#### Original Investigations

# Dopaminergic Supersensitivity After Neuroleptics: Time-Course and Specificity

Pavel Muller and Philip Seeman

Department of Pharmacology, Faculty of Medicine, University of Toronto, Toronto, Ontario, Canada, MSS 1A8

Abstract. It is known that a single dose of a neuroleptic can elicit dopaminergic supersensitivity in animals. On the other hand, the clinical syndrome of tardive dyskinesia takes many months or years to develop. To resolve this apparent discrepancy, it is possible that abclinical or latent tardive dyskinesia is fully compensated in most patients taking neuroleptics. In others, where the tardive dyskinesia is full-blown and prossly apparent, the dopaminergic supersensitivity may be decompensated. Such compensatory and decompensatory phases have been proposed earlier by Hornykiewicz (1974), in the case of Parkinson's Disease.

Dopaminergic supersensitivity persists for a period proportional to the length of the neuroleptic treatment. It is not yet clear whether the relation between the eight of treatment and the persistence of superensitivity holds for very long treatments, but in principle the relationship might account for the persistence of tardive dyskinesia after years of neuroleptic pretreatment.

**Ley words:** Tardive dyskinesia — Dopamine receptors
Stereotypy

## Time-Course of Development of Dopaminergic Supersensitivity

There appears to be a correlation between the timeourse of development of tolerance to a neuroleptic 7th the rate of development of dopaminergic supersensitivity. For example, according to Ezrin-Vaters and Seeman (1977), tolerance of catalepsy to aloperidol develops rapidly over the first five days and ten develops more slowly. While the development of this tolerance may to some extent be accounted for by arning from test to test, it correlates well with the rate of development (Lerner and Nosé, 1977; Asper et al., 1973) of dopaminergic supersensitivity.

Although the time-course of development of dopaminergic supersensitivity has received some attention, there is little or no information on the rate of development of dopamine/neuroleptic receptors in the first days of neuroleptic treatment. For example, Christensen et al. (1976) reported an increase in sensitivity to apomorphine-induced stereotypies within a day or two after single injection of chlorpromazine or haloperidol; similar results were reported on climbing behavior by Costentin et al. (1977) and Martres et al. (1977). However, detailed information on the timecourse of development of the receptor alterations after repeated neuroleptic administration has not yet been reported. The shortest treatment schedule hitherto reported was by Burt et al. (1977), who treated rats with haloperidol for 7 days and then withdrew them for five days. By that time it was found that the <sup>3</sup>H-haloperidol receptors had already achieved their maximum increase (Table 1).

This rapid development of dopaminergic supersensitivity in animals (albeit at massive doses) is faster than the rate of development of tardive dyskinetic symptoms in patients. This is one of the main reasons why Tarsy and Baldessarini (1977) feel that neuroleptic-induced dopaminergic supersensitivity (in animals) may not be an appropriate model for tardive dyskinesia.

According to Crane (1973), the development of tardive dyskinesia within the first 6 months of treatment is unusual and most of the patients with tardive dyskinesia developed their symptoms after neuroleptic treatment for one year or more. Tarsy and Baldessarini (1977) suggest, therefore, that the dopaminergic supersensitivity seen after repeated neuroleptic treatment of animals is a better model for acute dyskinesia. This dyskinesia appears within 2 – 5 days after the initiation of the neuroleptic treatment (Fig. 1).

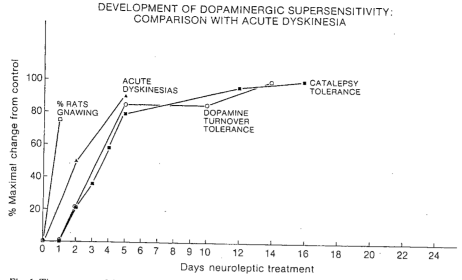


Fig. 1. Time-course of dopaminergic supersensitivity. Maximal observed change from the control was taken as 100 %. Rat gnawing: Christenson et al. (1976). Acute dyskinesias: Marsden et al. (1975). Turnover tolerance: Lerner and Nosé (1977). Catalepsy tolerance: Ezrin-Watersam (1977)

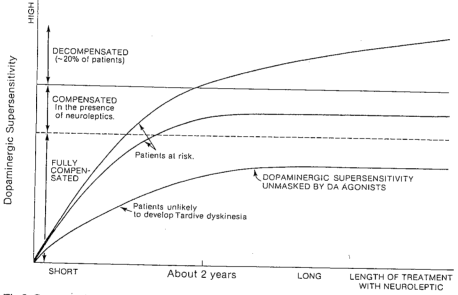
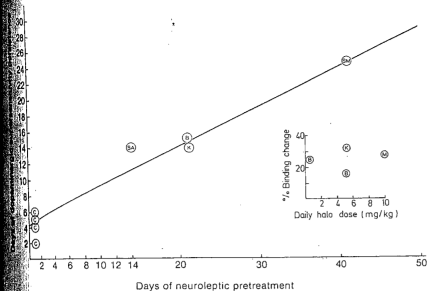


Fig. 2. Compensation of dopaminergic supersensitivity in tardive dyskinesia — a model. Decompensated dopaminergic supersensitivity leads to spontaneous appearance of dyskinetic symptoms. Dopaminergic supersensitivity compensated in the presence of neuroleptics will be clinically dormant until neuroleptics are discontinued or dose is lowered. Fully compensated dopaminergic supersensitivity could be precipitated to dopamine agonists or anticholinergic drugs

In order to demonstrate behavioural dopaminergic supersensitivity in rats which have received long-term neuroleptics, it is necessary to challenge them with either dopamine-mimetic drugs or anticholinergic drugs (Tarsy and Baldessarini, 1974; Gianutsos and Lal, 1976). This is because such rats do not spontaneously show stereotypy. Similarly, many patients on long-term neuroleptics may not spontaneously exhibit any obvious dyskinetic signs in the early stages. Such

patients may have a latent or subclinical dyskinesh which is fully compensated by certain adaptations the brain (see Fig. 2).

This suggestion of a latent compensated form tardive dyskinesia is analogous to the early compensated phase of Parkinson's Disease, as proposed Hornkiewicz (1974). In this early stage of Parkinson Disease, it is thought that the dopaminergic cell losg counterbalanced by several compensatory change, in



1.3. Correlation of the persistence of dopaminergic supersensitivity with length of neuroleptic pretreatment. C.: Christensen et al. (1976), the correlation of the persistence of dopaminergic supersensitivity with length of neuroleptic pretreatment. C.: Christensen et al. (1976), the correlation of the daily dose with maximal neuroleptic binding increase over controls, Muller and Seeman.

wrotransmitter function. Thus, in order to unmask clatent dyskinesia, it is necessary to challenge acutely ith L-Dopa or to block any cholinergic compensations anticholinergic drugs. It seems reasonable to think such compensatory mechanisms would effectively ask the latent tardive dyskinetic state for many onths or years. Thus, the apparent discrepancy in mecourse between the onset of dopaminergic super-instituty, which is a matter of days or weeks, and the sect of frank dyskinesia, which is a matter of months a years, may be accounted for by these compensatory echanisms.

### Platination of Neuroleptic-Induced Dopaminergic spersensitivity and of Tardive Dyskinesia

has been stated that tardive dyskinesia may be reversible or poorly reversible (Crane, 1973), also others report dissipation of the dyskinesia thin several months (Quitkin et al., 1977). On the ter hand, dopaminergic supersensitivity (in rats) and by about a month's treatment with neuro-lie disappears within 2-4 weeks after withdrawal 3).

According to Crane (1973), tardive dyskinesia symptoms are either irreversible or very poorly reversible.

Withdrawal after the dyskinetic symptoms are observed improves the prognosis (Crane, 1973; tim et al., 1977). When patients are under concurrent medical observation (Quitkin et al., 1914) and withdrawn within a median time of 1 month,

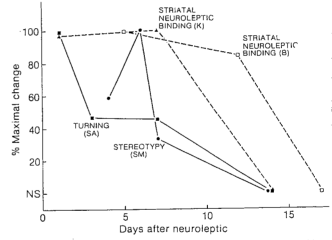


Fig. 4. The rate of decay of dopaminergic supersensitivity induced by neuroleptics. S.A.: Sayers et al. (1975). S.M.: Smith et al., (1976). K.: Kobayashi et al. (1978). B.: Burt et al. (1977)

the dyskinesia symptoms disappear within 2-3 months. The clinical statistics are further improved if the oldest patients (over fifty years of age) are not included; such patients are generally afflicted by a more persistent dyskinesia. According to Crane, patients over fifty have poorer prognosis in the reversal of tardive dyskinesia.

These observations suggest that tardive dyskinesia is considerably reversible, particularly in young patients. Similar observations on a different time scale are observed in the neuroleptic-treated rodents (Fig. 4). The result in Fig. 4 shows that the length of time needed for reversal of dopaminergic supersensitivity appears to

tensen rs and

ads to

ne**sia** n**s in** 

comed by son a cost in

correlate with the duration of neuroleptic administration, regardless of the type and dose of neuroleptic employed, and regardless of the type of supersensitive property monitored.

### Dose-Dependency in Long-Term Neuroleptic Treatment

The experiments with chronic neuroleptic showed little dose-dependency (Fig. 3, inset). It is probable that the effect could be dose-dependent at lower neuroleptic concentrations. Interestingly, according to Crane (1973), no consistent dose-dependency can be demonstrated for the incidence of tardive dyskinesia.

## Effects of Long-Term Neuroleptic Treatment on Dopamine Receptors (Table 1)

Soon after the development of the <sup>3</sup>H-haloperidol assay method for neuroleptic/dopamine receptors (Seeman et al., 1975; Burt et al., 1975), Muller and Seeman reported that these receptors increased after long-term neuroleptic treatment (Muller and Seeman, 1976; Burt et al., 1977). This has been recently more fully examined (Kobayashi et al., 1978; Muller and Seeman, 1977; Burt et al., 1977; Friedhoff et al., 1977). The maximum increase appears to be around 30% for the 3Hneuroleptic receptor (striatum) and about 65% for the <sup>3</sup>H-apomorphine sites. This increase is characteristic for all neuroleptics studied, except for clozapine in the study by Kobayashi et al. (1978). Clozapine has generally yielded conflicting results. Chronic treatment with this drug resulted in stimulated locomotion (Smith and Davis, 1976; Gianutsos and Moore, 1977) and stereotypy in the hands of Smith and Davis (1976) but not of Gnegy et al. (1977). Chronic clozapine pretreatment had no effect on dopamine turnover (von Stralendorff et al., 1976; Gianutsos and Moore, 1977) in different brain areas, but large decreases of dopamine turnover after long-term clozapine was observed in the striatum and the olfactory tubercle (Gianutsos and Moore, 1977). There was also no effect on the adenylate cyclase (Gnegy et al., 1977).

The increase in locomotor behaviour after repeated neuroleptics suggests that dopaminergic supersensitivity occurs in the mesolimbic areas. Locomotion has been shown to be associated primarily with the mesolimbic rather than striatal dopaminergic system (Jackson et al., 1975a, b; Costall and Naylor, 1975; Pijnenburg et al., 1976; Creese and Iversen, 1974). Jackson et al. (1975a) have shown that long-term penfluridol potentiates the locomotor response to dopamine administered to the nucleus accumbens but not to the striatum. These results are supported by our findings (Fig. 5) of dopaminergic supersensitivity in the mesolimbic areas. Klawans et al. (1977) did not show a

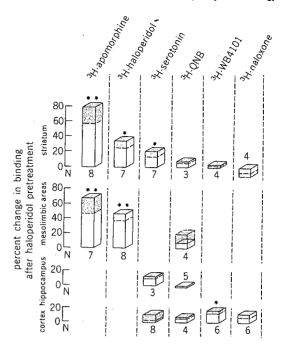


Fig. 5. Percentage changes in neurotransmitter receptors (rat brain regions) following long-term haloperidol treatment (10 mg/kg/day for at least 3 weeks). Height of the bars: mean percentage change Shaded portion: SEM. The number of completely independently as sayed membrane preparations (from 2 to 15 rats per preparation) used in each experiment (N) is indicated below each bar (\*P < 0.05.\*\* P < 0.02, using Wilcoxon's rank test). Long-term haloperidol treatment selectively increased the dopamine/neuroleptic binding without significant effects on acetylcholine receptors ( $^3$ H-QNB), alpha-adrenergic receptors ( $^3$ H-WB-4101), or opiate receptors ( $^3$ H-naloxone)

significant increase in <sup>3</sup>H-dopamine binding to the limbic areas, even though the results show a large trend towards such increase. Tolerance to catalepsy after repeated neuroleptic administration was demonstrated by some, but not others (Table 1).

### Effects of Long-Term Nonneuroleptic Drugs on Dopamine Transmission

From the drugs summarized in Table 2, opiates are the drugs closest to neuroleptics in terms of their effect on the dopaminergic system after repeated administration. Tolerance to the cataleptic effects as well as to dopamine turnover occurs with both opiates and neuroleptics (Gessa and Tagliamonte, 1975).

Reports of cross tolerance between neuroleptics and opiates have been published (Puri and Lal, 1974; Ezrin Waters and Seeman, 1977), even though the two studies do not agree whether the cross tolerance is one-way or two-way. Acutely, both opiates and neuroleptics induce catalepsy, even though there are differences in appearance of the animal, as well as different pathway, involved (Costall and Naylor, 1973); furthermore,

The second secon

Table 3. The effect of chronic administration of neuroleptics on non-dopaminergic neurotransmission

| % 4         | Drug/dose/day                    | $R_x$  | W                                     | Assay conditions  | References   |
|-------------|----------------------------------|--------|---------------------------------------|---|--|
| Noradrena   | ıline:                           | ······ | · · · · · · · · · · · · · · · · · · · |   | 1  |
| NS          | Chlorpromazine 5 mg/kg i.p.      | 7 d    | 3 d                                   | adenylate cyclase limbic  | Dolphin et al. (1977)                                |
| 60 %        | Chlorpromazine 5 mg/kg i.p.      | 7 d    | 3 d                                   | adenylate cyclase limbic;<br>adenylate cyclase limbic;<br>100 μΜ ΝΑ | Dolphin et al. (1977) Dolphin et al. (1977)          |
| NS          | Chlorpromazine 5 mg/kg i.p.      | 7 d    | 3 d                                   | clonidine-locomotion  | Dolphin et al. (1977)                                |
| 167%        | Haloperidol 3 mg/kg i.p          | 3 wks  | 4 d                                   | clonidine-locomotion  | Dustan and Jackson (1978)                            |
| 113%        | Haloperidol 10 mg/kg p.o.        | 3 wks  | 2 d                                   | cx 1 nM WB-4101 $\pm$ 1 $\mu$ M phenox.                             | Muller and Seeman (1977)                             |
| NS          | Haloperidol 10 mg/kg p.o.        | 3 wks  | 2 d                                   | st 1 nM WB-4101<br>± 1 μM phenox.                                   | Muller and Seeman (1977)                             |
| Serotonin:  |                                  |        |                                       |   |  |
| 100%        | Chlorpromazine 30 mg/kg i.p.     | 4 d    | 18 h                                  | locomotion with L-tryptophan  | Heal et al. (1976)                                   |
| NS          | Haloperidol 10 mg/kg i.p.        | 5 d    | 3 d                                   | locomotion with L-trypophan   | Heal et al. (1976)                                   |
| NS          | Spiroperidol 1 mg/kg i.p.        | 4 d    | 18 h                                  | locomotion with L-trypophan   | Heal et al. (1976)                                   |
| 120 %       | Haloperidol 10 mg/kg p.o.        | 3 wks  | 2 d                                   | st 3 nM 5-HT ± 100 nM   | Muller and Seeman (1977)                             |
| NS          | Haloperidol 10 mg/kg p.o.        | 3 wks  | 2 d                                   | hippo 3 nM 5-HT<br>± 100 nM 5-HT                                    | Muller and Seeman (1977)                             |
| NS          | Haloperidol 10 mg/kg p.o.        | 3 wks  | 2 d                                   | cx 3 nM 5-HT<br>± 100 nM 5-HT                                       | Muller and Seeman (1977)                             |
| Acetylcholi | ine:                             |        |                                       |   |  |
| NS          | Haloperidol 3 mg/kg i.p.         | 3 wks  | 4 d                                   | benztropine-locomotion  | Duston and Indian (1076                              |
| -17%        | Haloperidol 3 mg/kg in water     | 3 wks  | 4 d                                   | reserpine-blocked locomotion  | Dustan and Jackson (1976<br>Dustan and Jackson (1977 |
| 266%        | Haloperidol 3 mg/kg in water     | 2 wks  | 4 d                                   | atropine-locomotion   | Dustan and Jackson (1977                             |
| 58%         | Haloperidol<br>2.5-10 mg/kg i.p. | 24 d   | 3 - 5 d                               | pilocarp-blocked<br>locomotion                                      | Gianutsos and Lal (1976)                             |
| 73 %        | Haloperidol 2.5-10 mg/kg i.p.    | 3 wks  | 3-5 d                                 | dexetimide-locomotion   | Gianutsos and Lal (1976)                             |
| 1S          | Haloperidol 5 mg/kg i.p.         | 3 wks  | 14 d                                  | st 1 nM QNB<br>± 1 μM atrop.  | Kobayashi et al. (1978)                              |
| 12          | Haloperidol 10 mg/kg p.o.        | 3 wks  | 2 d                                   | st 1 nM QNB<br>± 100 nM scop.                                       | Muller and Seeman (1977)                             |
| -31 %       | Haloperidol 5 mg/kg i. p.        | 3 wks  | 14 d                                  | hippo 1 nM QNB<br>± 1 μM atrop.                                     | Kobayashi et al. (1978)                              |
| IS          | Haloperidol 5 mg/kg i.p.         | 3 wks  | 1 - 7 d                               | hippo 1 nM QNB<br>± 1 μM atrop.                                     | Kobayashi et al. (1978)                              |
| IS          | Haloperidol 10 mg/kg p.o.        | 3 wks  | 2 d                                   | hippo 1 nM QNB<br>± 0.1 μM scop.                                    | Muller and Seeman (1977)                             |
| IS          | Haloperidol 10 mg/kg p.o.        | 3 wks  | 2 d                                   | mli 1 nM QNB<br>± 0.1 μM scop.                                      | Muller and Seeman (1977)                             |
| IS          | Haloperidol 10 mg/kg p.o.        | 3 wks  | 2 d                                   | cx 1 nM QNB<br>± 0.1 µM scop.                                       | Muller and Seeman (1977)                             |
| ABA:        |                                  |        |                                       |   | 178  |
| IS          | Haloperidol 5 mg/kg i.p.         | 3 wks  | 1-14 d                                | st 10 nM GABA<br>± 0.5 nM GABA                                      | Kobayashi et al. (1978)                              |
| ;<br>1S     | Haloperidol 5 mg/kg i.p.         | 24 d   | 3-5 d                                 | hippo 10 nM GABA<br>± 0.5 µM GABA                                   | Kobayashi et al. (1978)                              |

there are biochemical differences (Kuschinsky and Hornykiewicz, 1972; Leysen et al., 1977) as well as pharmacological differences (Ezrin-Waters et al., 1976). Both groups of drugs block amphetamine-induced stereotypy (Sasame et al., 1972) as well as increase dopamine turnover (Ahtee and Kaariainen,

1973). Chronic neuroleptic treatment does not product any significant change in the striatal or cortical loxone binding (Fig. 5). The morphine pretreatment profile also differs from the neuroleptic profile in the morphine does not produce supersensitivity to apone phine (Table 2).

Long-ter stees '3H-ne mailive ad altered (Tal-Of the o ome simila limitation evidence of levity. Pher c-butaclar supersensiti.
The tole

neuroleptic exception ( on whether Moreover, comparable neuroleptic Chronic pictamine. dopaminer by stereoty et al., 1977 these ago dopamineour propo duces dor autorecept dopaminei preparatio

We hapomorph cally apon the <sup>3</sup>H-ha centration pretreatm predilectic pimozide al., 1976; the drop autorecep amine an (1977) and after chrotter met

Plects of rausmiss sveral s noradren neurolept (1977) re cyclase t

**postsy**naj

Long-term neuroleptic treatment consistently eleness <sup>3</sup>H-neuroleptic receptors (Table 1); dopamineensitive adenylate cyclase, however, is inconsistently lered (Table 1; Burkard and Bartholini, 1974).

Of the other drugs reviewed, the only drug showing one similarity to neuroleptics is ethanol. It causes imulation of locomotion (Table 2), but there is no vidence of ethanol-induced dopaminergic supersensityity. Phenobarbital, diazepam, promethazine, and butaclamol also do not produce dopaminergic persensitivity.

The tolerance in dopamine turnover (striatum) after reproleptics has been noted by several labs with one cception (Puri and Lal, 1974); there is no agreement on whether such tolerance occurs in the limbic areas. Moreover, the degree of tolerance in turnover is of comparable magnitude to the increase in <sup>3</sup>H-reproleptic receptors.

Chronic administration of dopamine agonists (ampletamine, L-Dopa or bromocryptine) induce apparent opaminergic 'behavioural-facilitation', as monitored by stereotypy (Klawans and Margolin, 1975; Klawans tal., 1977; Fuxe et al., 1973). Chronic treatment with hese agonists produces tolerance to their acute opamine-turnover-reducing effect. This agrees with our proposal that chronic agonist pretreatment produces dopaminergic subsensitivity of dopaminergic inforeceptors and thus produces apparent behavioural dopaminergic facilitation (Muller and Seeman, in preparation).

We have found that the binding of <sup>3</sup>Hpomorphine in the striatum is reduced in the chronially apomorphine- or amphetamine-treated rat, while the 3H-haloperidol sites (displaceable by a low concontration of pimozide) is not altered by the same pretreatment. Apomorphine is thought to have some medilection for presynaptic sites (Carlsson, 1975) and amozide prefers postsynaptic receptors (Gianutsos et 121976; Walters and Roth, 1976). We thus interpret drop in <sup>3</sup>H-apomorphine sites as a reduction in attoreceptors after repeated administration of amphetmine and apomorphine. The findings of Burt et al. (1977) and Friedhoff et al. (1977) did not detect changes ther chronic agonist pretreatment, possibly because ter methods did not distinguish between pre- and ictsynaptic binding.

### Meets of Long-Term Neuroleptics on Nondopaminergic

Styeral studies indicate that there may be a possible pradrenergic supersensitivity following repeated at oleptic administration (Table 3). Dolphin et al. (1917) report supersensitivity of the limbic adenylate class to 10 μM noradrenaline while the baseline

adenylate cyclase activity was unchanged by the treatment. Stimulation of locomotion with clonidine was more pronounced in the study of Dustan and Jackson (1976), but not in that of Dolphin et al. (1977). We have reported an increase in alpha-adrenergic receptors in the rat cortex but not in the striatum (Muller and Seeman, 1977; Fig. 5). Such a possible noradrenergic supersensitivity might be due to blockade of noradrenergic receptors by neuroleptics (U'Prichard et al., 1977; Andén et al., 1970; Keller et al., 1973).

Chronic neuroleptic treatment does not potentiate the locomotor response to L-tryptophan. Of the neuroleptics studied by Heal et al. (1976), chlorpromazine was the only one which caused apparent behavioural supersensitivity to L-tryptophan. In our binding studies, <sup>3</sup>H-serotonin binding was increased in the striatum but not in the cortex or hippocampus after repeated haloperidol treatment. Since we saw the same effect in the striatum of rats treated chronically with ethanol, we think that the neuroleptic-induced increase in the striatal binding might not be a specific effect (Muller and Seeman, 1977).

Table 3 shows no consistent changes in the choliner-gic or GABA sites after repeated haloperidol administration. The apparent cholinergic hyposensitivity of the cholinergic system reported by Gianutsos and Lal (1976) could be accounted for by an increase in tonic dopaminergic action, thus swinging the cholinergic-dopaminergic balance towards dopamine even if the sensitivity of the cholinergic system remained unchanged.

Acknowledgements. Supported by the Ontario Mental Health Foundation and the Medical Research Council of Canada.

#### References

Ahtee, L., Kaariainen, I.: The effect of narcotic analgesics on the homovanillic acid content of rat nucleus caudatus. Eur. J. Pharmacol. 22, 216-218 (1973)

Ahtee, L.: Catalepsy and stereotypies in the rats treated with methadone: relation to striatal dopamine. Eur. J. Pharmacol. 27, 221-230 (1974)

Anden, N.-E., Butcher, S. G., Corrodi, H., Fuxe, K., Ungerstedt, U.: Receptor activity and turnover of dopamine and noradrenaline after neuroleptics. Eur. J. Pharmacol. 11, 303-314 (1970)

Asper, H., Baggiolini, M., Burki, H. R., Lauener, H., Ruch, W., Stille, G.: Tolerance phenomena with neuroleptics. Catalepsy apomorphine stereotypies and striatal dopamine metabolism in the rat after single and repeated administration of loxapine and haloperidol. Eur. J. Pharmacol. 22, 287-294 (1973)

Bowers, M. B., Jr., Rozitis, A.: Regional differences in homovanillic acid concentrations after acute and chronic administration of antipsychotic drugs. J. Pharm. Pharmacol. 26, 743 – 745 (1974)

Burkard, W. P., Bartholini, G.: Changes in activation of adenylate cyclase and of dopamine turnover in rat striatum during prolonged haloperidol treatment. Experimentia 30, 685 (1974)

Burt, D. R., Creese, I., Snyder, S. H.: Antischizophrenic drugs: chronic treatment elevates dopamine receptor binding in brain. Science 196, 326-328 (1977)

- Burt, D. R., Enna, S. J., Creese, I., Snyder, S. H.: Dopamine receptor binding in the corpus striatum of mammalian brain. Proc. Natl. Acad. Sci. U.S.A. 72, 4655 4659 (1975)
- Carlsson, A.: Receptor-mediated control of dopamine metabolism. In: Pre- and postsynaptic receptors. E. Usdin and W. E. Bunney, eds., pp. 49-63. New York: Marcel Dekker 1975
- Christensen, A. V., Fjalland, B., Møller Nielsen, I.: On the supersensitivity of dopamine receptors induced by neuroleptics. Psychopharmacology 48, 1-6 (1976)
- Clouet, D. H., Iwatsubo, K.: Dopamine-sensitive adenylate cyclase of the caudate nucleus of rats treated with morphine. Life Sci. 17, 35-40 (1975)
- Costall, B., Naylor, R. J.: Neuroleptic and non-neuroleptic catalepsy. Arzneim. Forsch. 23, 674-683 (1973)
- Costall, B., Naylor, R. J.: The behavioural effects of dopamine applied intracerebrally to areas of mesolimbic system. Eur. J. Pharmacol. 32, 87-92 (1975)
- Costentin, J., Marcais, H., Protais, P., Baudry, M., Delabaume, S., Matres, M. P., Schwartz, J. C.: Rapid development of hypersensitivity of striatal dopamine receptors induced by alfamethylparatyrosine and its prevention by protein synthesis inhibitors. Life Sci. 21, 307 314 (1977)
- Cox, B., Ary, M., Lomax, P.: Changes in sensitivity to apomorphine during morphine dependence and withdrawal in rats. J. Pharmacol. Exp. Ther. 196, 637-641 (1975)
- Crane, G. E.: Persistent dyskinesia. Br. J. Psychiatry 122, 395-405 (1973)
- Creese, I., Iversen, S. D.: The role of forebrain dopamine systems in amphetamine induced stereotyped behaviour in the rat. Psychopharmacologia (Berl.) 39, 345-357 (1974)
- Dolphin, A., Sawaya, M. C. B., Jenner, P., Marsden, C. D.: Behavioural and biochemical effects of chronic reduction of cerebral noradrenaline receptor stimulation. Naunyn-Schmiedeberg's Arch. Pharmacol. 299, 163-173 (1977)
- Dustan, R., Jackson, D. M.: The demonstration of a change in adrenergic receptor sensitivity in the central nervous system of mice after withdrawal from long-term treatment with haloperidol. Psychopharmacology 38, 105-114 (1976)
- Dustan, R., Jackson, D. M.: The demonstration of a change in responsiveness of mice to physostigmine and atropine after withdrawal from long-term haloperidol pretreatment. J. Neural Transm. 40, 181-189 (1977)
- Engel, J., Liljequist, S.: The effect of long-term ethanol treatment on the sensitivity of the dopamine receptors in the nucleus accumbens. Psychopharmacology 49, 253-257 (1976)
- Ezrin-Waters, C., Muller, P., Seeman, P.: Catalepsy induced by morphine or haloperidol effects of apomorphine and anti-cholinergic drugs. Can. J. Physiol. Pharmacol. 54, 516-519 (1976)
- Ezrin-Waters, C., Seeman, P.: Tolerance to haloperidol catalepsy. Eur. J. Pharmacol. 41, 321-327 (1977)
- Ezrin-Waters, C., Seeman, P.: Haloperidol-induced tolerance to morphine catalepsy. Life Sci. 21, 419 422 (1977)
- Friedhoff, A. J., Bonnet, K., Rosengarten, H.: Reversal of two manifestations of dopamine receptor supersensitivity by administration of L-DOPA. Res. Commun. Chem. Pathol. Pharmacol. 16, 411-423 (1977)
- Fukui, K., Takagi, H.: Effect of morphine on cerebral contents of metabolites of dopamine in the normal and tolerant mice: its possible relation to analgesic action. Br. J. Pharmacol. 44, 45-51 (1972)
- Fuxe, K.: Tools in the treatment of Parkinson's disease: studies on new types of dopamine receptor stimulating agents. In: Progress in the treatment of Parkinsonism. Advances in neurology, Vol. 3, D. B. Calne, ed., pp. 273-279. New York: Raven 1973
- Gessa, G. L., Tagliamonte, A.: Effect of methadone and dextromoramide on dopamine metabolism: comparison with halo-

- peridol and amphetamine. Neuropharmacology 14, 913-92 (1975)
- Gianutsos, G., Drawbaugh, R. B., Hynes, M. D., Lal, H., Behavioural evidence for dopaminergic supersensitivity alter chronic haloperidol. Life Sci. 14, 887—898 (1974)
- Gianutsos, G., Lal, H.: Alteration in the action of cholinergic and anticholinergic drugs after chronic haloperidol: indirect evidence for cholinergic hyposensitivity. Life Sci. 18, 515-520 (1976)
- Gianutsos, G., Thornburg, J. E., Moore, K. E.: Differential actions of dopamine agonists and antagonists on γ-butyrolactone induced increase in mouse brain dopamine. Psychopharmacology 50, 225 229 (1976)
- Gianutsos, G., Moore, K. E.: Dopaminergic supersensitivity in striatum and olfactory tubercle following chronic administration of haloperidol or clozapine. Life Sci. 20, 1585-1592 (1977)
- Gnegy, M., Uzunov, P., Costa, E.: Participation of an endogenous Ca<sup>++</sup>-binding protein activator in the development of drug induced supersensitivity of striatal dopamine receptors. J. Pharmacol. Exp. Ther. 202, 558-564 (1977)
- Griffiths, P. J., Littleton, J. M., Ortiz, A.: Changes in monoamine concentrations in mouse brain associated with ethanol dependence and withdrawal. Br. J. Pharmacol. 50, 489 494 (1974)
- György, L., Pfeifer, K. A., Hajtman, B.: Modification of certain central nervous effects of haloperidol during long-term treatment in the mouse and rat. Psychopharmacologia (Berl.) 16, 223-233 (1969)
- Heal, D. J., Green, A. R., Boullin, D. J., Graham-Smith, D. G., Single and repeated administration of neuroleptic drugs to rats effect on striatal dopamine-sensitive adenylate cyclase and locomotor activity produced by transleypromine and L tryptophan or L-dopa. Psychopharmacology 49, 287-300 (1976)
- Hicks, J., Strong, R., Smith, R. C., Samorajski, T.: Behavioural supersensitivity to bromocriptine in mice. Society for Neuroscience abstract, III, p. 441, No. 1406 (1977)
- Hoffman, P. L., Tabakoff, B.: Alterations in dopamine receptor sensitivity by chronic ethanol treatment. Nature 268, 551-552 (1977)
- Hornykiewicz, O.: The mechanisms of action of L-DOPA in Parkinson's disease. Life Sci. 15, 1249-1259 (1974)
- Iwatsubo, K., Clouet, D. H.: Dopamine-sensitive adenylate cyclase of the caudate nucleus of rats treated with morphine of haloperidol. Biochem. Pharmacol. 24, 1499-1503 (1975)
- Jackson, D. M., Andén, N.-E., Engel, J., Liljequist, S.: The effect of long-term penfluridol treatment on the sensitivity of the dopamine receptors in the nucleus accumbens and the corpus striatum. Psychopharmacologia (Berl.) 45, 151-155 (1975)
- Jackson, D. M., Andén, N.-E., Dahlström, A.: A functional effect of dopamine in the nucleus accumbens and in some other dopamine-rich parts of the rat brain. Psychopharmacologia (Berl.) 45, 139-149 (1975)
- Keller, H. H., Bartholini, G., Pletscher, A.: Increase of 3-methoxy hydroxyphenylethylene glycol in rat brain by neuroleptic drugs. Eur. J. Pharmacol. 23, 183-186 (1973)
- Klawans, H. L., Crossett, P., Dana, N.: Effect of chronic amples amine exposure on stereotyped behaviour: implications for pathogenesis of L-Dopa-induced dyskinesias. In: Dopaminergic mechanisms. Advances in neurology, Vol. 9, D. Calne, T. N. Chase and A. Barbeau, eds., pp. 105-112. New York: Raven 1975
- Klawans, H. L., Margolin, D. I.: Amphetamine-induced dopamine gic hypersensitivity in guinea-pigs. Arch. Gen. Psychiatry 32 725-732 (1975)
- Klawans, H. L., Goetz, C., Nausieda, P. A., Weiner, W. J. Levodopa-induced dopamine receptor hypersensitivity. Ann Neurology 2, 125-129 (1977)

- Obayashi, R. M., Fields, J. Z., Hruska, R. E., Beaumont, K., Yamamura, H. I.: Brain neurotransmitter receptors and chronic antipsychotic drug treatment: a model for tardive dyskinesia. In: Animal models in psychiatry. E. Usdin, ed., pp. 405-409. New York: Pergamon 1978
- Kushinsky, K., Hornykiewicz, O.: Morphine catalepsy in rat: Felation to striatal dopamine metabolism. Eur. J. Pharmacol. 19, \$19-122 (1972)
- Nuchinsky, K.: Dopamine receptor sensitivity after repeated morphine administrations to rats. Life Sci. 17, 43 48 (1975)
- Lemer, P., Nosé, P.: Haloperidol: effect of long-term treatment on rat striatal dopamine synthesis and turnover. Science 197, 181 183 (1977)
- Lycen, J., Tollenaere, J. P., Koch, M. H. J., Laduron, P.: Differentiation of opiate and neuroleptic receptor binding in rat brain. Eur. J. Pharmacol. 43, 253 267 (1977)
- driden, C. D., Tarsy, D., Baldessarini, R. J.: Spontaneous and drug-induced movement disorders in psychotic patients. In:
  Psychiatric aspects of neurologic disease. D. F. Benson and D.
  Plumer, eds., pp. 219 265. New York: Grune & Stratton 1975
- Artres, M. P., Costentin, J., Baudry, M., Marcais, H., Protais, P., Schwartz, J. C.: Long-term changes in the sensitivity of pre- and postsynaptic dopamine receptors in mouse striatum evidenced by behavioural and biochemical studies. Brain Res. 136, 319—337 (1977)
- Tali, Z., Singhal, R. L., Hrdina, P. D., Ling, G. M.: Changes in brain cyclic AMP metabolism and acetylcholine and dopamine during narcotic dependence and withdrawal. Life Sci. 16, 1889—1894 (1975)
- Mer Nielsen, I., Fjalland, B., Pedersen, V., Nymark, M.: Pharmacology of neuroleptics upon repeated administration. Psychopharmacologia (Berl.) 34, 95-104 (1974)
- dler, P., Seeman, P.: Increased specific neuroleptic binding after chronic haloperidol in rats. Soc. Neurosci. Abstr. 2, 874 (1976)
- Milet, P., Seeman, P.: Brain neurotransmitter receptors after longterm haloperidol: dopamine, acetylcholine, serotonin, αnoradrenergic and naloxone receptors. Life Sci. 21, 1751 – 1758 (1977)
- Feenburg, A. J. J., Honing, W. M. M., Van Der Heyden, J. A. M., Yan Rossum, J. M.: Effects of chemical stimulation of the misolimbic dopamine system upon locomotor activity. Eur. J. Pharmacol. 35, 45-58 (1976)
- id. S. K., Lal, H.: Tolerance to the behavioural and neurochemical affects of haloperidol and morphine in rats chronically treated with morphine or haloperidol. Naunyn-Schmiedeberg's Arch. Pharmacol. 282, 155—170 (1974)
- Tui, S. K., Volicer, L., Cochin, J.: Changes in the striatal adenylate colored activity following acute and chronic morphine treatment and during withdrawal. J. Neurochem. 27, 1551—1554 (1976)
- tkin, F., Rifkin, A., Gochfeld, L., Klein, D. F.: Tardive dykinesia: are first signs reversible? Am. J. Psychiatry 134, 84—17 (1977)
- md, W. H., Gerald, M. C.: The effect of chronic administration and withdrawal of (+)-amphetamine on seizure threshold and and endogenous catecholamine concentrations and their rates of hosynthesis in mice. Psychopharmacology 51, 175-179 (1977)
- activity following reserpine and chronic chlorpromazine administration in rats. Life Sci. 17, 563 – 568 (1975)

- Sasame, H. A., Perez-Cruet, J.: Evidence that methadone blocks dopamine receptors in the brain. J. Neurochem. 19, 1953-1957 (1972)
- Sayers, A. C., Bürki, H. R., Ruch, W. Asper, H.: Neuroleptic-induced hypersensitivity of striatal dopamine receptors in the rat as a model of tardive dyskinesias. Effects of clozapine, haloperidol, loxapine and chlorpromazine. Psychopharmacologia (Berl.) 41, 97-104 (1975)
- Scatton, B., Glowinski, J., Julou, L.: Dopamine metabolism in the mesolimbic and mesocortical dopaminergic systems after single or repeated administrations of the neuroleptics. Brain Res. 109, 184-189 (1976)
- Seeber, U., Kuschinsky, K.: Dopamine-sensitive adenylate cyclase in homogenates of rat striata during ethanol and barbiturate withdrawal. Arch. Toxicol. (Berl.) 35, 247-253 (1976)
- Seeman, P., Chau-Wong, M., Tedesco, J., Wong, K.: Brain receptors for antipsychotic drugs and dopamine: direct binding assays. Proc. Natl. Acad. Sci. U.S.A. 72, 4376-4380 (1975)
- Smith, R. C., Davis, J. M.: Behavioural evidence for supersensitivity after chronic administration of haloperidol, clozapine and thioridazine. Life Sci. 19, 725-732 (1976)
- Tang, L. C., Cotzias, G. C.: L-3,4-Dihydroxyphenylalanine-induced hypersensitivity simulating features of denervation. Proc. Natl. Acad. Sci. U.S.A. 74, 2126-2129 (1977)
- Tarsy, D., Baldessarini, R. J.: Behavioural supersensitivity to apomorphine following chronic treatment with drugs which interfere with synaptic function of catecholamines. Neuropharmacology 13, 927-940 (1974)
- Tarsy, D., Baldessarini, R. J.: The pathophysiologic basis of tardive dyskinesia. Biol. Psychiatry 12, 431-450 (1977)
- U'Prichard, D. C., Greenberg, D. A., Snyder, S. H.: Binding characteristics of radiolabeled agonist and antagonist at central nervous system alpha noradrenergic receptors. Mol. Pharmacol. 13, 454-473 (1977)
- Von Stralendorff, V. B., Ackenheil, M., Zimmermann, J.: Akute und chronische Wirkung von Carpipramin, Clozapin; Haloperidol und Sulpirid auf den Stoffwechsel biogener Amine im Rattengehirn. Arzneim. Forsch. 26, 1096-1098 (1976)
- Von Voigtlander, P. F., Losey, E. G., Triezenberg, H. J.: Increased sensitivity to dopaminergic agents after chronic neuroleptic treatment. J. Pharmacol. Exp. Ther. 193, 88-94 (1975)
- Walters, J. R., Roth, R. H.: Dopamine neurons: an in vivo system for measuring drug interactions with presynaptic receptors. Naunyn-Schmiedeberg's Arch. Pharmacol. 296, 5-14 (1976)
- Worms, P., Scatton, B.: Tolerance to stereotyped behaviour and to decrease in striatal homovanillic acid levels after repeated treatment with apomorphine dipivaloyl ester. Eur. J. Pharmacol. 45, 395-396 (1977)
- Yarborough, G.: Supersensitivity of caudate neurones after repeated administration of haloperidol. Eur. J. Pharmacol. 31, 367-369 (1975)
- Zarcone, V., Barchas, J., Hoddes, E., Montplaisir, J., Sack, R., Wilson, R.: Experimental ethanol ingestion: sleep variables and metabolites of dopamine and serotonin in the cerebrospinal fluid. In: Alcohol intoxication and withdrawal. Vol. 2, M. M. Gross, ed., pp. 431-451. New York: Plenum 1975

Received February 24, 1978