Schizophrenia is a major therapeutic challenge of modern medicine, and one of the last frontiers of brain research. The illness is defined by delusions, hallucinations, disorganized behavior, and cognitive difficulties such as memory loss. It occurs in ~1% of the world population and usually first appears in early adulthood. Although antipsychotic medications have dramatically improved the lives of patients with schizophrenia, the causes of the illness remain unknown.

Of the many contemporary theories of schizophrenia, the most enduring has been the dopamine hypothesis. As originally put by Van Rossum in 1967 (ref. 1, p. 321), “When the hypothesis of dopamine blockade by neuroleptic agents can be further substantiated, it may have fargone consequences for the pathophysiology of schizophrenia. Overstimulation of dopamine receptors could be part of the aetiology…” Indeed, this speculative sentence by Van Rossum foreshadows the title of the important work by Abi-Dargham et al. (2) in this issue of PNAS: “Increased baseline occupancy of D₂ receptors by dopamine in schizophrenia.”

The discovery of the antipsychotic/dopamine receptor (3, 4), now commonly known as the dopamine D₂ receptor, led to repeated confirmation that it is the primary site of action for all antipsychotics (3–5), including clozapine and quetiapine (6). All these drugs have different potencies at the receptor. The potency depends on the drug’s dissociation constant at D₂, which, in turn, relates to the rate of release of the drug from the D₂ receptor. For example, the dopamine D₂ receptor releases clozapine and quetiapine more rapidly than it does any of the other antipsychotic drugs (7, 8).

Given the tight correlation between the clinical potency and the D₂-blocking action of the antipsychotic medications, dopamine overactivity could be the common denominator in the psychotic element of schizophrenia. This possibility has been actively investigated. Dopamine overactivity can be presynaptic (an excess of dopamine release from dopamine nerve terminals) or postsynaptic (an increase in the density of D₂ receptors or an increase in postreceptor action). The innovative report by Abi-Dargham et al. (2) sheds light on both pre- and postsynaptic aspects by using an indirect method to measure the levels of endogenous dopamine in patients and controls.

Although numerous postmortem studies have consistently revealed D₂ receptors to be elevated in the striata of patients with schizophrenia (9), the majority of the postmortem tissues examined have come from patients who have been treated with antipsychotics, raising the probability that the drugs themselves contributed to the elevation of D₂ receptors. To measure the density of D₂ receptors in never-medicated patients with schizophrenia, D₂-selective ligands have been used with in vivo brain imaging methods (10–12). The results have not been consistent. Data with [11C]methylspiperone show elevated D₂ receptors in schizophrenia (ref. 10, but see also ref. 12), whereas data with [11C]raclopride do not show such elevation (ref. 11 and discussed later in this paper). One major reason for this discrepancy is the quantitatively different effects of endogenous dopamine on [11C]methylspiperone and [11C]raclopride (see references in ref. 7).

Hence, one way to resolve this discrepancy is to measure D₂ receptors after partial depletion of endogenous dopamine in patients. The work of Abi-Dargham et al. (2) provides this resolution. Fig. 1 summarizes the principle used by Abi-Dargham et al. Fig. 1 (Top) illustrates that the radiobenzamide (5)-(−)-3-[123]Jido-2-hydroxy-6-methoxy-N-{(1-ethyl-2-pyrrolidinyl)methyl}benzamide ([123]IBZM) binds to the same number of D₂ receptors in control and schizophrenia individuals. That is, the “binding potential” was the same in both sets of subjects. However, after partial depletion of endogenous dopamine by oral ingestion of α-methylparatyrosine over 2 days, the binding of [123]IBZM rose by 19% in schizophrenia but only by 9% in control subjects (Fig. 1, Bottom). In fact, when Abi-Dargham et al. examined the number of D₂ receptors after partially removing the obscuring effect of endogenous dopamine, the D₂ receptors were significantly elevated in schizophrenia patients as compared with control subjects. When the authors examined the data by subgroups, the results of increased receptors reached significance for previously medicated patients, but exhibited only a trend for patients who had never been medicated with antipsychotic drugs. Despite this lack of statistical significance in this latter group of patients, the empirical findings of Abi-Dargham et al. indicate that an increase in dopamine D₂ receptors must occur, because it is not possible for patients to show a greater increase yet not have a higher number of D₂ receptors. Thus, the paper by Abi-Dargham et al. provides support for both an increase in the level of dopamine as well as an increase in the number of D₂ receptors in schizophrenia, compared to control subjects.

Schizophrenia, as compared with control subjects, also is associated with an increased releasability of dopamine (13, 14). A high release rate of dopamine reduces the binding of radiobenzamides to tissues (15, 16), but enhances the binding of radiospiperone (17, 18). Competition with endogenous dopamine, as well as dopamine-induced internalization of the D₂ receptors, may account for the lessened binding of radiobenzamides to the tissue (13, 14), because the benzamides are generally water-soluble and have less ready access to vesicle-associated receptors. Radiospiperone compounds, by contrast, are highly lipid-soluble and readily permeate cell membranes to reach internalized receptors.

In addition to the two schizophrenia-associated factors of increased D₂ receptors and increased dopamine release, there is a third factor. Dopamine D₂ receptors exist in monomer, dimer, and oligomeric forms (19). The D₂ monomer, but not the D₂ dimer, is selectively labeled by a photolabel of radiospiperone (19). This finding is in contrast to a benzamide photolabel (for nemonapride), which readily binds to both monomers and dimers of D₂ (19). This important distinction between benzamides and butyrophoenenes may
control individual has three D₂ receptors, two in the dimer form and one in the monomer form. It is proposed that in schizophrenia, under the influence of increased release of endogenous dopamine, all three exist in the monomer form. Thus, radioligand binding would show no difference, but the binding of radioligand would be higher in the schizophrenic brain (as compared to controls) because of an increased number of monomers.

The dopamine hypothesis has been much criticized. For instance, although therapeutic doses of most antipsychotics occupy 60% to 80% of the D₂ receptors in patients, clozapine and quetiapine have been apparent exceptions, exhibiting clinical efficacy with only 10% to 45% occupation of D₂ receptors (see references in ref. 7). It therefore has been suggested that the dopamine hypothesis of schizophrenia be extended into a serotonin-dopamine hypothesis. However, recent work on imaging both D₂ and serotonin-2 receptors in patients taking antipsychotics fails to find evidence for a contribution from the occupation of serotonin receptors (20). For example, the threshold for clinical antipsychotic action remains at 65% occupation of D₂ receptors in first-episode patients, whether one uses haloperidol, which has no serotonin-receptor blocking action, or risperidone or olanzapine, which block all serotonin-2 receptors but at doses far below those needed for clinical efficacy. Similarly, the threshold for extrapyramidal signs, which is ~80% D₂ occupancy, remains unaltered despite the presence of 100% block of serotonin-2 receptors for risperidone or olanzapine. It should also be noted that therapeutic doses of clozapine and quetiapine transiently occupy high levels of D₂ receptors in patients, but the effect lasts for only the first few hours (6). Thus, the D₂-occupying properties of clozapine and quetiapine are remarkable only for their short duration of action; they otherwise support the dopamine hypothesis of schizophrenia, as originally outlined by Van Rossum (1).

There is more to schizophrenia than psychosis. The psychological abnormalities and cognitive difficulties in schizophrenia precede and outlive the psychosis. The hypothesis of dopamine dysregulation is the best explanation for the psychotic episode in schizophrenia; the pathophysiology of other psychological and cognitive abnormalities in schizophrenia remains unclear. A combination of susceptibility genes (21) and other factors contributes to schizophrenia, and the net result dysregulates the dopamine neurotransmission system, leading to high release of dopamine, more D₂ receptors, and an apparent predominance of monomer forms of D₂. This dopamine dysregulation leads to the psychotic episode. Further research needs to uncover underlying mechanisms that predispose the brain to the dysregulation of the dopamine system (22). Until then, the dopamine hypothesis remains the main path to the origin and treatment of clinical signs and symptoms of psychosis in schizophrenia.

Fig. 1. Method and findings of Abi-Dargham et al. (2) to reveal an increased occupancy of dopamine D₂ receptors in schizophrenia. (Top) The number of dopamine D₂ receptors, measured by the \([123I]\)IBZM binding potential (green triangles with I), were the same in the brain striata of control and schizophrenia subjects. The levels of synaptic dopamine (pink triangles with D), which is higher in schizophrenia subjects than in controls, normally occupies most of the D₂ receptors, masking the difference between control and schizophrenia individuals. (Bottom) After partial depletion of endogenous brain dopamine by oral ingestion of \(\alpha\)-methylparatyrosine over 2 days, the binding of \([123I]\)IBZM rose in both the control and schizophrenia subjects, but that for the patients rose significantly higher.


