is evidence of cognitive vulnerability predating the onset of a diagnosable mood disorder. Mapping the temporal association between cognitive deficits and the course of bipolar illness may assist with a more accurate early diagnosis and would have important implications for establishing critical periods for intervention and prevention.

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
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71 Deficits in early visual information processing as  
 72 measured by visual backward masking (VBM), have  
 73 been one of the most reliable neurocognitive findings  
 74 in bipolar patients [10, 11, 23]. VBM is a procedure in  
 75 which two brief stimuli are presented in rapid suc-  
 76 cession and the subject is required to identify the  
 77 location of the first stimulus (the target) which is  
 78 disrupted by the later stimulus (the mask) [31]. Under  
 79 certain conditions, performance on the task is medi-  
 80 ated by magnocellular visual pathways projecting to  
 81 the dorsal occipito-parietal and frontal regions [8, 17],  
 82 which have been implicated in structural and func-  
 83 tional imaging studies of patients manifesting and at  
 84 risk for mood disorders [15, 21].

85 The best way to distinguish between primary changes  
 86 associated with the illness from abnormalities secondary  
 87 to the burden of illness is through the use of a high risk  
 88 study design. A number of studies have reported visual  
 89 processing and perception deficits among the unaffected  
 90 relatives of patients with schizophrenia [9, 17]. There  
 91 have been two studies of early visual information pro-  
 92 cessing using a VBM task in subjects at risk for bipolar  
 93 disorder. In one study, Keri and colleagues reported that  
 94 performance on a VBM task was similar between the  
 95 unaffected relatives of bipolar patients and controls [18].  
 96 In a pilot study comparing performance on a VBM task  
 97 between a small group of offspring of bipolar parents  
 98 and controls [22], we reported that while all study sub-  
 99 jects did worse on progressively shorter (more difficult)  
 100 levels of the task, the affected remitted (lifetime  
 101 depressive episodes) offspring made more errors at the  
 102 hardest level of the task compared to unaffected high risk  
 103 and control subjects.

104 These preliminary findings suggested to us that early  
 105 information processing deficits may arise very early in  
 106 the course of bipolar illness, prior to any full-blown  
 107 episodes of mania. The current study was a further  
 108 investigation of early visual information processing  
 109 using a VBM task in a larger similarly recruited high-risk  
 110 sample. We predicted that deficits in early information  
 111 processing would not be present in the unaffected high  
 112 risk subjects, but that differences in performance may be  
 113 present early in the course of illness in the remitted af-  
 114 fected high risk subjects. To examine for cognitive vul-  
 115 nerability in well but at risk offspring we included a  
 116 control group of well offspring of well parents. In addi-  
 117 tion, to examine burden of illness effects we included a  
 118 clinical comparison group of referred bipolar patients.

## 119 Methods

### 120 ■ High risk families

#### 121 Bipolar parents

123 Consenting offspring participating in a longitudinal study of the  
 124 children of bipolar parents described elsewhere [4–6] participated

125 in this neurocognitive study. Briefly, offspring were identified  
 126 through their bipolar parent. Proband parents had been identified  
 127 through their participation in genetic studies [32]. In accordance  
 128 with research protocol, each proband completed a SADS-L [7]  
 129 interview conducted by two research psychiatrists and met DSM-IV  
 130 criteria for either bipolar I or bipolar II disorder. For this study, the  
 131 other parent had no lifetime history of a major psychiatric disorder  
 132 on the basis of SADS-L interviews.

133 The proband's response to long-term lithium was assessed in  
 134 accordance with research protocol [1, 12]. Differences in long-term  
 135 response to lithium is thought to identify a more homogenous  
 136 subgroup of bipolar patients [1, 2] who have differences in clinical  
 137 course and possibly therefore differences in underlying patho-  
 138 physiology. Briefly, lithium responders had to have a highly  
 139 recurrent illness prior to lithium, with no subsequent mood epi-  
 140 sodes while on therapeutic lithium (plasma level of at least  
 141 0.7 mmol/l). Lithium non-responders had to have at least two  
 142 major recurrences associated with therapeutic lithium levels.  
 143 Diagnosis and lithium response of the proband was based on blind  
 144 consensus of at least two research psychiatrists utilizing all relevant  
 145 clinical information.

### High risk offspring

146  
 147  
 148 The offspring completed KSADS-PL [16] interviews conducted  
 149 blind to familial association by a child and adolescent psychiatrist.  
 150 DSM-IV diagnoses were made based on a blind consensus review,  
 151 which included at least two additional psychiatrists, one being a  
 152 child and adolescent psychiatrist. As part of the high-risk study,  
 153 offspring were re-assessed annually or at any time symptoms  
 154 developed. For this study, only those offspring deemed unaffected  
 155 for a lifetime mood disorder or affected with a lifetime bipolar  
 156 spectrum disorder (recurrent major depression, bipolar I or II  
 157 disorder, bipolar nos or cyclothymia) in clinical remission were  
 158 included. Remission was based on the absence of reported and  
 159 observed clinically significant residual signs or symptoms (with no  
 160 impairment of functioning) for at least two consecutive months  
 161 prior to testing (in accordance with DSM-IV full remission crite-  
 162 ria).

163 We included high risk subjects with a diagnosis of recurrent  
 164 unipolar depression, given the parent history of clear-cut bipolar  
 165 disorder and the wealth of evidence supporting the view that  
 166 depression in first degree relatives of bipolar probands represent  
 167 latent bipolar disorder especially if recurrent and/or early in the  
 168 course of illness [3]. Bipolar not otherwise specified (bpnos) in this  
 169 study referred to offspring with a lifetime history of recurrent  
 170 major depression (full DSM-IV criteria) and full-threshold hypo-  
 171 manic episodes, but which fell short of the DSM-IV duration cri-  
 172 teria.

### ■ Comparison groups

#### Low risk offspring

173  
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 176 In order to compare to well individuals not at increased genetic risk  
 177 for bipolar disorder, we selected psychiatrically well offspring of  
 178 well parents recruited through two local schools in Ottawa already  
 179 participating as control families in the ongoing longitudinal study  
 180 previously described [4]. These families were initially identified on  
 181 the basis of a demographic screening questionnaire the parent(s)  
 182 completed that enquired about family composition and about the  
 183 medical and psychiatric lifetime history of both parents and their  
 184 children. Agreeable families completed a screening interview to  
 185 verify the information on the psychiatric and medical health status  
 186 of both parents. Identical to the high risk offspring, consenting  
 187 children from control families completed a KSADS-PL interview

188 conducted by a child and adolescent psychiatrist and on the basis  
189 of a blind consensus review, were determined to be free from  
190 lifetime DSM-IV major psychiatric disorders (Axis I and II) at the  
191 time of neurocognitive testing.

## 192 Referred bipolar patients

193

194 Our main goal was to investigate whether subjects at genetic risk  
195 for mood disorders have a vulnerability in early information pro-  
196 cessing identifiable by abnormalities on VBM testing, therefore we  
197 investigated the offspring of bipolar parents, both affected and  
198 unaffected. In order to test the assay sensitivity of VBM, we also  
199 included a group of referred bipolar patients unselected for family  
200 history. Clinical patient populations are associated with higher  
201 burden of illness than are high risk populations, including an in-  
202 creased risk of neurocognitive impairment. Therefore, we recruited  
203 a consecutive series of patients referred to subspecialty outpatient  
204 clinics in Ottawa and Hamilton. Based on prospective longitudinal  
205 psychiatric assessment these patients met DSM-IV criteria for a  
206 bipolar disorder (bipolar I, II, nos) on the basis of consensus review  
207 and were in clinical remission (not acutely ill with non-impairing  
208 residual signs or symptoms) for a minimum of two consecutive  
209 months (in accordance with DSM-IV) at the time of testing.

## 210 ■ Exclusion criteria for all subjects

211 The exclusion criteria for high risk and control subjects and re-  
212 ferred patients included: (1) history of closed head injury resulting  
213 in loss of consciousness; (2) untreated active medical illness; (3)  
214 identified learning disability or diagnosis of ADHD; (4) lifetime  
215 history of substance dependence; (5) Prior electroconvulsive (ECT)  
216 treatment.

217 All participants in this study were properly informed about the  
218 study and signed a written consent form approved by the respon-  
219 sible research ethics board.

## 220 ■ VBM task

221 The masking task was presented on an IBM compatible micro-  
222 computer with an SVGA3 monitor and circuitry capable of milli-  
223 second timing and following a method that we have previously  
224 employed in adults with established bipolar disorder and in our  
225 pilot high risk study [22, 24]. Visual angles subtended by the  
226 stimuli were approximately 0.57° on the vertical and horizontal  
227 dimensions. The target stimuli were letters (O, S, U, C) presented at  
228 one of four possible target locations (up, down, left or right,  
229 approximately 2.2° of visual angle away from fixation); the mask  
230 consisted of overlapping X's and Os. Subjects identified the location  
231 of the target by pointing in the correct direction using a joystick;  
232 they were not required to identify the target. This condition opti-  
233 mizes transient channel responses. Five blocks of 16 target location  
234 practice trials were presented prior to the task, followed by three

blocks with increasingly short target-to-mask intervals and ending  
with a block of variable target-to-mask intervals identical to the  
format of the upcoming task.

235 Trials began with presentation of a fixation point followed  
236 400 ms later by the appearance of a target 14 ms and then by a 14-  
237 ms mask. The target-mask inter-stimulus intervals (ISIs) were 14,  
238 29, 43, 57, 86 and 114 ms. There were 48 non-practice trials at each  
239 ISI, distributed evenly across the four possible target locations, with  
240 an overall total of 368 trials including all practice trials. Percentage  
241 of incorrect responses and mean reaction time (RTs) at each ISI  
242 constituted the dependent measures.  
243  
244  
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## ■ Statistical analyses

246 We performed chi-square tests and one-way ANOVA tests for  
247 comparison of categorical and continuous demographic and clinical  
248 variables, respectively. Repeated measures analyses of variance  
249 compared RTs and error rates between groups at each ISI. In case  
250 of a significant interaction, we performed one way ANOVA for each  
251 inter-stimulus interval. Since already the nominal *P* values for these  
252 comparisons were non-significant, there was no need to control for  
253 multiple comparisons. We also calculated Cohen's *d* effect size for  
254 pairwise differences between groups.  
255

256 Analyses of variance were also used to compare performance of  
257 high risk subjects with presence versus absence of family history of  
258 response to long-term lithium, personal history of psychotic  
259 symptoms and medication at the time of testing. Pearson's *r* was  
260 used to test for associations between RTs or error rates and age or  
261 clinical scales scores. For some of the exploratory analyses (ie  
262 differences in VBM performance between subjects with versus  
263 without lifetime history of psychotic symptoms, differences be-  
264 tween medicated and non-medicated subjects at the time of testing,  
265 association between symptom scores and VBM performance), we  
266 combined the referred remitted patients and affected remitted high  
267 risk offspring in order to increase power. Multiple linear regression  
268 models were utilized to determine whether parent diagnosis, parent  
269 clinical course, parent lithium response, parent lifetime psychotic  
270 symptoms, parent hospitalization were associated with error rates  
271 in the high- risk offspring. For these analyses we looked at error  
272 rates at 14 ms, the most difficult level of the task.

## Results


### ■ Sample description

273 The study sample consisted of 54 unaffected high risk  
274 offspring, 36 affected remitted high risk offspring, 79  
275 unaffected offspring of well parents (controls) and 23  
276 referred remitted bipolar patients (referred patients)  
277 in this study (refer to Table 1). There was a trend for  
278 differences in proportions of females between groups,  
279 with the largest proportion of females in the affected  
280  
281

**Table 1** Descriptive characteristics of the subjects by group

	High risk affected subjects	High risk unaffected subjects	Referred bipolar patients	Controls	<i>P</i>
<i>N</i>	36	54	23	79	
Age (years)	22.0 (4.2)	18.5 (5.0)	17.87 (2.8)	17.3 (5.8)	0.01
Sex <i>n</i> (%) Females	26 (72.2)	25 (46.3)	12 (52.2)	49.0 (62.0)	0.08
HAMD	2.1 (2.3)	0.9 (1.5)	6.1 (6.8)	0.6 (1.0)	0.01
BDI	4.6 (3.6)	3.5 (3.7)	NA	3.7 (3.1)	NS
CGAS/GAF	84.8 (9.3)	87.3 (7.7)	74.3 (7.5)	90.6 (4.7)	0.01

Values are means and standard deviations in parentheses  
NA not available

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**Table 2** Clinical characteristics of the high risk affected and bipolar patient groups

Variables	Affected remitted high risk subjects (n = 36)	Referred remitted bipolar patients (n = 23)	Significance
Mean age onset index mood episode years (standard deviation)	16.0 (3.4)	14.8 (2.1)	NS
Age at testing years (standard deviation)	22.0 (4.4)	17.9 (2.8)	$P < 0.01$
Mood diagnosis number (%)	BP I = 2 (5.6) BP II = 12 (33.3) BP NOS = 2 (5.6) Cyclothymia = 1 (2.8) Unipolar = 19 (52.8)	BP I = 9 (39.1) BP II = 4 (17.4) BP NOS = 10 (43.5) Cyclothymia = 0 Unipolar = 0	$P < 0.01$
Nature of the clinical course episodic/remitting number (%)	21 (58.3)	2 (10.5)	$P < 0.01$
Mood stabilizer ever number (%)	Yes = 12 (33.3) Anticonvulsant = 2 Atypical antipsychotic = 3 Lithium = 9 Combination = 1	Yes = 21 (81) Anticonvulsant = 5 Atypical antipsychotic = 16 Lithium = 7 Combination = 2	$P < 0.01$
Mood stabilizer at testing number (%)	Yes = 12 (33.3) Anticonvulsant = 1 Atypical antipsychotic = 0 Lithium = 8 Combination = 0	Yes = 19 (100) information about medication at the time of testing missing in 4 subjects Anticonvulsant = 1 Atypical antipsychotic = 11 Lithium = 1 Combination = 6	$P < 0.01$
Psychotic symptoms ever number (%)	Yes = 6 (16.7)	Yes = 12/19 (63.2) information about psychosis missing in four subjects	$P < 0.01$
Hospitalization ever number (%)	Yes = 6 (16.7)	Yes = 8 (34.8)	NS

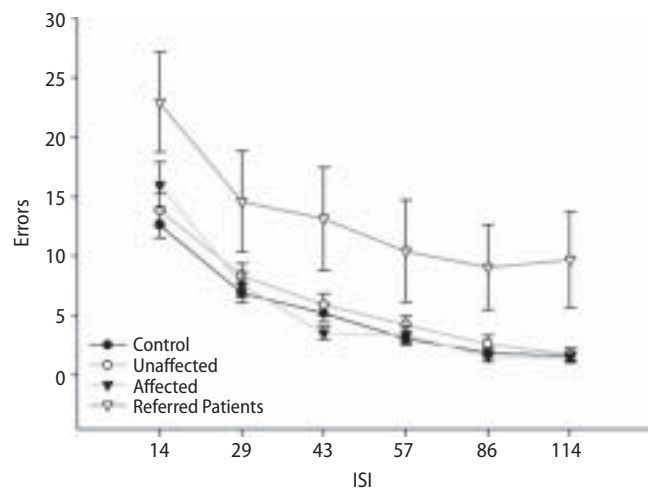
282 high risk group and lowest proportion of females in  
283 the unaffected high risk group. There were differences  
284 between groups in age, with affected high risk subjects  
285 being the oldest and control subjects the youngest.  
286 The global functioning scores were lower in the re-  
287 ferred patient group, whilst these were comparable  
288 between the high risk and control groups.

289 There were significant clinical differences between  
290 the high risk affected remitted group and the referred  
291 remitted patient group (Table 2). Specifically, the re-  
292 ferred patients had more cases of bipolar I disorder, a  
293 higher proportion of cases with a non-episodic illness  
294 course and higher lifetime rates of psychotic features.  
295 Finally, the referred patients had higher rates of life-  
296 time exposure to mood stabilizing medication and  
297 more of the patients were treated with mood stabili-  
298 zers at the time of neurocognitive testing.

299 Neither Hamilton depression rating scale scores  
300 (HAMD) nor the beck depression rating scale scores  
301 (BDI) correlated with numbers of errors or median  
302 response times for any of the groups. Therefore we  
303 did not use these as covariates for the final analyses.  
304 There was no association between age and median  
305 response times or error rates in a combined sample.  
306 Therefore we did not use age as a covariate.

### 307 ■ Error rates between the groups

308 There was a significant main effect of group, such  
309 that the referred patients showed the highest numbers  
310 of errors across all ISIs ( $F = 6.32$ ,  $df = 3$ ; 188,  
311  $P < 0.001$ ), see Fig. 1. The number of errors increased  
312 with shortening of the ISI in all groups (main effect of



**Fig. 1** Error rates across ISIs between groups on the VBM task, four groups (controls, high risk affected, high risk unaffected, referred bipolar patients)

313 ISI  $F = 173.93$ ,  $df = 5$ ; 940,  $P < 0.001$ ). There was no  
314 interaction between ISI and group ( $F = 1.51$ ,  $df = 15$ ;  
315 940,  $P = 0.10$ ).

316 Excluding the patient group resulted in compar-  
317 able error rates between the affected high risk, unaf-  
318 fected high risk and control groups ( $F = 0.51$ ,  
319  $df = 2$ ; 166,  $P = 0.51$ ). There was a main effect of ISI  
320 with more errors at shorter ISIs in all groups  
321 ( $F = 175.43$ ,  $df = 5$ ; 830,  $P < 0.001$ ) and significant  
322 interaction between group and ISI ( $F = 2.02$ ,  $df = 10$ ;  
323 830,  $P = 0.03$ ). Visual inspection of Fig. 1 reveals that  
324 this interaction was likely driven by affected high risk  
325 subjects performing the worst at ISI of 14 ms, but the

326 best among the three groups at the ISI of 43 ms. In  
 327 post hoc comparisons, none of these differences were  
 328 statistically significant even at uncorrected  $p$  levels  
 329 (ISI 14 ms, error rates between unaffected, affected,  
 330 control subjects,  $F = 1.33$ ;  $df = 2$ ; 166,  $P$  uncor-  
 331 rected = 0.27, effect sizes for affected versus control  
 332 subjects Cohen's  $d$  ES = -0.31, affected versus  
 333 unaffected high risk subjects Cohen's  $d$  ES = -0.2,  
 334 ISI 43 ms error rates between unaffected high risk,  
 335 affected high risk, control subjects,  $F = 1.56$ ;  $df = 2$ ;  
 336 166,  $P =$  uncorrected 0.21, effect sizes—affected ver-  
 337 sus control subjects Cohen's  $d$  ES = 0.3, affected  
 338 versus unaffected HR subjects Cohen's  $d$  ES = 0.43.).

### 339 ■ Median response times

340 There were no significant differences in median re-  
 341 sponse times between controls, unaffected high risk,  
 342 affected high risk and referred patients ( $F = 0.64$ ,  $df = 3$ ;  
 343 188,  $P = 0.59$ ). There was a main effect of ISI  
 344 ( $F = 306.31$ ,  $df = 5$ ; 940,  $P < 0.001$ ), with longer median  
 345 response times at shorter ISIs and an interaction be-  
 346 tween ISI and group ( $F = 3.73$ ,  $df = 15$ ; 940,  $P < 0.001$ ),  
 347 such that referred patients had lower reaction times at  
 348 easier levels of the task (see Fig. 2).

349 Excluding the patient group resulted in comparable  
 350 median response times between the affected high risk,  
 351 unaffected high risk and control groups ( $F = 0.13$ ,  $df =$   
 352 2; 166,  $P = 0.88$ ). There was no ISI by group interaction  
 353 ( $F = 0.19$ ,  $df = 10$ ; 830,  $P = 0.99$ ). The prolongation of  
 354 median response times with shortening of the ISIs re-  
 355 mained significant ( $F = 231.82$ ,  $df = 5$ ; 830,  $P < 0.001$ ).

### 356 ■ Exploratory analyses

357 Twenty-nine (54%) of the unaffected high risk sub-  
 358 jects and 21 (58%) of the affected high risk subjects

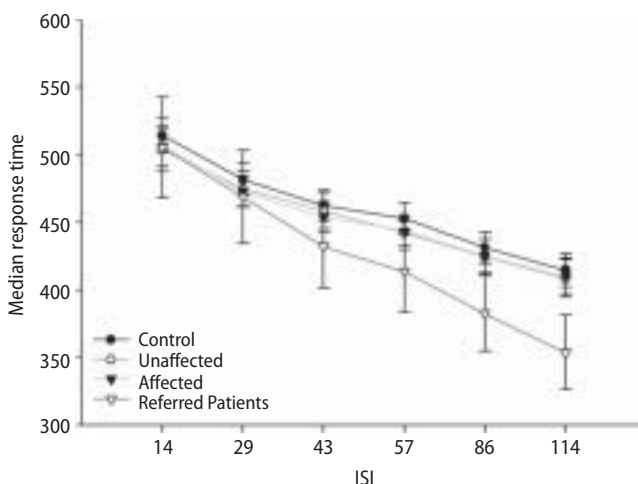


Fig. 2 Median response times across ISIs between groups on the VBM task, four groups (controls, high risk affected, high risk unaffected, referred bipolar patients)

359 derived from lithium responsive families. There  
 360 were no differences in numbers of errors or median  
 361 response times between high risk subjects from  
 362 lithium responsive compared to lithium non-  
 363 responsive families (error rates:  $F = 0.00$ ,  $df = 1$ ; 86,  
 364  $P = 0.96$ ; median response times  $F = 0.61$ ,  $df = 1$ ;  
 365 86,  $P = 0.44$ ), nor was there any interaction between  
 366 lithium response in parents and status of the off-  
 367 spring (affected, unaffected) (error rates  $F = 1.08$ ,  
 368  $df = 1$ ; 86,  $P = 0.30$ ; median response times  
 369  $F = 1.12$ ,  $df = 1$ ; 86,  $P = 0.29$ ). Furthermore, there  
 370 was no ISI by group interaction for error rates or  
 371 median response times.

372 In a multiple linear regression model, none of the  
 373 clinical variables pertaining to the parent clinical  
 374 phenotype, including parent diagnosis, parent clinical  
 375 course, parent lithium response, parent lifetime psy-  
 376 chotic symptoms and lifetime number of hospital-  
 377 izations, were associated with error rates in the high  
 378 risk offspring.

379 There were no differences in either response time  
 380 or error rates between those high risk subjects with a  
 381 lifetime diagnosis of recurrent unipolar versus bipolar  
 382 disorder (error rates  $F = 0.41$ ,  $df = 1$ ; 34,  $P = 0.53$ ;  
 383 median response times  $F = 0.05$ ,  $df = 1$ ; 34,  $P = 0.82$ ).

384 In establishing why referred bipolar patients  
 385 showed poorer performance, we looked at following  
 386 variables: severity of symptoms, lifetime history of  
 387 psychosis, medication at testing. Since the HAMD  
 388 scores fell within a narrow range, we combined all  
 389 subjects in order to increase power to detect even  
 390 small differences. There was no association between  
 391 HAMD scores and either error rates ( $r = -0.08$ ,  
 392  $P = 0.42$ ) or median response times ( $r = 0.09$ ,  
 393  $P = 0.36$ ).

394 Affected remitted subjects (high risk + referred  
 395 patients) with a prior history of psychosis made more  
 396 errors relative to affected remitted subjects (high  
 397 risk + referred patients) without a prior history of  
 398 psychosis ( $F = 4.56$ ,  $df = 1$ ; 53,  $P = 0.03$ ), with no ISI  
 399 by group interaction ( $F = 0.90$ ,  $df = 5$ ; 265,  $P = 0.90$ ).  
 400 There were no differences between affected subjects  
 401 (high-risk + referred patients) with a prior history of  
 402 psychosis compared to those without psychosis in  
 403 median response times ( $F = 0.56$ ,  $df = 1$ ; 53,  
 404  $P = 0.46$ ), although there was an interaction between  
 405 ISI and group ( $F = 2.93$ ,  $df = 5$ ; 265,  $P = 0.01$ ) such  
 406 that a history of psychosis was associated with in-  
 407 creased response times at shorter ISIs.

408 There were no differences between affected sub-  
 409 jects (high risk + referred) taking medication com-  
 410 pared to affected subjects (high risk + referred) not  
 411 medicated at the time of testing (error rates  $F = 1.90$ ,  
 412  $df = 1$ ; 53,  $P = 0.17$ ; median response times  $F = 0.25$ ,  
 413  $df = 1$ ; 53,  $P = 0.62$ ) and no interaction between  
 414 medication and ISI for either error rates or median  
 415 response times (error rates  $F = 0.48$ ,  $df = 5$ ; 265,  
 416  $P = 0.79$ ; median response times  $F = 0.90$ ,  $df = 5$ ;  
 417 265,  $P = 0.48$ ).

418 **Discussion**

419 The major aim of this study was to examine whether or  
 420 not a genetic risk for bipolar disorder was associated  
 421 with abnormalities in early information processing. The  
 422 main finding from this study was no evidence of an  
 423 early visual information processing deficit prior to the  
 424 onset of bipolar disorder in a prospectively studied  
 425 cohort of high risk individuals. That is, there was no  
 426 significant difference in error rates or median response  
 427 times on a visual backward masking task between  
 428 unaffected high risk and control subjects. This finding  
 429 replicates our pilot study [22] and suggests that a deficit  
 430 in early information processing is not a vulnerability  
 431 trait in those at familial risk for bipolar disorder.

432 Consistent with our previous report, affected  
 433 remitted high risk subjects early in the course of  
 434 bipolar disorder, had a different pattern of respond-  
 435 ing on the VBM task compared to unaffected high risk  
 436 and control subjects. That is, the affected remitted  
 437 high risk subjects made more errors than the unaf-  
 438 fected high risk and control subjects at the shortest  
 439 ISI (the hardest level of the task), but they made less  
 440 errors than the other two groups at ISI of 43 ms. In  
 441 post hoc analyses, these differences were not statisti-  
 442 cally significant and their biological significance re-  
 443 mains unknown.

444 The findings in this study are in keeping also with our  
 445 recent structural imaging investigations showing no  
 446 differences in subgenual cingulate volumes or in con-  
 447 centrations of neurochemicals in the anterior cingulate,  
 448 as measured by magnetic resonance spectroscopy be-  
 449 tween high risk (unaffected and affected) and control  
 450 subjects [13, 14]. Taken together these data suggest that  
 451 there is no evidence of a major neurological deficit in  
 452 high risk individuals or in affected individuals early in  
 453 the course of uncomplicated bipolar disorder. These  
 454 findings also concur with findings of Keri and colleagues  
 455 [18] and with other high risk studies reporting generally  
 456 less severe developmental abnormalities compared to  
 457 schizophrenia prior to and during the early stages of  
 458 bipolar disorder [19, 28, 29].

459 In the case of a negative finding, it is crucial to  
 460 demonstrate assay sensitivity or in other words to  
 461 demonstrate that we can detect an abnormality if it  
 462 exists. To this goal, we included a referred patient  
 463 group typically associated with greater burden of ill-  
 464 ness. Compared to high risk affected subjects, the  
 465 patient group had a greater proportion of medicated  
 466 subjects, greater proportion of subjects with psychotic  
 467 mood disorders, and thus greater likelihood of having  
 468 neurocognitive impairments. Indeed the referred pa-  
 469 tients made more errors on the VBM task and showed  
 470 a different pattern of timed reaction during the task.  
 471 This finding demonstrates that we were able to detect  
 472 VBM impairment in the patient group and reassures  
 473 us that the lack of deficit in the high risk and control  
 474 offspring is a true negative finding.

We also attempted to elucidate which variables may  
 underlie the VBM deficits observed among some of the  
 affected high risk subjects and the referred bipolar pa-  
 tients. Exploratory analyses revealed an association  
 between a prior lifetime history of psychotic symptoms  
 and increased error rates. None of the other variables  
 provided an explanation to these differences. In par-  
 ticular, we detected no differences in median response  
 times or error rates between medicated and unmedi-  
 cated subjects at the time of testing. The significantly  
 worse performance among referred patients was not  
 caused by presence of unipolar patients among the high  
 risk subjects, as there were no differences in any VBM  
 measures between bipolar and unipolar offspring of  
 bipolar parents. This is not surprising given the very  
 high likelihood that depression in the high risk popu-  
 lation is genetically related to the bipolar disorder  
 segregating in the family [3]. Likewise severity of  
 symptoms according to HAMD was no associated with  
 VBM performance. The lack of association between  
 symptom levels and VBM was likely due to narrow  
 range of symptom scores, since we selected subjects in  
 remission at the time of scanning.

Limitations of the current study include the con-  
 founding of illness course with prior and current  
 exposure to mood stabilizers. Of relevance here is the  
 fact that none of the subjects with established illness  
 had prior ECT or a history of substance dependence.  
 Secondly, it may be that more sensitive and specific  
 neurocognitive tasks will be able to detect subtle  
 differences in cognition in high-risk subjects or dur-  
 ing the early stages of bipolar illness. This high risk  
 cohort was highly selected and derived largely from  
 intact middle class families therefore may not gener-  
 alize to other high-risk populations. Our post hoc  
 exploratory analyses of variables underlying VBM  
 deficits among referred bipolar patients are pre-  
 liminary and require future replications.

A clear strength of this study lies in the fact that  
 the parents of the high risk and control offspring were  
 comprehensively assessed and that the offspring were  
 followed and repeatedly prospectively studied.  
 Therefore the stability and accuracy of the parent and  
 offspring diagnoses, and of the assessment of clinical  
 status at the time of testing, is at the best possible  
 clinical standard. In addition, the high risk subjects  
 were largely psychotropic drug naïve and all subjects  
 were free from other potential confounders such as  
 prior substance use disorder or learning disabilities  
 and major neurological problems.

There is accruing evidence that bipolar disorder  
 evolves through recognizable clinical stages from  
 non-specific prodromes to early stages characterized  
 largely by depressive episodes followed on average  
 several years later by the onset of activated episodes  
 [4]. Therefore the temporal mapping of neurocogni-  
 tive deficits in the evolution of the illness is impor-  
 tant. From this study and other findings, it appears  
 that neurocognitive functioning is largely intact in



534 uncomplicated bipolar disorder until the clinical onset of illness. Furthermore, these findings highlight the importance of early intervention in order to prevent burden of illness effects. This appears especially true in individuals who experience psychotic symptoms. More research is needed to characterize the early stages of bipolar illness and to understand the mechanism underlying burden of illness effects.

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