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The Pipeline and Future of Drug Development in Schizophrenia

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Abstract

While the current antipsychotic medications have profoundly impacted the treatment of schizophrenia over the past 50 years, the newer atypical antipsychotics have not fulfilled initial expectations, and enormous challenges remain in long-term treatment of this debilitating disease. In particular, improved treatment of the negative symptoms and cognitive dysfunction in schizophrenia which greatly impact overall morbidity is needed. In this review we will briefly discuss the current pipeline of drugs for schizophrenia, outlining many of the strategies and targets currently under investigation for the development of new schizophrenia drugs. Many of these compounds have great potential as augmenting agents in the treatment of negative symptoms of cognition. In addition, we will highlight the importance of developing new paradigms for drug discovery in schizophrenia and call for an increased role of academic scientists in discovering and validating novel drug targets. Indeed, recent breakthroughs in genetic studies of schizophrenia are allowing for the development of hypothesis-driven approaches for discovering possible disease-modifying drugs for schizophrenia. Thus, this is an exciting and pivotal time for the development of truly novels approaches to drug development and treatment of complex disorders like schizophrenia.

Introduction

Since the discovery of the antipsychotic effect of chlorpromazine more than 50 years ago¹, the number of antipsychotic medications available for the treatment of schizophrenia has tremendously increased. With the development of the first generation antipsychotics, or typical antipsychotics, it was for the first time possible to treat the "positive" symptoms of schizophrenia, such as delusions and hallucinations, leading to the deinstitutionalization of the world's mentally ill. The typical antipsychotic drugs are generally not thought to be effective at treating the "negative" symptoms, such as anhedonia and lack or motivation, and cognitive dysfunction of schizophrenia and have a high burden of extrapyramidal side effects (EPS)².

The reintroduction of clozapine in the United States in 1989, issued in the current era of atypical or second generation antipsychotics. Atypical antipsychotic drugs are differentiated from typical antipsychotic drugs by virtue of a relative lack of EPS and serum prolactin elevation as compared with typical antipsychotic drugs. Clozapine itself has become the 'gold standard' antipsychotic medication because of its absence of debilitating extrapyramidal side-effects and its demonstrated clinical superiority in treatment-resistant schizophrenia³ and suicidality⁴. Whether clozapine has any significant beneficial effect on negative symptoms and cognition is unclear^{2,5}. Clozapine, however, is associated with its own set of serious side-effects including weight gain, diabetes and an increased risk of seizures and agranulocytosis⁶.

The documented superiority of clozapine^{3,7} over other antipsychotic drugs has led to an intense effort over the past 15-20 years to develop clozapine-like atypical antipsychotics that are safer and better tolerated than clozapine. As such, multiple atypical and pseudo-atypical antipsychotic drugs, including risperidone, olanzapine, quetiapine and ziprasidone have been developed. Expectations that these agents comprised a breakthrough in the treatment of schizophrenia, especially with regards to improvements in negative symptoms and cognition were initially high⁸. These expectations, however, have not been realized^{9,10}. While there is evidence that most of the new medications offer, at best, modest advantages over the typical antipsychotic drugs with regard to improvement in negative symptoms, cognitive impairment and functional capacity, the improvements are not consistent among studies^{2,11,12}. In addition, the atypical antipsychotics carry their own substantial side effect burden, specifically weight gain and the metabolic syndrome^{13,14}.

While the introduction of antipsychotic medications has had a profound effect on the treatment of schizophrenia over the past 50 years, and the atypical antipsychotic drugs have provided a larger and more diverse armamentarium of treatment options, the advances that have been made since the discovery of the antipsychotic properties of chlorpromazine have been small and incremental. Thus, enormous challenges remain in long-term treatment of this debilitating disease and continuing with the current paradigms of drug discovery is unlikely to produce significant advances^{15,16}. It is therefore important to continue pursue diverse molecular targets for discovering new

antipsychotic compounds and to devise novel paradigms for drug discovery in schizophrenia.

A fundamental barrier to the discovery and development of novel treatments for schizophrenia remains that our level of understanding of the biological processes involved in schizophrenia is not sufficient to predict the therapeutic value of novel drug targets¹⁶. Thus, unvalidated targets are frequently left unpursued by the pharmaceutical industry and, frequently, companies have focused on alterations of existing medications (i.e. separating enantiomers or marketing active metabolites; e.g. 9-OH-risperidone or paliperidone)¹⁷, finding additional compounds that hit known and validated targets ("me too" drugs; e.g. ORG-5222 or asenapine)¹⁸, and on gaining approval on new indications for already marketed drugs¹⁹ (e.g. clozapine for suicide in schizophrenia⁴). These methods, however, cannot continue indefinitely as the number of such possibilities is limited and thus it is critical to find new approaches to drug development. Interestingly, many of the atypicals will soon be going off patent, beginning with the launch of generic risperidone in 2007. Thus, there is significant interest and urgency within the pharmaceutical industry and among schizophrenia basic scientists and clinicians in developing safer and more-effective treatments for schizophrenia.

In this review we will briefly discuss the current pipeline of drugs for schizophrenia (Table 1) and will outline many of the strategies and targets currently under investigation for the development of new schizophrenia drugs. In addition, we will highlight the importance of developing new paradigms for drug discovery in

schizophrenia and call for an increased role of academic scientists in discovering and validating novel drug targets^{15,16}.

Symptom Domains in Schizophrenia

It has been proposed that new therapeutics in schizophrenia should target narrower ranges of symptoms rather than to try to develop the perfect "monotherapy" for a complex disorder²⁰. This proposal is grounded in the complexity of schizophrenia which is characterized by severe and variable symptoms in a number of symptom domains, including positive symptoms such as hallucinations, delusions, and disorganized thought, negative symptoms of cognitive impairments in attention, working memory, and a variety of executive functions. All of the currently approved drugs for the treatment of schizophrenia, however, were developed and are most efficacious at treating the positive symptoms of the disease while the negative symptoms and cognitive impairments actually contribute disproportionately more to the long-term disability in patients with schizophrenia²¹.

It is clear that patients who exhibit significant negative symptoms have particularly poor function capacity and quality of life^{22,23} and while there was optimism that the atypicals comprised a breakthrough in the treatment of negative symptoms⁸, this prospect has not been realized to a clinically significant degree^{9,10}. Despite the

limitations of current medications and the morbidity associated with negative symptoms, no drug has received Food and Drug Administration (FDA) approval for an indication of negative symptoms. As such, the National Institutes of Mental Health (NIMH) has recently released a consensus statement on the negative symptoms of schizophrenia²⁴ highlighting that negative symptoms represent a distinct and clinically important entity that should be a focus of future drug development efforts.

In addition to negative symptoms, schizophrenia is also characterized by significant cognitive impairments. For example, patients with schizophrenia have been documented to have problems with attention, working memory and learning and a variety of executive-level functions including abstract thinking and problem solving^{25,26}. Indeed, a meta-analysis of cognitive deficits suggested that indices of cognitive deficits are much better predictors of functional outcome than indices from any other symptom domain²⁷. However, these cognitive deficits have been relatively unimproved by currently approved antipsychotic drugs, though some evidence exists for the superiority of atypicals such as olanzapine and risperidone over typicals^{5,28,29}. Due to the remaining need for improved treatment of the cognitive impairments in schizophrenia the National Institutes of Mental Health (NIMH) has begun a joint academic and industry initiative termed MATRICS (Measurement and Treatment Research to Improve Cognition in Schizophrenia) to facilitate the development of better treatments targeted at cognition³⁰.

Thus, while psychopharmacologic research in schizophrenia aims for the development of new antipsychotic drugs with a more rapid onset of action, lower risk of side effects and improved efficacy in the domains of negative and cognitive symptoms it is unlikely that a single drug will have the desired effect across all of these domains. Optimal treatment of schizophrenia in the near future will likely rely on polypharmacy with individualized treatment aimed at the multidimensional nature of this disorder.

Current Antipsychotics and Drugs in Phase III Clinical Trials

There are a number of theories regarding the mechanism of action of antipsychotic drugs^{16,31,32}, though the precise mechanism remains incompletely understood. Briefly, it is Important that all of the currently approved antipsychotic drugs have at least some affinity for the dopamine D₂ receptor and for the typicals there is a strong correlation between the therapeutic doses and their binding affinity for D₂ receptors^{33,34}. In addition, positron emission tomography (PET) studies have demonstrated that antipsychotic effects are associated with a striatal D₂ receptor occupancy of 65-70%^{35,36} with occupancy levels greater than 80% associated with increased risk of EPS³⁶. The basis of the "atypicality" of newer medications, likewise is incompletely understood, though a primary theory is the serotonin-dopamine antagonism theory³⁷ which posits that a higher ratio of serotonin 5-HT_{2A} receptor affinity to dopamine D₂ receptor affinity explains the enhanced efficacy and reduced EPS burden seen with the atypicals. This hypothesis is consistent with the atypical features

of risperidone, olanzapine, quetiapine, ziprasidone and the recently approved paliperidone (9-OH-risperidone). A third class of antipsychotics are the dopamine partial agonists, with aripiprazole being the only one currently approved for clinical use³². It is thought that the relative lack of EPS seen with clinical use of aripiprazole is due to its functional selectivity at D₂ receptors protecting against excessive blockade of the D₂ system^{32,38,39}. Thus, while the mechanism of action of currently available antipsychotics is not fully known, D₂ receptor occupancy (either by antagonism or functionally-selective agonism) is important for the treatment of the positive symptoms of schizophrenia with some modulation of this D₂ blockade, likely increased dopamine transmission in the cortex and hippocampus, being important for both why SGAs and aripiprazole have a reduced EPS burden and somewhat higher efficacy at treating negative symptoms and cognitive dysfunction⁴⁰.

The drugs in the pipeline for the treatment of schizophrenia that are currently in Phase III clinical trials appear to have the same mechanism of action of already available agents. Asenapine (formerly known as ORG-5222; Organon/Pfizer; now discontinued from clinical development) and iloperidone (Titan Pharmaceuticals) are antagonists at D₂ and 5-HT_{2A} and many other receptors (Table 2) and bifeprunox (Lundbeck/Solvay) is a D₂ partial agonist. An NDA has been submitted for bifeprunox for treatment of schizophrenia (<u>http://www.google.com/search?sourceid=navclient&ie=UTF-8&rls=DMUS,DMUS:2006-43,DMUS:en&q=bifeprunox+fda</u>). A novel medication bexarotene—a retinoid-X-receptor activator is currently listed as being in Phase III clinical trials as an add-on medication

for schizophrenia (<u>http://clinicaltrials.gov/ct/show/NCT00141947?order=6</u>) As such, with the possible exception of bexarotene, all compounds currently in Phase III clinical trials represent "me-too" drugs that are not significantly different from currently available medications, though there is some clinical benefit to having additional drugs available as individuals may have differential responses to medications and have varying tolerance to side effects. As these drugs will not represent significant advances in the treatment of schizophrenia, this review will focus primarily on compounds at earlier stages of development.

The current "gold standard" antipsychotic, clozapine, interestingly has relatively weak affinity for the D₂ dopamine receptors but has moderate to high affinity and antagonist/inverse agonist activity for many other neurotransmitter receptors, including other dopamine receptors (D₁, D₃, D₄), various serotonin receptors (5-HT_{2A}, 5-HT_{2C}, 5-HT₆, 5-HT₇), muscarinic acetylcholine receptors (M₁, M₂, M₃, M₄, M₅), and adrenergic receptors (α_1 , α_2)³¹. Additionally, clozapine's active metabolite N-desmethyl-clozapine, is a potent partial agonist at dopamine⁴¹ and muscarinic^{42,43} receptors. This extremely complex pharmacological profile is thought to underlie both clozapine's superior clinical efficacy and its spectrum of serious side effects³. As such, much effort in antipsychotic drug development over the past two decades has been to create clozapine-like drugs that bind to fewer targets and thus reduce the side effect burden by targeting only the appropriate receptors. As will be reviewed below, attempts thus far to target clozapine's 'magic receptor', however, have been largely been unsuccessful. Indeed, it appears that the paradigm of 'one-disease one-target' that became the dominant approach in the

pharmaceutical industry with the advent of molecular biological techniques, while ideal from a scientific and practical perspective, may not be suitable for complex psychiatric diseases such as schizophrenia. In recent years, a number of authors have proposed that designing selectively non-selective drugs that interact with several molecular targets (coined 'magic shotguns')³¹ will lead to more effective medications for a variety of complex disorders^{31,44,45}.

We will briefly review the individual molecular targets that may have a role in these 'magic shotguns' or in a polypharmacy approach targeting the various clinical symptom domains of schizophrenia **[Figure 3 – Domains]**. Where available, references to key review articles are provided. In addition, we will highlight how selective compounds have generally been ineffective as monotherapy for schizophrenia. Indeed, many of the drugs in Phase I, Phase II and preclinical development for the treatment of schizophrenia represent a shift from targeting D₂ and 5-HT_{2A} receptors to targeting to other monoaminergic receptors and other neurotransmitter receptors, however, though results in small clinical trials have generally been less than encouraging.

Additional Dopaminergic Approaches

In addition to the key role of dopamine D₂ receptors in antipsychotic function, compounds selective for other dopamine receptors have been explored as potential treatments for schizophrenia.

Dopamine D₁ receptors

Significant evidence exists for the importance of dopamine D₁ receptors in the pathophysiology of schizophrenia, particularly having a role in cognitive dysfunction⁴⁶. In drug-naïve patients with schizophrenia, a decreased level of D₁ receptor-like binding in the prefrontal cortex on PET imaging was correlated with the severity of negative symptoms and cognitive dysfunction⁴⁷. In addition, chronic blockade of D₂ receptors results in a down-regulation of D₁ receptors in the prefrontal cortex and consequently produces severe impairments in working memory in non-human primates⁴⁸. This downregulation of D₁ receptors may explain why long-term treatment with typical antipsychotic drugs may contribute to the cognitive dysfunction in schizophrenia. In fact, direct blockade of D₁ receptors with selective antagonists, predicted to have antipsychotic effects in early preclinical models, showed no antipsychotic efficacy in clinical trials and may have exacerbated symptoms in some patients^{49,50}. Thus, current efforts are focused on a possible role of D₁ receptor agonists in treating the cognitive dysfunction in schizophrenia. Indeed, short-term administration of the D₁ selective agonist, ABT-431, reversed the cognitive deficits in monkeys treated chronically with a D₂ receptor antagonist⁴⁸. Other studies have also shown cognitive enhancement with a partial agonist of the D₁ receptor and selective, full D₁ receptor agonists in non-human

primates^{51,52}. Thus, novel compounds targeted at stimulating D₁ receptor signaling either directly or indirectly may be of immense value in treating cognitive deficits in schizophrenia, though some potential pitfalls may need to be overcome. First, in addition to insufficient D₁ receptor activity, excessive D₁ activity such as that resulting from acute stress may also be deleterious to cognition⁵³. In addition, chronic treatment with a D₁ receptor agonist may actually lead to downregulation of the D₁ receptor potentially worsening cognition in the long-term. Thus, an optimized level of D₁ receptor activation may be required to realize full cognitive benefits⁵³, which may be accomplished by partial agonists or an intermittent pattern of administration^{48,54}.

Dopamine D₃ receptors

D₃ receptors are structurally similar to D₂ receptors and thus most antipsychotics have relatively high affinity at this site⁵⁵. As such, significant effort has been placed on investigating the potential role of the D₃ receptor as a target for drug development in schizophrenia. Indeed, a post-mortem study of drug-free patients with schizophrenia demonstrated elevated D₃ receptor levels in contrast to normal D₃ receptor partial agonist was able to block the increase in locomotor activity in mice induced by N-methyl-D-aspartate (NMDA) glutamate receptors antagonists, such as phencyclidine or ketamine, a frequently used preclinical model of psychosis⁵⁷. As such, multiple selective dopamine D₃ agents are currently in clinical trials for the treatment of schizophrenia. For example, A-437203 is currently undergoing Phase II trials, although

clinical data are not yet available⁵⁵ as is SB-773812 (Clinical Trials @.gov identifier NCT00259870). Development of another agent, PNU-177864, which is a partial agonist at D₃ receptor appears to have been stopped due to safety concerns⁵⁸. Thus, the potential antipsychotic efficacy of selective D₃ receptor agonism and antagonism remains unknown at this time, though some data suggests the benefit of D₃ receptor partial agonists in the treatment of Parkinson's disease and drug addiction⁵⁹. Additional preclinical studies have also suggested a role of D₃ receptor antagonists in improving negative symptoms⁶⁰ and working memory⁶¹, though clinical evidence is unavailable.

Dopamine D₄ receptors

When the dopamine D₄ receptor was initially cloned it was also found that clozapine had higher affinity for this receptor than for D₂ receptors creating significant speculation that the D₄ receptor may be clozapine's "magic receptor"⁶². Further support of a role of D₄ receptors in schizophrenia came from postmortem studies showing higher levels of D₄ receptors in the forebrain, though these results have not been entirely consistent among studies⁶³. Even so, clinical trials of D₄ antagonists have not demonstrated any appreciable efficacy in the treatment of acute schizophrenia. For example, randomized, controlled trials of L-745870 and sonepiprazole found no differences in clinical responses compared with placebo-treated patients with schizophrenia^{64,65}. In addition, a trial of finanserin, a potent antagonist at both D₄ and serotonin 5-HT_{2A} receptors, also found no evidence of antipsychotic efficacy versus placebo in patients with schizophrenia⁶⁶. These clinical trial failures have suggested

that selective D₄ receptor antagonism alone is not responsible for the antipsychotic efficacy of clozapine; however, it is possible that D₄ receptor blockade in collaboration with action at other neurotransmitter receptors may be clinically beneficial. Indeed, studies of the physiological roles for the D₄ receptor are finding that D₄ receptors may play an important role in impulsivity and working memory⁶³. For example, recent findings demonstrated that D₄ receptors in hippocampal neurons can decrease NMDA receptor activity⁶⁷ and inhibit glutamatergic signaling in the frontal cortex⁶⁸. In addition, D₄ antagonists were observed to reverse phencyclidine-induced cognitive impairment in monkeys⁶⁹, together suggesting a suggesting that D₄ receptor-selective agents may be valuable in the treatment of the cognitive deficits in schizophrenia.

Catechol-O-methyltransferase

Catechol-*O*-methyltransferase (COMT) is a postsynaptic enzyme that methylates and thereby deactivates synaptically released catecholamines, particularly dopamine⁷⁰. Historically, monoamine oxidase was considered the primary enzyme for the initial deactivation of synaptic dopamine⁷¹, though mounting evidence suggests that COMT may be especially important for the breakdown of dopamine, particularly in the prefrontal cortex⁷². For example, COMT knockout mice show increased baseline levels of dopamine, but not other catecholamines such as norepinephrine, specifically in the frontal cortex⁷³. In addition, the COMT knockout mice also showed enhanced memory performance⁷³, suggesting a potential role of COMT inhibition in improving cognition. Indeed, a selective, reversible inhibitor of COMT, tolcapone, has been reported to

improve working memory in rodents⁷⁴ and has been shown to improve cognitive dysfunction in patients with advanced Parkinson's disease⁷⁵, though use is limited due to a risk of liver failure⁷⁶. Other COMT inhibitors are currently being investigated for treatment of the cognitive dysfunction in schizophrenia.

Interestingly, a common single nucleotide polymorphism (SNP) in the gene encoding COMT (val108/158met) results in the transcription of a variant of the COMT enzyme with approximately 40% less enzymatic activity in humans⁷⁷. The reduced activity associated with the met variant presumably results in greater availability of dopamine in the prefrontal cortex and, thus, may be linked to some aspects of cognition in humans. Furthermore, accumulating evidence predicts that patients with schizophrenia who have the met allele may have improved cognitive response to clozapine⁷⁸. The potential of pharmacologic inhibition of COMT in the long-term treatment of the cognitive dysfunction in schizophrenia, however, remains to be determined.

Serotoninergic Approaches

As serotonin receptors have been postulated to play a critical role in the action of the atypical antipsychotic drugs, we will briefly review a few of the serotonin receptors that continue to be targets in drug development for schizophrenia.

Serotonin 5-HT_{2A} receptors

Since the report that the atypicals, as a group, bind with higher affinity to 5-HT_{2A} receptors than to dopamine D₂ receptors^{79,80}, selective 5-HT_{2A} receptor antagonists have been extensively explored as putative antipsychotic drugs. Unfortunately, however, the 5-HT_{2A} selective compound M-100907, was discontinued after two Phase III trials found M-100907, although more effective than placebo, failed to reduce symptoms to the same extent as haloperidol⁸¹. A phase 2 study of the 5-HT_{2A/2C} antagonist SR46349B (eplivanserin) showed efficacy similar to haloperidol and better than placebo⁸². Thus, it is now clear that while selective 5-HT_{2A} receptor antagonists may have antipsychotic properties, they are not superior to D₂ antagonists. It is likely that the predominant role of 5-HT_{2A} receptors in antipsychotic action is to modulate dopaminergic tone, particularly along the mesocortical pathway^{83,84}. However, these studies also provide insight into why compounds with more complex pharmacologic profiles are likely superior to the "magic bullet" approach in the treatment of complex diseases such as schizophrenia^{31,45}.

Serotonin 5-HT_{1A} receptors

In addition to antagonism of the 5-HT_{2A} receptor, the agonist effects of clozapine on 5-HT_{1A} receptors have been postulated to contribute to its superior efficacy⁸⁵. Research has also demonstrated that 5-HT_{1A} receptor agonism may actually result from 5-HT_{2A} receptor antagonism suggesting that 5-HT_{1A} agonism alone may produce an

atypical antipsychotic drug when coupled with weak D_2 antagonism. Indeed, aripiprazole, a D_2 receptor partial agonist, may owe some of its atypical properties to its net effect of weak D_2 receptor antagonism, 5-HT_{2A} receptor antagonism and 5-HT_{1A} receptor agonism^{32,38,86}. As such, 5-HT_{1A} receptor modulation is most likely to play a role in regulating dopaminergic tone similarly to 5-HT_{2A} receptors⁸⁴, thus contributing to atypicality. Particularly, 5-HT_{1A} receptor agonism has been suggested to enhance dopamine levels in the prefrontal cortex⁸⁷, which may be related to the modest efficacy of many atypicals in treating the negative symptoms and cognitive dysfunction of schizophrenia. Thus far, attempts to develop medications combining 5-HT_{1A} receptor agonism with other receptor binding activities have not full replicated the superior clinical profile of clozapine, again highlighting the need for compounds with more complex pharmacologic profiles³¹.

Serotonin 5-HT₄ receptors

Serotonin 5-HT₄ receptors are found at high densities in the hippocampus, frontal cortex and amygdala, suggesting a role of these receptors in cognitive functions⁸⁸. Indeed, 5-HT₄ receptors have been shown to be markedly decreased in patients with Alzheimer's disease⁸⁹. 5-HT₄ receptor agonists have shown promise in the improvement of cognitive function by enhancing cholinergic transmission in the hippocampus⁸⁸, thus are being developed for the treatment of Alzheimer's disease. Interestingly, a recent study showed that the activation of 5-HT₄ receptors in a neuronal culture inhibited the secretion of β -amyloid peptide and enhanced neuronal survival⁹⁰.

While 5-HT₄ receptor-selective agonists are mostly being studied for their role in the treatment of Alzheimer's disease, they may also be of benefit in the treatment of the cognitive dysfunction in schizophrenia.

Serotonin 5-HT₆ receptors

As several atypical antipsychotics, including clozapine and olanzapine exhibit high nanomolar affinity for 5-HT₆ receptors⁹¹, significant efforts have been made to understand its possible role in schizophrenia and other neuropsychiatric disorders⁹². Studies in rodents have suggested a role of the 5-HT₆ receptors in the control of cholinergic neurotransmission⁹³, and the selective 5-HT₆ receptor antagonist SB-271046 has been shown to improve memory retention in the water maze test of spatial learning and memory⁹⁴. Thus, 5-HT₆ receptors may have an important future role in the treatment of cognitive deficits in neuropsychiatric illnesses such as Alzheimer's disease and schizophrenia.

Other Monoaminergic Approaches

*α*₂ adrenergic receptors

In the prefrontal cortex, α_2 adrenergic receptors appear to play an important role in cognitive functioning⁹⁵. Indeed, treatment with the α_2 adrenergic receptor agonists clonidine and guanfacine has been shown to improve cognitive performance in small trials of patients with schizophrenia^{96,97}. In addition, patients randomized to risperidone plus guanfacine showed significant improvement on tasks of working memory and attention compared with patients receiving typical antipsychotics plus guanfacine⁹⁷. Thus, α_2 adrenergic receptor activity is likely to be important in the development of new drugs for schizophrenia that can improve cognition. Complicating the picture, however, is the fact that clozapine and other atypicals have potent antagonist properties at α_2 adrenergic receptors⁹⁸, which may contribute to the atypicality of atypicals by preferentially enhancing dopaminergic transmission in the frontal cortex over subcortical dopaminergic pathways⁹⁹. Indeed, combined treatment of a selective α_2 adrenergic receptor activity similar to clozapine¹⁰⁰. Thus, balancing α_2 adrenergic receptor activity to achieve both antipsychotic and pro-cognitive efficacy may be challenging.

Cholinergic Approaches

Acetylcholine is known to play an important role not only in motor function, but also in various domains of cognition, particularly attention, learning, and memory¹⁰¹. Indeed, cholinergic dysfunction has been shown to be central to the pathophysiology of Alzheimer's disease and has also been postulated to contribute to the cognitive deficits of various neuropsychiatric disorders, including schizophrenia¹⁰². Cholinesterase

inhibitors, such as donezepil and rivastigmine, are currently the main pharmacologic approach to the treatment of Alzheimer's disease and have been shown to slow the cognitive decline in this neurodegenerative disease¹⁰³. As such, it has been proposed that cholinesterase inhibitors may also be useful in the treatment of the cognitive dysfunction in schizophrenia¹⁰⁴. Indeed, there have been multiple small randomized controlled trials of cholinesterase inhibitors in patients with schizophrenia, though results have been disappointing¹⁰⁵. In addition to cholinesterase inhibitors, significant efforts are underway to explore the modulation of various subtypes of both muscarinic and nicotinic acetylcholine receptors in the treatment of schizophrenia.

Muscarinic acetylcholine receptors

Of the five known muscarinic acetylcholine receptors (M_1 – M_5), the M_1 receptor has been most closely linked to cognition and schizophrenia¹⁰⁶. For example, decreased M_1 receptor binding has been reported in postmortem studies of the prefrontal cortex, hippocampus, and striatum from patients with schizophrenia¹⁰⁶, suggesting that M_1 receptor agonism might by beneficial in treating the cognitive dysfunction in schizophrenia¹⁰⁶. Indeed, the salutary actions of clozapine on cognition have been hypothesized to be due in part action at M_1 receptors¹⁰⁷. However, studies have variably reported clozapine to be both an agonist and an antagonist at M_1 and other muscarinic receptors¹⁰⁶. Interestingly, the major active metabolic of clozapine, Ndesmethylclozapine, has been reported to be a potent M_1 agonist that preferentially binds to M_1 receptors versus clozapine¹⁰⁸ although more comprehensive studies fail to demonstrate selectivity N-desmethylclozapine for M_1 receptors⁴². In addition. Ndesmethylclozapine has high affinities for 5-HT_{2A} and 5-HT_{2C} receptors, and is a partial $D_{2/3}$ receptor agonist^{41,43}, suggesting that this metabolite of clozapine may also have antipsychotic and cognition-enhancing properties. Indeed, N-desmethylclozapine (ACP-104) and other M₁ receptor agonists are in clinical trials as potential treatments of the cognitive dysfunction in schizophrenia. Xanomeline, a non-selective muscarinic agonist with potent actions at a variety of non-muscarinic GPCRs including 5-HT_{1A} and 5-HT_{2A} receptors¹⁰⁹ improved cognition and psychotic-like symptoms in Alzheimer's disease, but was discontinued due to poor tolerability¹¹⁰. The relatively non-selective actions of xanomeline at a number of GPCRs (http://pdsp.med.unc.edu/pdsp.php) should engender caution among schizophrenia researchers for embracing positive data from xanomeline studies as being specifically indicative of a role for M1 receptors in schizophrenia. Overall, evidence suggests M₁ receptor agonists could be useful in treating various symptom domains in schizophrenia, though the roles of the other muscarinic receptor subtypes are less clear.

Nicotinic acetylcholine receptors

It is well known that the smoking rates in individuals with schizophrenia are significantly higher than in the general population and some have suggested that these individuals may be 'self-medicating' with nicotine¹¹¹. Indeed, nicotine administration has been shown to improve various measures of cognition may ease some of the side effects of antipsychotic medications¹¹¹. Thus, considerable research has explored the

potential use of nicotinic agents for the treatment of schizophrenia, specifically selective agonists and antagonists at various subunits of the nicotinic acetylcholine receptor. For example, the α 7 nicotinic receptor subtype modulates auditory gating, a process known to be deficient in schizophrenia¹¹² and agonists at α 7 receptors such as 3-2,4-dimethoxybenzylidene anabaseine (DMXB-A) can normalize the auditory gating deficits in rodents¹¹³. Moreover, DMXB-A had a positive effect on a cognitive battery in a small proof-of-concept trial in humans¹¹⁴, and additional clinical trials of α 7 receptor agonists are underway. However, long-term use of α 7 agonists may induce the desensitization of nicotinic receptors, leading to a limited duration of efficacy¹¹².

It has also been suggested that $\alpha 4\beta 2$ nicotinic receptors are involved in cognition, and agonists of $\alpha 4\beta 2$ receptors such as RJR 2403 can produce significant and long-lasting improvement of memory in rats¹¹⁵. Thus, nicotinic $\alpha 4\beta 2$ receptor agonists may be of therapeutic benefit for the treatment of the cognitive deficits in schizophrenia. In addition, allosteric modulators of nicotinic receptors are being explored as therapeutic agents. For example, galantamine is a positive allosteric modulator of nicotinic receptors in addition to being an acetylcholinesterase inhibitor¹¹². The allosteric interaction of galantamine with nicotinic receptors can enhance the channel activity induced by a receptor agonist, either endogenous acetylcholine or theoretically a coadministered subtype-selective agonist.

Glutamatergic Approaches

Since the 1950s, the *N*-methyl-D-aspartate (NMDA) glutamate receptor antagonists phencyclidine (PCP) and ketamine were known to produce a large range of schizophrenia-like symptoms including psychotic symptoms, negative symptoms and cognitive dysfunction¹¹⁶. Thus, it has been hypothesized for decades that some deficiency in NMDA function might play a role in the pathophysiology of schizophrenia¹¹⁶, suggesting that drugs that can augment NMDA receptor activity may have therapeutic potential in schizophrenia. It is also important to note, that a competing hypothesis suggests that a hyperactivity of glutamatergic neurotransmission is involved in the psychopathology of schizophrenia, leading to seemingly contradictory pharmacologic approaches being explored¹¹⁶. Below, we briefly review various approaches being explored for modulating NMDA receptor neurotransmission and discuss approaches aimed at other glutamatergic mediators.

NMDA glutamate receptors

NMDA glutamate receptors are ligand-gated ion channels with a primary glutamate binding site and an allosteric glycine binding site¹¹⁶. Interestingly, the opening of the NMDA channel appears to require both glutamate and glycine binding and can be modulated by multiple substances, including Mg²⁺, polyamines, and protons, at various allosteric sites¹¹⁶. Thus, there are multiple potential sites to target for enhancing NMDA receptor activity; however, direct agonists of the glutamate binding site of the NMDA receptor may not be clinically feasible due to the risk of excess

excitation causing neurotoxicity and seizures. Therefore, the allosteric sites on the NMDA receptor complex, particularly the glycine binding site have been targeted for development of pharmacotherapy in schizophrenia.

Compounds that target the glycine site of the NMDA receptor complex have been studied in multiple small clinical trials and include the amino acids glycine, Dcycloserine, D-serine and D-alanine¹¹⁷. In most of these studies, the test compound was administered along with either a typical or atypical antipsychotic, and there appears to be significant benefits reducing negative symptoms and cognitive impairment in patients with schizophrenia¹¹⁷. Of the four agents, D-cycloserine has been the least efficacious, likely due to it being a partial agonist that acts as an antagonist at high doses. Interestingly, when used concurrently with clozapine, glycine¹¹⁸ and D-serine¹¹⁹ have been reported to be ineffective while D-cycloserine seemed to worsen symptoms¹²⁰, possibly because clozapine may already enhance glycine and glutamate neurotransmission. Overall, agonists at the glycine allosteric site of the NMDA glutamate receptor hold promise in the treatment of the negative and cognitive symptoms of schizophrenia, possibly as an augmentation of currently existing antipsychotics.

Glycine transporter

Another strategy being explored to boost NMDA activity at the glycine allosteric site is to increase synaptic glycine by inhibiting the glycine transporter. The use of

glycine transport inhibitors would have the advantage of avoiding the very high doses of glycine and D-serine that are needed. Indeed, preclinical data suggest that inhibition of glycine reuptake represents a feasible approach to enhance NMDA receptor activity and possibly be therapeutic in schizophrenia¹¹⁶. For example, selective, high-affinity inhibitors of the glycine transporter, including Org-24598¹²¹, N-[3-(4'-fluorophenyl)-3-(4'phenylphenoxy)propyl]sarcosine¹²² and SSR-504734 have been found to reverse PCPinduced hyperactivity and dopaminergic hyperreactivity in rodents^{121,122}. Clinical trials to date, however, have only studied the low potency glycine transport inhibitor sarcosine (N-methyl glycine). In a clinical trial of sarcosine added to the stable antipsychotic regimen of patient with schizophrenia, there was a highly significant reduction in negative symptoms, along with smaller but significant reductions in positive and cognitive symptoms¹²³. Interestingly, a subsequent study with patients on clozapine, found no improvement of symptoms with the addition of sarcosine, a result similar to studies with the NMDA glycine site agonists¹²⁴. These results strongly suggest a role of glycine transport inhibitors in the treatment of schizophrenia, though results of trials with selective, high-potency inhibitors are anticipated.

Metabotropic glutamate receptors

Agents acting at metabotropic glutamate receptors (mGluR) are currently in preclinical development. Specifically, there are two main groups of mGluRs being studied in schizophrenia, Group I receptors include mGluR1 and mGluR5 and Group II receptors include mGluR2 and mGluR3¹²⁵. Group I receptors increase presynaptic

glutamate release while Group II receptors inhibit presynaptic glutamate release, however agonists at each are being explored as potential treatments in schizophrenia demonstrating the duality of glutamatergic hypotheses in the pathophysiology of schizophrenia^{126,127}. Indeed, both approaches have shown efficacy in preclinical models of schizophrenia¹¹⁶, however, development of selective agents at mGluR subtypes has been an issue. Allosteric modulators of mGluRs hold promise as therapeutic agents and indeed, several groups have recently developed highly selective allosteric potentiators of these receptors^{127,128}. These selective allosteric modulators of mGluRs compounds may prove beneficial in the treatment of schizophrenia and preliminary positive results with an mGluR2 agonist in Phase II trials have been reported (http://www.prnewswire.com/cgi-bin/micro_stories.pl?ACCT=916306&TICK=LLY&STORY=/www/story/12-07-

<u>2006/0004487009&EDATE=Dec+7,+2006</u>).

Other ionotropic glutamate receptors

Another glutamatergic approach to drug development in schizophrenia has been the development of compounds that stimulate AMPA (alpha-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid) and kainate glutamate receptors. AMPA receptors help to activate NMDA receptors while NMDA receptors are required for proper incorporation of AMPA receptors into the postsynaptic membrane, a process involved in synaptic plasticity¹²⁹. Indeed, administration of the AMPA/kainate receptor antagonist, LY-293558, partially reversed the impairment of working memory induced by subanesthetic doses of ketamine in rats¹³⁰. Further preclinical data suggest that AMPA/kainate receptor antagonists may have antipsychotic efficacy and possible utility in the treatment of cognitive deficits in schizophrenia, though further research is indicated⁴⁰.

In apparent contrast to the postulated utility of AMPA/kainate receptor antagonists as antipsychotics, allosteric potentiators of AMPA receptor function, a class of compounds termed ampakines, are also being studied as potential treatments for schizophrenia¹²⁹. Ampakines may avoid the desensitization frequently seen with direct AMPA agonists and can enhance glutamatergic transmission, facilitating long-term potentiation, learning and memory in rodents¹²⁹. In a clinical trial of schizophrenia patients on clozapine, coadministration of the ampakine CX-516 yielded significant improvements in memory and attention¹³¹, however, a trial of CX-516 as monotherapy in schizophrenia showed no clear beneficial effects¹³². Importantly, higher potency ampakines are currently under clinical development as both monotherapy for schizophrenia and adjunctive treatment for cognitive dysfunction, though results of trials are not yet available. An initial trial with Org24448 has been planned for cognition enhancement in schizophrenia (NCT00425815) though progress has not yet been reported for this compound. Overall, it remains unclear if modulation of AMPA receptors by agonists, antagonists or allosteric modulators such as ampakines has therapeutic value in the treatment of schizophrenia although this is a highly active area of current research.

Other Approaches

Cannabinoid receptors

A recent meta-analysis demonstrated a statistically significant correlation of prior cannabis use and the development of schizophrenia¹³³, adding to a large amount of evidence implicating the endogenous cannabinoid system in schizophrenia¹³⁴. The endogenous cannabinoid system contains at least two cannabinoid receptors, the CB₁ and CB₂ receptors. A selective CB₁ receptor antagonist, SR-141716 showed activity in preclinical models of antipsychotic efficacy^{135,136}, however, in a recent clinical trial, SR-141716 failed to antipsychotic efficacy versus placebo⁸². Whether further clinical trials with cannabinoid receptor antagonists in schizophrenia are warranted is debatable.

Neurokinin receptors

Neurokinin 1 (NK₁) and neurokinin 3 (NK₃) receptors have been explored as potential targets for neuropsychiatric drug development¹³⁷⁻¹³⁹. NK₁ receptor antagonists may have efficacy in the treatment of depression, though a recent clinical trial of the NK₁-selective antagonist, aprepitant, for depression did not show efficacy versus paroxetine¹⁴⁰. NK₃ receptor antagonists, however, have been investigated as potential antipsychotic agents as NK₃ receptors appear to regulate midbrain dopaminergic function¹⁴¹. As such, several NK₃ receptor antagonists, including osanetant (Sanofi-Synthélabo) and talnetant (Glaxo Smith Kline), have been in development as potential

treatments for schizophrenia. In a recent clinical trial, osanetant showed statistically significant improvement in positive symptoms and global assessment versus placebo and was similar to haloperidol⁸², however an informal report of a follow-up study indicated negative results and the compound was discontinued¹⁴¹. No clinical trial data have been published to date talnetant and trials in schizophrenia appear to have been discontinued; thus whether NK₃ receptor antagonists may serve as novel antipsychotics either as monotherapy or as augmentation for the treatment of negative symptoms or cognition remains to be determined.

Neurotensin receptors

Neurotensin (NT) is a neuropeptide that, for decades, has been implicated in the pathophysiology of schizophrenia as it is closely associated with, and modulates dopaminergic and other neurotransmitter systems¹⁴². Indeed, significant preclinical data suggested a potential use of NT receptor agonists as novel therapeutic agents for the treatment of schizophrenia¹⁴². For example, administration of NT agonists, such as PD-149163, can reverse amphetamine-induced effects on hyperactivity and prepulse inhibition without inducing catalepsy¹⁴³. Thus, NT receptor agonists likely have potential in the treatment of schizophrenia; however, there have been no published clinical trials of NT agonists. Interestingly, there is also seemingly contradictory evidence indicating that neurotensin antagonists may have antipsychotic potential as there may be pathologically increased NT tone in schizophrenia¹⁴². A recent clinical trial, however, showed no antipsychotic efficacy of a potent and selective NT₁ receptor antagonist, SR-

48692, compared with placebo⁸². Thus, NT antagonists may not be useful for the treatment of schizophrenia; however, clinical trials of NT agonists are needed to explore this novel treatment strategy for schizophrenia.

Additional approaches

A number of other approaches for the development of novel therapeutics for the treatment of schizophrenia have been described including, cyclooxygenase-2 (COX2) inhibitors, phosphodiesterase 10A (PDE10A) inhibitors, neurosteroids, and secretin. COX-2 inhibitors such as celecoxib have been hypothesized improve cognitive performance by reducing inflammatory processes in the central nervous system¹⁴⁴. Indeed, in one small trial, there was some significant benefit with the addition of celecoxib to risperidone¹⁴⁵. PDE10A is a recently identified phosphodiesterase expressed at high levels in the brain and PDE10A inhibitors have been shown to antagonize the effects of both amphetamine and phencyclidine in rodents suggesting antipsychotic potential¹⁴⁶. Secretin is a gastrointestinal peptide that has poorly defined roles in the brain, however, recent studies have suggested a possible therapeutic benefit in autism and transient improvement of symptoms in schizophrenia¹⁴⁷, though repeated intravenous administration is likely to limited therapeutic potential. Neurosteroids, such as dehydroepiandrosterone (DHEA) have been implicated in neuroprotection and enhancement of NMDA receptor neurotransmission suggesting therapeutic potential in schizophrenia⁴⁰. Indeed, a double-blind study of DHEA as an adjunct to antipsychotic treatment in chronic schizophrenic patients with prominent

negative symptoms suggests some efficacy at improving negative symptoms, especially in women¹⁴⁸, though further studies are needed.

Moving Towards the Future

As is apparent from the preceding sections, most of the current strategies for developing novel compounds for the treatment of schizophrenia have not been All currently available medications target D₂ dopamine receptors—a successful. paradigm that has dominated drug development for the past 20 years—and many have been identified by activity in preclinical models that were devised based on pharmacologic manipulation (such as psychostimulant-induced behaviors). Indeed. these models have helped identify additional antagonists at dopamine D₂ and serotonin 5-HT_{2A} receptors, and are helping to identify novel neurotransmitter approaches to modulate dopamine. In addition, the development of highly selective agents for various neurotransmitter receptor targets has been and will continue to be extremely valuable in the elucidation of brain physiology and understanding of the pathophysiology of complex disorders such as schizophrenia; however, future drug discovery approaches will have to be truly revolutionary and based on a better understanding of the pathogenesis of the disease. Interestingly, we are in an era of increased knowledge and enormous spending in biomedical research but a dearth of advances in therapeutics. This decrease in the introduction of fundamentally new drugs into clinical practice is evidence of attempts to make a fundamental shift in the basic paradigms

used for drug discovery. Thus, this is an exciting and pivotal time for the development of truly novels approaches to drug development and treatment of complex disorders like schizophrenia. Below, we will discuss some of the exciting advances in our understanding of the pathophysiology of schizophrenia, will highlight some novel strategies currently being explored for drug development, and will stress the need for and increased role of academic scientists in target identification and validation.

Models of the Pathophysiology of Schizophrenia

A major critique of current drug discovery approaches for schizophrenia is that adequate treatments cannot be developed because the underlying causes of major mental illnesses remain incompletely understood^{21,31,149}. Indeed, while there have been enormous advances in our understanding of the basic biological processes contributing to many human diseases, a detailed understanding of the processes underlying schizophrenia and other complex mental disorders remains elusive. With the sequencing of the human genome and the development of genomics-based technologies, there are unprecedented opportunities for gaining fundamental new insights into these complex diseases¹⁵⁰. Currently, at least three highly-overlapping hypotheses of the underlying pathophysiology of schizophrenia drive drug discovery efforts¹⁶.

The first hypothesis, and the one that accounts for all of the current antipsychotic medications and the vast majority of compounds in the pipeline, is the *signal*

transduction hypothesis that posits that basic alterations in receptor-medicated signal transduction induces schizophrenia-like psychopathology. Therefore, normalizing the altered signaling with medications targeting receptor and post-receptor molecules should be efficacious in treating schizophrenia^{116,151}. Indeed, targeting these neurotransmitter receptors sites has, to this point, been the predominate focus of psychopharmacological research, and this strategy has led to significant advances in our understanding of the pathophysiology of schizophrenia and brain function as a whole. Future efforts, however, should move beyond the current strategies of solely targeting the synaptic neurotransmission at the receptor level to the development of agents that can affect more diverse cellular functions including intracellular signaling pathways and the mechanisms involved in synaptic plasticity.

Secondly, *the molecular-genetic hypothesis* posits that it is strong effects of susceptibility genes underlying the pathophysiology of schizophrenia¹⁵², and suggests that targeting drugs at these genes or their associated anatomic and functional pathways might yield novel and more effective treatments for schizophrenia^{153,154}. Indeed, significant progress has been made in recent years on elucidating various susceptibility genes in schizophrenia, including dysbindin, neuregulin 1, COMT, DISC1 and others¹⁵⁵. Interestingly, many of these genes appear to be related to the control of synaptic plasticity and glutamate transmission (particularly NMDA receptor function). These recent breakthroughs in genetic studies of schizophrenia begin to allow for hypothesis-driven approaches for developing of actual disease-modifying drugs for

schizophrenia. In addition, individualized treatment strategies could be developed that are focused on subgroups of schizophrenia patients with specific susceptibility alleles.

A third hypothesis, the neural network hypothesis, proposes that schizophrenia results from the strong effects of altered neuronal integration. Thus, this hypothesis predicts that drugs that fundamentally reset the tone of networks of neuronal interactions will prove efficacious in treating schizophrenia^{149,156}. Indeed, significant evidence exists suggesting that schizophrenia is a neurodevelopmental disorder associated with abnormal connectivity resulting from defects in synaptic pruning and migration of neurons¹⁵⁷. Thus, if alterations in synaptic pruning are the primary process underlying the pathophysiology of schizophrenia, possibly due to inherited genetic alterations in genes such as DISC1 or dysbindin, then effective treatment strategies should target the underlying deficits [Figure1 - Synapse]. In addition, successful treatment of schizophrenia may then require early recognition and treatment during or even before an obvious prodromal stage. However, if the underlying defects are due to abnormal migration of cortical neurons and subsequent dysregulation of cortical development, it may be impossible to ameliorate such deficits via simple pharmacological approaches.

Challenges of future drug discovery

While there has been significant progress in our understanding of the underlying pathophysiology of schizophrenia, a great deal of additional research is needed before

we can begin the systematic development of drugs that may address the root cause of the disease. Thus, before our full understanding of those causes, the development of novel drugs with superior efficacy and improved side effect profiles is still essential. It has been suggested that "selectively non-selective" drugs, or "magic shotguns," can already be developed and may fill a clinical need for improved therapeutics^{31,44}. Indeed, genomics-based screening approaches are being used to identify novel drug candidates based on their ability to either mimic the gene expression "signature" of gold-standard drugs like clozapine, or based on their ability to normalize the expression of genes that are altered in schizophrenia⁴⁵. Alternatively, high-throughput behavioral screenings may prove useful for the identification of novel medications for schizophrenia⁴⁵, though hurdles exist in finding animal behavioral models with good predictive value. This lack of predictive, reliable and efficient animal models has severely hindered progress in discovering novel therapeutics for schizophrenia, highlighting a need for increased collaboration between scientists in academic settings and industry.

Indeed, the translation of newly gained knowledge into fundamentally new therapeutics is a major challenge facing the biomedical research community. Fiscal pressures that govern research efforts in industry make it increasingly difficult for companies to invest significant resources in exploratory projects and basic research that capitalize on translating discoveries of basic science into marketable products. Because of this, companies frequently launch expensive drug discovery and development programs based on intriguing but often poorly validated targets for novel therapeutic approaches. For the treatment of schizophrenia, for example, preclinical

models are highly effective at predicting whether or not a candidate molecule would have 'atypical' properties, but are only fair at predicting overall efficacy and are ineffective at predicting efficacy greater than 'conventional treatment'. In addition, none of the available animals models accurately predicts the propensity of various antipsychotic drugs to induce weight gain and associated side-effects, although some of this can be predicted based on a knowledge of *in vitro* receptor pharmacology¹⁵⁸. Moreover, in terms of the negative and cognitive symptom domains in schizophrenia, none of the commonly used animal models are highly predictive, although preclinical memory models may be useful for predicting ability to enhance cognition. Thus, we advocate that academic-based scientists should be more aggressively involved in contributing to the drug discovery process, particularly by focusing on target validation¹⁵ [Figure 2 - Validation]. This challenge will require increased collaboration with the pharmaceutical industry as well as priority by the NIMH to fund such endeavors.

Conclusions

In the past 20 years, new therapies for schizophrenia have primarily emerged from a quest to discover new drugs that lack the extrapyramidal side effects of the typical antipsychotic drugs. Indeed, the atypicals have been beneficial to patients due to their improved therapeutic margin and their somewhat better efficacy at treating the negative symptoms of schizophrenia. However, the atypicals have significant side effect profiles as well, including weight gain and diabetes—likely due to off-target

actions at therapeutically irrelevant receptors^{158,159}. Thus, we have reached a significant bottleneck in the drug discovery process due to incomplete understanding of the mechanisms of action of the currently available antipsychotics as well as poorly defined pathophysiology for this complex and likely heterogeneous disorder. Interestingly, as many of the atypicals will soon go off patent, there is increased urgency within the pharmaceutical industry to develop new, novel treatments for schizophrenia.

We predict that the future of pharmacologic treatment of schizophrenia will likely start with the continued use of polypharmacy and augmentation strategies aimed at treating the multiple symptom domains of schizophrenia. This may be followed by the development of selectively-nonselective single compounds that can target multiple domains at once while simultaneously decreasing side effects, eliminating potential pharmacokinetic interactions and improving medication compliance^{31,45}. The long-term goal, of course, will be to develop "cure therapeutics"¹⁵⁴ which will likely require a substantial shift in the current paradigm of drug development and significant advances in our understanding of the pathophysiology of schizophrenia. Thus, it is important to continue pursue diverse molecular targets and increase efforts at validating novel Indeed, recent breakthroughs in genetic studies of schizophrenia have targets. provided renewed excitement that novel targets for the development that may target underlying disease processes. This shift in our approach to drug development will require considerable contributions from academic-based researchers as well as bold and potentially risky endeavors by the pharmaceutical industry.

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Table 1. Approximate ^a Pipeline of Drugs in Development for Schizophrenia						
ry Target Name ^b G	eneric Name	Originator	World Status			
dc						
A						
Multiple	clozapine	Novartis	Launched			
D2, D3	nemonapride	Astellas	Launched			
D2, 5-HT2A	olanzapine	Eli Lilly	Launched			
D2, 5-HT2A	quetiapine	AstraZeneca	Launched			
D2, 5-HT2A	risperidone	Johnson & Johnson	Launched			
D2, 5-HT2A	paliperidone	Johnson & Johnson	Launched			
D2, 5-HT2A	sertindole	Lundbeck	Launched			
D2, 5-HT2A	ziprasidone	Pfizer	Launched			
tial, 5-HT1A agonist	aripiprazole	Otsuka	Launched			
Multiple	asenapine	Organon/Pfizer	Discontinued by Pfizer ^d			
tial, 5-HT1A agonist	bifeprunox	Solvay	Phase III			
D2, 5-HT2A	iloperidone	Titan Pharmaceuticals	Phase III			
D2, 5-HT2A	blonanserin	Dainippon	Phase III			
-X-receptor activator	bexarotene	Non-industry source	Phase III			
Multiple AC	CP-104 (NDMC ^e)	Acadia	Phase II			
AMPA 1	Org-24448	Cortex Pharmaceuticals	Phase II			
Unspecified	TGOF02N	Fabre-Kramer	Phase II			
D2, 5-HT2A	ocaperidone	Johnson & Johnson	Phase II			
	•					
nischarin	idazoxan	Potomac Pharma	Phase II			
allosteric modulator	D-serine	Prestwick Pharmaceuticals	Phase II			
5-HT2A/2C	SR46349B	Sanofi-Aventis	Phase II			
mGluR2	unknown	Eli Lilly	Phase II			
NK3	osanetant	Sanofi-Aventis	Phase II			
5-HT1A agonist	SLV-313	Solvay	Phase II			
T transport inhibitor	SLV-310	Solvay	Phase II			
D2, 5-HT2A lurasi	done hydrochloride	Sumitomo	Phase II			
D2, D3 partial ap	lindore fumarate	Wyeth	Phase II			
sigma1 opioid	E-5842	Esteve	NDR ^f			
D2 partial (-)-	3PPP, Maryland	Non-industrial source	NDR			
agonist, 5-HT1A	SDZ-MAR-327	Novartis	NDR			
	ridone hydrochloride	Ferrer	NDR			
1, D2, 5-HT2A	ZD-3638	AstraZeneca	NDR			
CB1 CBD	cannabis derivative	GW Pharmaceuticals	Phase I			
α7 nAChR	MEM-3454	Memory Pharmaceuticals	Phase I			
	ALX-5407	NPS Pharmaceuticals	Phase I			
Unspecified	YKP-1358	SK Corporation	Phase I			
Unspecified	CRD-101	Curidium	Phase I			
D3, 5-HT1A	BTS-79018	Abbott	NDR			
R, AChR agonist Unspecified ur nischarin allosteric modulator 5-HT2A/2C mGluR2 NK3 5-HT1A agonist T transport inhibitor D2, 5-HT2A lurasi D2, D3 partial ap sigma1 opioid D2 partial (-)- agonist, 5-HT1A abaper 1, D2, 5-HT2A CB1 CBD α7 nAChR cine Transporter Unspecified Unspecified	dexefaroxan idine, Polifarma idazoxan D-serine SR46349B unknown osanetant SLV-313 SLV-310 done hydrochloride lindore fumarate E-5842 3PPP, Maryland SDZ-MAR-327 ridone hydrochloride ZD-3638 cannabis derivative MEM-3454 ALX-5407 YKP-1358 CRD-101	Pierre Fabre Polifarma Potomac Pharma Prestwick Pharmaceuticals Sanofi-Aventis Eli Lilly Sanofi-Aventis Solvay Solvay Sumitomo Wyeth Esteve Non-industrial source Novartis Ferrer AstraZeneca GW Pharmaceuticals Memory Pharmaceuticals NPS Pharmaceuticals SK Corporation Curidium	Phase II Phase I Phase I Phase I Phase I Phase I Phase I Phase I Phase I Phase I Phase I			

Table 1. Approximate^a Pipeline of Drugs in Development for Schizophrenia

D2, 5-HT1A	SSR-181507 SSR-125047	Sanofi-Aventis Sanofi-Aventis	NDR NDR
sigma1 opioid D3	AVE-5997EF	Sanofi-Aventis	NDR
D4	NGD 94-1	Schering-Plough	NDR

Preclinical

Glutamate antagonistADX-2 seriesAddexPreclinicalGlycine TransporterGlyT-1 inhibitors, OrganonAkzo NobelPreclinicalGlycine TransporterGlyT-1 inhibitors, Organon-2Akzo NobelPreclinicalGlycine TransporterGlyT-1 inhibitors, Organon-3Akzo NobelPreclinicalPeptidergic receptorABS-201Argolyn BiosciencePreclinicalDA antagonist, GABA agonistBL-1020BioLineRxPreclinicalUnspecifiedBGC-20-761BTGPreclinicalUnspecifiedGPCR allosteric modulatorsEli LillyPreclinicalD1D1 agonist D2 antagonistEli LillyPreclinicalUnspecifiedR-1678Hoffmann-La RochePreclinicalUnspecifiedneurolepticsIntra-Cellular TherapiesPreclinicalMGluR2mGluR2mGluR2 agonistMerck & CoPreclinicalGlycine TransporterSSR-504734Sanofi-AventisPreclinical				
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Glycine TransporterGlyT-1 inhibitors, Organon-3Akzo NobelPreclinicalPeptidergic receptorABS-201Argolyn BiosciencePreclinicalDA antagonist, GABA agonistBL-1020BioLineRxPreclinicalUnspecifiedBGC-20-761BTGPreclinicalUnspecifiedGPCR allosteric modulatorsEli LillyPreclinicalD1D1 agonist D2 antagonistEli LillyPreclinicalUnspecifiedcalcineurin modulatorsGaleneaPreclinicalUnspecifiedR-1678Hoffmann-La RochePreclinicalUnspecifiedschizophrenia therapyIntegragenPreclinicalUnspecifiedneurolepticsIntra-Cellular TherapiesPreclinicalmGluR2mGluR2 agonistRMG-40083RemergentPreclinical				
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α7 nAChR RMG-40083 Remergent Preclinical	•			
0		mGluR2 agonist	Merck & Co	
Glycine Transporter SSR-504734 Sanofi-Aventis Preclinical			5	
			Sanofi-Aventis	
Glycine Transporter SSR-103800 Sanofi-Aventis Preclinical	Glycine Transporter	SSR-103800	Sanofi-Aventis	Preclinical
Unspecified schizophrenia therapy Sequenom Preclinical			Sequenom	Preclinical
Glycine Transporter Org-24461 Servier Preclinical			Servier	Preclinical
α7 nAChR TC-5280 Targacept Preclinical	α7 nAChR		Targacept	
D1 BSF-78438 Abbott NDR	D1		Abbott	
D2, 5-HT2A, mAChR Org-23366 Akzo Nobel NDR	D2, 5-HT2A, mAChR		Akzo Nobel	
D3 BP4.879a Bioprojet NDR	D3	BP4.879a	Bioprojet	NDR
Unspecified CDD-0304 Cognitive Pharmaceuticals NDR	Unspecified		Cognitive Pharmaceuticals	
Unspecified neuroleptics CuraGen NDR	Unspecified	neuroleptics		
neuregulin 1 schizophrenia therapy deCODE genetics NDR			deCODE genetics	
sigma1 opioid sigma antagonists Esteve NDR	sigma1 opioid	sigma antagonists		
D3 SB-277011 GlaxoSmithKline NDR				NDR
5-HT4 5-HT4/D2 antagonists Johnson & Johnson NDR	-			NDR
Glycine Transporter GlyT-1 inhibs, Gliatech Merck & Co NDR			Merck & Co	
D1, D2, 5-HT2A GMC-283 Merck KGaA NDR	D1, D2, 5-HT2A			
D2 Y-931 Mitsubishi Pharma NDR				
D2 dopamine antags Neurogen Corporation NDR	D2	dopamine antags		NDR
Unspecified neuroleptic Orion Pharma NDR	Unspecified		Orion Pharma	
D3 PD-157533 Pfizer NDR			Pfizer	
D3, D2 PD-157695 Pfizer NDR			Pfizer	NDR
D2, 5-HT1A, D3 PD-158771 Pfizer NDR	D2, 5-HT1A, D3		Pfizer	
D4 PD-165167 Pfizer NDR				
D4 PD-172760 Pfizer NDR			Pfizer	
D3 U-99194A Pfizer NDR				
D4 U-99363E Pfizer NDR				
D3 PNU-177864 Pfizer NDR			-	
α7 nAChR PNU-282987 Pfizer NDR	α7 nAChR	PNU-282987		
Unspecified clozapine-DHA, Protarga Sankyo NDR				
D2, 5-HT, D4 HMR-2934 Sanofi-Aventis NDR				
Unspecified P-1704 Sanofi-Aventis NDR	Unspecified			
D1 LE-300 Sanofi-Aventis NDR				
sigma1 opioid MS-377 Schering AG NDR	sigma1 opioid			
D4 SPI-376 Spectrum Pharmaceuticals NDR	D4	SPI-376	Spectrum Pharmaceuticals	NDR

ciama 1 aminid		Taiaha	NDD
sigma1 opioid	NE-100	Taisho	NDR
mGluR	mGluR agonists	Taisho	NDR
α7 nAChR	TC-1698	Targacept	NDR
Discontinued ⁹			
D2, 5-HT2A, 5-HT1A agonist	1192U90	GlaxoSmithKline	Discontinued
D4	belaperidone	Abbott	Discontinued
sigma1 opioid	E-6276	Esteve	Discontinued
D3	RGH-1756	Gedeon Richter	Discontinued
sigma1 opioid	rimcazole	GlaxoSmithKline	Discontinued
Unspecified	EMD-66352	Merck KGaA	Discontinued
D2	SDZ-HDC-912	Novartis	Discontinued
D4	sonepiprazole	Pfizer	Discontinued
D2	(S)-amisulpride	Sanofi-Aventis	Discontinued
5-HT2A	fananserin	Sanofi-Aventis	Discontinued
sigma1 opioid	SR-31742A	Sanofi-Aventis	Discontinued
sigma1 opioid	MS-355	Schering AG	Discontinued
D2	remoxipride	AstraZeneca	Discontinued

^aThis is an approximation of the pipeline of drugs being developed for schizophrenia, adapted from BL Roth and PJ Conn: IOM White Paper, 2006. Attempts were made to make this table as accurate as possible, though due to the scarcity of published material the authors can accept no responsibility for the currency and accuracy of this table. Subsections of the table are in no particular order. ^bCompounds are assumed to be antagonists at each listed target unless otherwise specified.

^cOnly includes the atypical antipsychotics

^dOrganon may be continuing development,

(http://www.medicalnewstoday.com/medicalnews.php?newsid=57683)

^eNDMC, N-desmethylclozapine

^fNDR, no development reported, compounds are listed as to their last known Phase of development.

⁹Recently discontinued compounds

Receptor ^b	haloperidol	clozapine	N-DMC ^c	risperidone	paliperidone (9-OH- risperidone)	iloperidone	asenapine (Org-5222)
D ₁	122	266	14	244	41	129	2.9
D_2	2.1	141	101	2.4	1.6	11	1.4
D_3	5.4	347	153	8	3.5	11	1.8
D_4	3.9	23	64	5.8	54	14	1.8
D_5	124	255	284	290	29	319	23
5-HT _{1A}	2067	134	14	423	617	93	32
5-HT _{2A}	83	9.3	11	0.34	1.1	1.9	0.28
5-HT _{5A}	2247	3857	351	206	278	N/A	4
5-HT ₆	5133	13	12	2057	2414	63	1.4
5-HT ₇	626	37	60	5.6	2.7	112	0.72
α_{1A}	12	1.6	105	5	2.5	N/A	4.4
α_{2A}	1932	90	138	151	3.9	162	8.5
M_1	>10000	14	68	>10000	>10000	4898	24
M_2	>10000	104	415	>10000	>10000	3311	79
M_3	>10000	32	96	>10000	>10000	>10000	39
M_4	>10000	18	170	>10000	>10000	8318	>10000
M_5	657	28	35	>10000	>10000	>1000	9.5
H_1	1698	1.3	3.4	20	19	12 ^c	0.16
H_2	1003	153	375	120	121	N/A	23

Table 2. Approximate K_i^a values (in nM) for selected current and pipeline antipsychotics

^aAveraged from cloned human receptor data from the Psychoactive Drug Screening Program database (http://pdsp.med.unc.edu/pdsp.php), and references therein.
 ^bAbbreviations: 5-HT (serotonin), D (dopamine), M (muscarinic acetylcholine), H (histamine), alpha (α-adrenergic)

^cN-desmethylclozapine ^dCloned human receptor data not available, data from human brain tissue

Primary Symptom Domains	Potentially Druggable Clinical Targets	Possible Pharmacologic Targets
Positive Symptoms	Hallucinations Delusions Formal Thought Disorder	Dopamine D_2 antagonists Dopamine D_2 partial agonists Dopamine D_3 antagonists/agonists Muscarinic M_1 agonists Glutamate modulators Cannabinoid CB ₁ antagonists Neurokinin NK ₃ antagonists Neurotensin NT1 agonists PDE10A inhibitors Glycine transport inhibitors mGluR2 positive modulators
Negative Symptoms	Blunted Affect Anhedonia Avolition Alogia Asociality	Dopamine D_1 agonists Dopamine D_3 antagonists/antagonists Serotonin 5-HT _{2A} antagonists Serotonin 5-HT _{1A} partial agonists NMDA modulators Glycine transport inhibitors Neurokinin NK ₃ antagonists Neurosteroids
Cognitive Deficits	Working Memory Attention/Vigilance Verbal Learning/Memory Visual Learning/Memory Reasoning/Problem Solving Information Processing Speed Social Cognition	Dopamine D_1 agonists Dopamine D_3 agonists COMT inhibitors Serotonin 5-HT _{2A} antagonists Serotonin 5-HT _{1A} partial agonists Serotonin 5-HT ₄ partial agonists Serotonin 5-HT ₆ antagonists Cholinesterase inhibitors Muscarinic M ₁ agonists Muscarinic M ₄ agonists Nicotinic α 7 agonists and modulators Nicotinic α 4 β 2 agonists NMDA positive modulators AMPA positive modulators Glycine transport inhibitors mGluR2/3 positive modulators Neurokinin NK ₃ antagonists COX2 inhibitors

Table 3. Possible Pharmacolog	gic Targets	in Schizophrenia
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Figure Legends

Figure 1. Hypothetical Roles of Schizophrenia Genes at a Glutamatergic Synapse. Pictured is a hypothetical schematic of various putative schizophrenia susceptibility gene products and how they may affect neurotransmitter signaling at a glutamatergic synapse. The schizophrenia genes include: DISC-1 (disrupted in schizophrenia-1), Dysbindin, NRG1 (neuregulin-1), RGS4 (regulator of G protein signaling 4), COMT (catechol-*O*-methyltransferase), PDE4B (phosphodiesterase 4B), G72, and DAAO (Damino acid oxidase). Other abbreviations are: Glu (glutamate), DA (dopamine), NMDA (*N*-methyl-D-aspartate glutamate receptor), 5-HT_{2A} (serotonin receptor 2A), mGluR5 (metabotropic glutamate receptor 5), D₁ (dopamine receptor 1), ErbB4 (ErbB-type tyrosine kinase receptor B4), cAMP (cyclic adenosine monophosphate), G_q/G_s (G proteins), PSD95 (postsynaptic density protein 95), D-ser (D-serine). Adapted from Harrison and Weinberger¹⁵² and Roth¹⁶.

Figure 2. Continuous Target Validation in Academia. Identifying and validating novel targets is a significant rate-limiting step in new drug development which has led to few new drugs with truly novel mechanisms of action. Translation of newly gained knowledge into fundamentally new therapeutics is a major challenge facing the biomedical research community and is limited by fiscal pressures governing research efforts in industry that make it difficult for companies to invest significant resources on risky exploratory projects and basic research. Thus, increased research in the academic setting is needed to identify selective compounds for novel targets that may

be used for testing hypotheses at these targets and for proof of concept experiments. Continuous validation of targets by academic scientists at each step in the drug development process, including proof of concept experiments in the clinical setting, may facilitate the development of fundamentally new therapeutics for schizophrenia. Adapted from BL Roth and PJ Conn: IOM White Paper, 2006.



