

## Retropulsion, GABA, and Possible Hallucinatory Behavior in Mice

Several hypotheses<sup>1,2</sup> have appeared which associate CNS levels of various aromatic amines with hallucinatory behaviors. Psychopharmacological tests of these hypotheses often use abnormal postures, twitches<sup>3</sup>, or gaits<sup>3</sup> in rodents as indices of hallucinations. The present paper describes one such index – retropulsion in mice – which was observed after administration of amino oxycetic acid (AOAA), an agent which increases<sup>4</sup> endogenous CNS levels of GABA by competitively inhibiting –  $\alpha$  – ketoglutarate transaminase<sup>5</sup>.

**Experimental.** One hundred fifteen 16-day-old C57BL/6J mice were s.c. injected with either saline (control) or AOAA (experimental). The dose of 20 mg/kg of AOAA (dissolved in unbuffered saline solution) has repeatedly<sup>6</sup> been demonstrated to increase brain GABA in juvenile C57BL/6J mice. After injections, subjects were placed in individual clear plastic cages and observed for 5 h. Although the experimenter was aware of which mice received AOAA, a subsequent double blind procedure has verified the results described below.

**Results.** All 92 experimental subjects exhibited hypothermia and immobility within 15 min of AOAA injections. Approximately half the mice also showed mild tremors and/or a complete extension of the tail in an anterior-posterior direction. The 23 control animals typically assumed a sleeping posture with their tails partially wrapped around their bodies.

After 15–30 min, 56 of the 92 experimental subjects expressed clonic or clonic-tonic convulsions, and 39 experimental subjects showed definite retropulsion (walking backwards) which began approximately 90 min after injections. The Table gives a more detailed breakdown of these data. Although it was not necessary for a mouse to convulse before exhibiting retropulsion, the two behaviors were correlated ( $C^2 = 0.31$ ;  $\chi^2 = 9.96$ ;  $df = 3$ ;  $p < 0.02$ ). Although the convulsive phase of the syndrome was stereotyped, the retropulsions were highly individualistic; some mice crawled backwards, while others moved with hopping-like motions. In most cases, only the forelimbs were involved, although some animals also used the hind

limbs. None of the 23 control subjects exhibited convulsions or retropulsions.

**Discussion.** Previous studies<sup>8</sup> have described the convulsive effects of AOAA but, to my knowledge, the present paper is unique in its observation of retropulsion in response to this agent. Insofar as could be ascertained, retropulsion in mice has only been reported twice in the psychopharmacology literature, and both reports associated it with hallucinogenic behavior. WOOLEY<sup>3</sup> described retropulsion in mice as resulting from an apparent 'hallucination of sliding down an inclined plane' due to their LSD treatment. He also referenced<sup>9</sup> a study in which LSD induced Siamese fighting fish to swim backwards.

The time course of retropulsion corresponds rather well with several other behavioral effects of AOAA. Increased threshold for the auditory startle reflex<sup>9</sup>, elevation of the electroconvulsive threshold<sup>8</sup>, and protection from certain behavioral aftereffects of juvenile acoustic trauma<sup>10</sup> are all maximal approximately 90 min after injection of a comparable dose of AOAA. Although it requires approximately 5 h<sup>8</sup> for AOAA to produce maximal elevations of brain GABA, the rate of GABA increase is highest 90 min after injection.

Present theories concerning the mechanisms of psychotomimetic agents are often couched in terms of their effects on indole amines and catecholamines. If retropulsion is a valid index of hallucinatory activity, and is causally related to the effects on brain GABA, altered states of consciousness may be more dependent upon relative levels of all the biologically active central amines, and not just of the aromatic amines.

**Zusammenfassung.** Amino-oxycetsäure, eine Substanz welche das endogene Gehirnniveau an GABA erhöht, führte in 16 Tage alten Mäusen zu LSD-ähnlichen Verhaltensweisen wie Hypothermie; zu Krämpfen und Rückwärtslaufen.

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Relationship of convulsion and retropulsion in 92 mice treated with AOAA

Maximum severity of convulsions	No mice showing associated retropulsion	No mice showing no retropulsion
Clonic-tonic	20	15
Clonic	11	10
No convulsion	8	28

<sup>1</sup> D. W. WOOLEY, *The Behavioral Bases of Psychoses* (Wiley, New York 1962).

<sup>2</sup> S. J. CORNE and R. W. PICKERING, *Psychopharmacology* 11, 65 (1967).

<sup>3</sup> D. W. WOOLEY, *Proc. natn. Acad. Sci., USA* 41, 338 (1955).

<sup>4</sup> D. P. WALLACH, *Biochem. Pharmacol.* 5, 166 (1960).

<sup>5</sup> N. M. VAN GELDER, *Biochem. Pharmacol.* 15, 533 (1966).

<sup>6</sup> P. Y. SZE, in *Physiological Effects of Noise* (Eds. B. L. WELCH and A. S. WELCH; Plenum Press, New York 1970), p. 259.

<sup>7</sup> S. SIEGEL, *Nonparametric Statistics* (McGraw-Hill, New York 1956), p. 196.

<sup>8</sup> K. KURIYAMA, E. ROBERTS and M. K. RUBINSTEIN, *Biochem. Pharmacol.* 15, 221 (1966).

<sup>9</sup> R. P. BOBBIN, G. GONZALEZ and P. S. GUTH, *Nature, Lond.* 223, 70 (1969).

<sup>10</sup> K. R. HENRY, unpublished manuscript (1971).

## Phenitron and Marijuana Induced Hypothermia

Phenitron [3-(Hexahydro-1H-azepin-1-yl)-3'-nitropropionophenone hydrochloride], at a dose of 10–15 mg/kg i.p., has been reported to prevent the development of catalepsy in dogs exposed to the smoke of burning hashish. Animals first rendered cataleptic with hashish reverted to a normal behavioural state 6–10 min after administration of

phenitron (20 mg/kg i.p.). The concentration of tetrahydrocannabinols in the hashish was not determined but the dose administered was equivalent to 2–3 times the threshold dose<sup>1</sup>.

GRUNFELD and EDERY<sup>2</sup> have reported profound behavioural changes in dogs and monkeys following i.v. ad-