

www.neuropsychopharmacology.org

Reply to Pradeep J Nathan and Ben J Harrison

Reply: Pronounced Cognitive Deficits Following an Intravenous L-Tryptophan Challenge in First-degree Relatives of Bipolar Patients Compared to Healthy Controls

S Sobczak¹, A Honig*, and Wim J Riedel¹

Department of Psychiatry, Institute of Brain and Behaviour, Maastricht University, University Hospital Maastricht (AZM), Maastricht, The Netherlands

Neuropsychopharmacology (2003) 28, 2214-2216, advance online publication, 10 September 2003; doi:10.1038/sj.npp.1300270

Sir

Thank you very much for your interest in our paper 'Pronounced cognitive deficits following an intravenous L-Tryptophan challenge in first-degree relatives of bipolar patients compared to healthy controls' (Sobczak et al, 2003b). We appreciate a critical review of the interpretation of our research findings. Next, we will give our response on the three major points of the letter.

GENERAL METHODOLOGICAL CONSIDERATION

If in an experiment, independent variable *X* is manipulated leading to a change in dependent variable Y, experimental methodology only permits conclusions attributing the change in Y to varying levels of X. Suppose that the manipulation of X induces changes in Y, but these are known to be actually accomplished by Z, in such a way that it would have been more appropriate to manipulate Z rather than X? One should attempt to either manipulate/control Z or measure its known indicator(s). According to experimental methodology, one may only speculate but not conclude about the potential mediating influence of Z. Obviously, X refers to tryptophan (Trp) and Y to cognitive performance variables. Z refers primarily to serotonin (5-HT), as the fact that manipulating Trp primarily brings about changes in 5-HT would be beyond dispute (but see point 1). The aim of this experiment and a series of other experiments was to study the role of serotonergic vulnerability, expressed as responsivity to serotonergic

manipulations in healthy first-degree relatives of bipolar patients (FH). We deliberately did not aim at assessing dopaminergic effects as we assumed that the role of dopamine in bipolar disorders is almost beyond dispute. Therefore, it is only of methodological interest if we should have considered dopaminergic factors in the present experiment. Obviously, an experiment similar to the present, in which dopaminergic turnover would be varied, is also of great interest. Perhaps the best approach would be when the two are combined.

However, several other downstream effects of manipulating Trp should be considered as well. 5-HT is known to have inhibitory influences on noradrenaline, acetylcholine, and dopamine turnover (Robbins, 1997). In particular, inhibitions of acetylcholine transmission brought about by increased 5-HT turnover might explain the large effects of Trp on delayed recall memory (Little et al, 1995). On the basis of that, speculation about other potentially mediating neurotransmitters to explain the effect of Trp on memory would point much more into the direction of acetylcholine than dopamine.

POINT 1: SPECIFICITY OF L-TRP CHALLENGE

The specificity of Trp as a serotonergic challenge has been debated several times. Indeed, it has been suggested that increasing Trp in the blood lowers tyrosine and phenylalanine uptake due to competition for the amino-acid transporter (van Praag et al, 1987). Dopamine lowers prolactin release, and the increase in prolactin after Trp is suggested to be due to a decrease in this inhibiting activity of dopamine. Prolactin is only an indirect parameter of brain dopaminergic activity. It has been shown that 5-HT also has direct prolactin-stimulating effects via the 5-HT_{2a/2c} receptors (Di Renzo et al, 1989, van de Kar et al, 1989). The only way to investigate the central effects on these neurotransmitters of Trp infusion is to assess 5-hydroxyindoleacetic acid (5-HIAA) and homovanillic acid (HVA) in

E-mail: adriaan.honig@spsy.azm.nl

Received 04 June 2003; accepted 09 June 2003

Online publication: 16 June 2003 at http://www.acnp.org/citations/

Npp061603030248/default.pdf

^{*}Correspondence: Dr A Honig, Department of Psychiatry, Institute Brain and Behaviour, Maastricht University, University Hospital Maastricht (AZM), PO Box 5800, Maastricht, The Netherlands, Tel: +31 43 3877537, Fax: +31 43 3875444,



the cerebrospinal fluid. These are invasive methods and, to our knowledge, such a study has never been described. Another indirect method is measurement of HVA in blood. For each HVA measurement, 1 ml blood is needed. As the amount of blood taken from subjects is limited and other blood parameters (ie hormones, amino acids) are also important to assess, we chose not to include HVA

Large neutral amino acids (LNAAs; leucine, valine, isoleucine, Trp, tyrosine, phenylalanine) compete for transporter uptake at the level of the blood-brain barrier. The ratios of Trp/LNAAs, phenylalanine/LNAAs, and tyrosine/ LNAAs might give another indication of the availability of serotonergic, respectively dopaminergic, precursors in the brain. In the present study, the ratio of Trp/LNAAs increased significantly to a value of 2.3 from a baseline value of 0.13. This is an increase of 1669%. The ratio tyrosine/LNAAs decreased after Trp loading, but this was not significant (F(1,42) = 0.017, NS). The ratio phenylalasignificantly was lower (F(1,42) = 88.98, P < 0.05). The ratio decreased from 0.11 at baseline to 0.0013. This is a decrease of 103%. As the increase in the ratio of Trp is much higher than the decrease in the ratio of phenylalanine, and the ratio of tyrosine did not change at all, we suggest that the effects of the Trp loading are more likely attributable to central serotonergic effects than to dopaminergic effects.

5-HT acts as a neuromodulator, and important functional interactions between brain 5-HT and dopaminergic systems are known. 5-HT inhibits dopaminergic activity in the mesolimbic system but stimulates this activity in nigrostriatal structures (Manji and Potter, 1997). Thus, we cannot fully exclude that other neurotransmitters are involved in the cognitive effects of Trp and 5-HT.

Despite this we suggest that the cognitive effects of Trp challenge are primary serotonergic mediated because the detrimental effects of Trp on attention and planning in healthy FH have also been found after acute tryptophan depletion (ATD) (Sobczak et al, 2002c). Thus, it seems that both an increase and decrease in Trp impair the same cognitive domains that are regulated by the frontal lobe (Sobczak et al, 2003b). The specificity of ATD has been pointed out by Klaassen et al (1999). They found a decrease in mood and impaired memory performance after ATD but not after lysine depletion. This supports the hypothesis that ATD affects brain 5-HT functioning and not brain protein metabolism in general.

POINT 2: COGNITIVE DEFICITS ON BASELINE ARE MEDIATED BY DOPAMINE

The second suggestion made by Nathan and Harrison (2003) is that the baseline cognitive deficits in FH may be a result of dysregulation of dopaminergic activity in mesocortical regions.

It must be emphasized that the FH subjects in this study were free of any clinical psychiatric symptoms. Thus, cognitive impairments were not related to altered mood states (Sobczak et al, 2003b). Therefore, if there was a dopaminergic dysfunction in these subjects at baseline, it would only have affected cognition and not mood.

Independent of Trp, cognitive performance was more impaired in relatives of type I bipolar patients (FH I) compared to relatives of type II patients (FH II). This suboptimal baseline performance in FH I resulted in more pronounced cognitive deficits after Trp (ie on planning). These findings agree with an association of serotonergic vulnerability and cognitive impairments in FH I subjects. FH II subjects were characterized by mood changes after ATD and Trp challenge (Sobczak et al, 2002a, b). Taking these findings together, we speculate that FH II subjects share more symptoms of primary affective disorders in which 5-HT plays a prominent role, whereas FH I subjects show characteristics of primary psychotic disorders in which cognitive deficits persist and functional deficits in other neurotransmitters like noradrenaline, acetylcholine and dopamine, or even structural brain abnormalities, may also be involved.

POINT 3: FUTURE STUDY ON THE ROLE OF 5-HT AND **DOPAMINE**

Indeed the search for biological markers of bipolar disorders is just in its infancy. The present findings must first be replicated in a larger research population including relatives of type I and type II bipolar patients. Of interest is whether there is a biological distinction between type I and type II patients and their relatives. The vulnerability to serotonergic, dopaminergic, and noradrenergic dysfunction should be investigated using ATD, specific serotonergic challenge procedures, and a phenylalanine/tyrosine depletion test. Abnormalities in cholesterol and fatty acids have also been associated with altered brain neurotransmitter activity and psychopathology (Sobczak et al, 2003a; Hibbeln and Salem, 1995; Swartz, 1990). Thus, research into the interaction of neurotransmitters, cholesterol, and fatty acids in humans and their association with psychopathology is still a challenge.

REFERENCES

Di Renzo G, Amoroso S, Taglialatela M, Canzoniero L, Basile V, Fatatis A et al (1989). Pharmacological characterization of serotonin receptors involved in the control of prolactin secretion. Eur J Pharmacol 162: 371-373.

Hibbeln JR, Salem N (1995). Dietary polyunsaturated fatty acids and depression: when cholesterol does not satisfy. Am J Clin Nutr 62: 1-9.

Klaassen T, Riedel WJ, Deutz NEP, van Someren A, van Praag HM (1999). Specificity of the tryptophan depletion method. Psychopharmacology 141: 279-286.

Little JT, Broocks A, Martin A, Hill JL, Tune LE, Mack C et al (1995). Serotonergic modulation of anticholinergic effects on cognition and behavior in elderly humans. Psychopharmacology 120: 280-288.

Manji HK, Potter WZ (1997). Monoaminergic systems. In Young LT, Joffe RT (eds). Bipolar Disorders. Biological Models and their Clinical Application. Marcel Dekker Inc., New York.

Nathan PJ, Harrison BJ (2003). L-Tryptophan challenge and cognitive deficits in bipolar disorder: evidence for hyperserotonergic and hypodopaminergic mechanisms (letter). Neuropsychopharmacology (in press).

Robbins TW (1997). Arousal systems and attentional processes. Biol Psychol 45: 57-71.

- Sobczak S, Honig A, Christophe A, Maes M, Helsdingen RWC, De Vriese S et al (2003a). Lower high-density lipoprotein cholesterol and increased ω6 polyunsaturated fatty acids in first-degree relatives of bipolar patients. Psychol Med in press.
- Sobczak S, Honig A, van Duinen MA, Riedel WJ (2002a). Mood, prolactin and cortisol responses following intravenous Ltryptophan challenge; evidence for serotonergic vulnerability in first-degree relatives of bipolar patients. Int J Neuropsychopharmacol 5: 249-254.
- Sobczak S, Honig A, Nicolson NA, Riedel WJ (2002b). Effects of acute tryptophan depletion on mood and cortisol release in firstdegree relatives of type I and type II bipolar patients and healthy matched controls. Neuropsychopharmacology 27: 834-842.
- Sobczak S, Honig A, Schmitt JAJ, Riedel WJ (2003b). Pronounced cognitive deficits following an intravenous L-tryptophan challenge

- in first-degree relatives of bipolar patients compared to healthy controls. Neuropsychopharmacology 28: 711-719.
- Sobczak S, Riedel WJ, Booij L, Aanmet Rot M, Deutz NEP, Honig A (2002c). Cognition following acute trytophan depletion: differences between first-degree relatives of bipolar disorder patients and matched healthy control volunteers. Psycho Med 32: 503-515.
- Swartz CM (1990). Albumin decrement in depression and cholesterol decrement in mania. J Affect Disord 19: 173-176.
- Van de Kar LD, Lorens SA, Urban JH, Bethea CL (1989). Effect of selective serotonin (5-HT) agonists and 5-HT2 antagonist on prolactin secretion. Neuropharmacology 28: 299-305.
- Van Praag HM, Lemus C, Kahn R (1987). Hormonal probes of central serotonergic activity: do they really exist? Biol Psychiatry