Antipsychotic Polypharmacy in Treatment of Schizophrenia; Should or Should Not?

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Antipsychotics have been utilized as the standard treatment for schizophrenia regardless of illness phase where antipsychotic monotherapy (APM) is routinely recommended as the gold standard rather than antipsychotic polypharmacy (APP). However, approximately 20 to 40% of patients with schizophrenia do not respond to APM based on randomized controlled clinical trials and large practical clinical trials indicating that the subgroup of patients with schizophrenia would need differential treatment approaches beyond traditional treatment strategies such as APM. Numerous studies have supported the use of APP in particular for patients with certain clinical situations including: failure to show efficacy or tolerability from treatment with APM, need for different treatment for targeting specific symptom domains, severe illness, failure to treatment with clozapine, skepticism about following treatment guidelines, or cross titration periods. Furthermore, recent large cohort studies and practical clinical trials have proposed more benefits of APP rather than APM in terms of rehospitalization, mortality, and specific symptoms. APP has recently become more widely utilized and recognized as one of the next treatment strategies to clinicians for patients with schizophrenia. Some experts have already proposed the revision of treatment guidelines incorporating APP as evidence-based treatment option for certain patients with schizophrenia. Taken together, APP now deserves an evidence-based and acceptable treatment strategy, not an empirical or preferential treatment approach for treatment of schizophrenia in contemporary clinical practice.

Key Words: Antipsychotic Agents; Polypharmacy; Schizophrenia; Cohort Studies

INTRODUCTION

Schizophrenia is a chronic and devastating mental illness needing a maintenance treatment for prevention of relapse and recurrence in a naturalistic practice setting. A number of antipsychotics have been developed and used to treat patients with schizophrenia, but the clinical outcomes of schizophrenia after proper treatment with antipsychotic agents are still unsatisfactory today.¹⁻⁵ Antipsychotic monotherapy (APM) has been the gold standard for schizophrenia treatment,¹⁻⁵ however, empirically 10 to 60% of schizophrenia patients respond poorly or only partially to APM in real practice.¹⁰⁻¹⁵ Based on recent individual patient data from randomized controlled trials (RCTs),¹⁵ approximately 2 out of 10 patients (19.8%) starting APM failed to show any symptom improvement after acute phase treatment for more than 4 weeks, however, it increased up to 4-7 out of 10 patients (43-67%) when applying different criteria for response to AP therapy (25/50% more reduction in Positive And Negative Syndrome Scale, PANSS).

Antipsychotic polypharmacy (APP) has been widely utilized for compensation of inadequate AP treatment response in real practice. APP is defined as the use two or more AP for treatments for schizophrenia for any reasons.¹⁶ Clinicians try APP in the treatment of schizophrenia for a number of reasons including; pharmacodynamic synergy expecting differential neurotransmitter receptor affinity and occupancy broadening the range of re-

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Phenчor antipsychotic polypharmacy to clinicians by which rigorous systematic data search such as PRISMA (Preferred Reporting Items for Systematic reviews and Meta-analyses) was not utilized. However, extensive and careful data search and review were performed by the author to produce unbiased data on the use of APP.

Published articles were identified from PubMed using the keywords 'antipsychotic,' 'polytherapy,' 'schizophrenia,' 'combination,' or 'polypharmacy.' There are a countless references on the use of antipsychotic polypharmacy to clinicians who rigorously use systematic data search such as PRISMA (Preferred Reporting Items for Systematic reviews and Meta-analyses) was not utilized. However, extensive and careful data search and review were performed by the author to produce unbiased data on the use of APP.

APP has been known to be utilized for 10 to 20% of schizophrenia cases in an outpatient basis, while 40% of schizophrenia cases in an inpatient basis. However, there has been wide variety of uses in epidemiological findings of APP due to a lack of established criteria on the number of APs, duration of AP combination, and other clinical situations related with AP use in treatment of schizophrenia.

According to the recent systematic review based on operational criteria using 147 studies,17 there were substantial differences in the prevalence and yearly trends of APP in North America (16%) with the lowest APP rates compared to those from Oceania (16.4%), Asia (32%) and Europe (23%).

Specifically for Asian regions, the recent Asian collaborative schizophrenia study18 including 15 countries (n=3,357) found that approximately 43% of participants were on APP which was substantially higher rates compared to that (32%) of previous systematic study.17 The mean dose of AP by chlorpromazine equivalents (CPZeq) was 424 mg/d. Among participating countries, Vietnam and Japan had the highest rates of APP usage, 59.2% and 57.4%, respectively. Such high prevalences of APP has been also consistently found in numerous independent studies in different geographic regions such as Korea, China, Brazil, Canada, America, and Japan.18-23

There has been no clear determination of how many number of APs are commonly used in APP, which is found to create a variable range of AP numbers. According to the extensive meta-analysis,17 17.8% were taking two APs and 0.2% were taking ≥3 antipsychotics. In a multicenter study24 using real world data (n=851), 19.2% of patients were on two APs, while 1.2% of patients were on three APs. In another recent study25 investigating the AP usage pattern (n=280), 44.1%, 24.4%, 1.4% and 0.7% of patients were on two, three, four and five APs, respectively. Currently available independent studies and meta-analysis tentatively showed that two APs were most frequently used in APP.

**WHY DO CLINICIANS UTILIZE APP IN ROUTINE PRACTICE?**

APP is not clearly defined by any consensus among experts as well as in treatment guidelines utilized for clinicians over the world and thereby it has been still one of major debates in the treatment of schizophrenia.

In general polypharmacy is defined to use five or more therapeutic agents for treating certain illness under indication,26 however, the specific criteria of APP yet not clearly been established. According to the recent study, it was found that APP was mostly defined simply as the use of two or more APs for treatment of schizophrenia.8 Indeed there should be numerous and diverse reasons for clinicians to utilize APP for treating their patients with schizophrenia in routine practice.

Given inadequate efficacy of APs for treatment of patients with schizophrenia, clinicians frequently and mainly use APP to ameliorate positive and/or negative symp-
Various reasons for choosing APP in routine practice.

Some clinicians are not willing to adhere to the recommended treatment guidelines proposing the use of APM due to insufficient persuasive reasons for the limited use of APP since inadequate clinical evidence exists from RCTs. Other reasons are also diverse including: cross titration, pressured feelings of clinicians themselves, treatment resistance cases, medical cost issues, prevention of relapse and recurrence, reduction of hospitalization, avoidance of high dose APM, preference of clinicians/patients, use of different pharmacological profile of individual AP for synergistic effects in treatment or counteracts against side effects, longstanding habits of clinicians for using cocktail therapy, differential choice of treatment strategy in acute/maintenance treatment phase, faster treatment response, and combination of different formulation of APs. Fig. 1 illustrates the various reasons for choosing APP in routine practice.

WHAT BENEFITS DO WE EXPECT FROM APP?

Despite many concerns about APP, we can also expect and consider its benefits in the treatment of schizophrenia. In the most recent meta-analysis using data of RCTs (n=31) comparing APP vs. APM in schizophrenia, overall psychotic symptom reduction, and study-defined responses after AP treatment were compared between APP and APM. Based on the results, APP was found to be superior to APM regarding total symptom reduction with a large effect size difference (SMD=−0.53), while it failed to show superiority over APM regarding study-defined multiple response rates (≥20% PANSS/BPRS reduction, ≥25% PANSS reduction, and ≥20% PANSS reduction or CGI-I of 1 or 2), which is in contrast to that of their previous meta-analysis. Indeed, a significantly greater response rate (at least 50% reduction of the PANSS score or BPRS score or CGI-I of 1 or 2; RR=0.76 and NNT=7) was found in the previous meta-analysis including only 6 relevant studies, while it was not replicated in the recent meta-analysis. In detail, APP was superior in inpatient only studies (n=4), Chinese studies (n=4) and non-North American/European studies (n=5), while the number of high-quality studies was too inadequate to address separate analysis. Interestingly, this superiority of APP over APM regarding both total symptom reduction and study-defined responses also became non-significant when analyzing only high-quality studies, indicating that such efficacy differences between APP and APM were mostly found in low-quality studies rather than by high-quality studies, expectation and selection biases in the studies included, and still lack of high-quality evidence comparing for efficacy between APP vs. APM.

According to the recent large-scale, non-interventional, retrospective-prospective parallel arm study comparing APP (addition of a second AP after >60 days of APM, n=7,901) vs. APM (switch to a new AP after >60 days of APM, n=5,480) in Hungary, significantly more psychiatric hospitalizations were found with APM than APP (hazard ratio, HR=1.69). Further, the most recent nationwide cohort study compared the risk of psychiatric rehospitalization between APP vs APM including 62,250 schizophrenia patients analyzing 29 different APP and APM treatment types for 10 years. Based on the results, clozapine plus aripiprazole polypharmacy ranked as the lowest risk of psychiatric rehospitalization in the total cohort, compared to that of clozapine monotherapy, with a difference of 14%. Also, such differences between clozapine plus aripiprazole polypharmacy vs clozapine monotherapy were more evident in a subgroup analysis regarding the first episode schizophrenia showing a difference of 22% favoring APP over APM. In addition, aggregated data analysis also showed that any APP had 7% to 13% lower risks of psychiatric rehospitalization as compared with any APM.

 Likewise a recent, large naturalistic study including acute-phase schizophrenia in Japan (n=1,543) also showed the clinical benefit of APP when the primary and secondary APM strategies failed in clinical practice where approximately 59% of the patients overall were responders to an initial or a second APM. As a next step treatment, APP

![Fig. 1. Various reasons for choosing APP in routine practice.](image-url)
Antipsychotic Polypharmacy

(first or third AP combination to the second AP) was administered to the non-responders (n=581, 37.7%) where 522 (89.8%) showed a CGI-I score ≤3, while only 10.2% of the remaining patients showed a CGI-I score of ≥4. The responder rate of 89.8% observed in APP was much higher than the reported response rate to clozapine (40%) in the previous meta-analysis including treatment-resistant schizophrenia (TRS).32

Interestingly, a 6-month, recent RCT (n=127)33 tried to evaluate clinical benefits and risks between the continuation of APP and switching to APM in an outpatient clinic. The primary endpoint was time to all-cause discontinuation and it was shorter for patients in the APM switching group than in the APP continuation group and the APM switching group showed more frequent treatment discontinuation than the APP continuation group. Overall, 86% (n=48) in the APP continuation group were still on both APs, while 69% (n=40) in the APM switching group were still on the same treatment indicating a 17% difference in AP change favoring APP over APM. Furthermore, those who stopped one AP were more associated with earlier change of their treatment in the APM switching group than in the APP continuation group; it was also notable that a significant portion of these individuals resumed their previous APP regimen. Despite a failure to show superiority of APP over APM since two thirds of patients successfully switched to APM, it clearly demonstrated that a subgroup of patients may experience practical benefits from APP rather than APM and that patients should be allowed to recover the previous APP if an adequate trial with APM comes to an end with unsatisfactory results or no utility.

WHAT CONCERNS SHOULD WE HAVE IN THE USE OF APP IN ROUTINE PRACTICE?

1. APP is strongly associated with high-dose antipsychotic treatment

The proposed rationales for the use of APP includes scientific perspectives on differential PD of currently available SGAs. A combination of two or more APs with different PD would bring about optimal occupancy of dopamine D2 receptor in certain portion of patients needing higher levels of D2 occupancy or result in a diverse range of multiple receptor activities beyond D2 receptor.34 These are associated with the augmentation of therapeutic effect in patients with TRS, hastening of treatment response, or targeting some specific comorbid conditions such as anxiety, sleep disturbance, and cognitive dysfunction, etc.

Another aspect is more practical and persuasive to clinicians in the use of APP. Combinations of low-dose, two or more, APs possessing differential affinities and occupancy of multiple receptors is expected to achieve significant reduction of SEs which are susceptible to high dose APM and to attain already existing efficacy. However, currently available evidence suggests that APP is strongly associated with high/excessive dosing trends in routine practice.35

According to the previously mentioned large 10-year cohort study,36 the daily doses of APs were substantially increased in both calculations “DDD” and “CPZeq” during the 10 year periods. The yearly DDD in AMP patients also significantly increased, however, it was found to double in APP patients indicating that APP is mainly utilized for augmenting inadequate efficacy rather than countering SEs. Such huge differences in total AP dose between APM and APP might be possibly explained by several aspects; prior inadequate or under-treatment of first-episode schizophrenia in 1996 and conversion to a sufficient level of treatment in 2005, trends toward aggressive SGA use based on increasing clinical experiences with SGAs since high-dose FGA use is vulnerable to SEs, intensive marketing of SGAs by manufacturers since it was the early period of SGAs launch in the market, and earlier control of psychotic symptoms to reduce hospitalization for compensation of insufficient admission capacity.

The recent study analyzed the claim-data system regarding community mental health outpatients who had been treated with the same pharmacologic regimen for at least 3 months in Canada (n=435).22 The mean prescribed daily dose (PDD/DDD ratio) was significantly higher in the APP group than in the APM group (1.94 vs 0.94), further to say the PDD/DDD ratio was also extremely excessive in the APP group compared to the APM group since a 1.5 PDD/DDD ratio is considered to be a cutoff for excessive dosing. The proportion of excessive dosing was also approximately three times higher in the APP group than in the APM group irrespective of primary diagnosis. It was more profound in patients who were treated with SAG plus SGA polypharmacy and when the dose of initial AP was high. More interestingly the mean PDD/DDD ratio for individual SGA increased when it was a part of APP regardless of primary diagnosis as well. A previous study36 also found a strong association between persistent excessive dosing and APP on admission. In another previous study,37 the PDD/DDD ratio was 2.6 in the APP group, while it was 1.3 in the APM group, which is in line with recent and previous similar studies.22 According to the recent East Asian study investigating the medication trend in 2005 (n=194) and 2010 (n=201),38 it was found that the rate of high-dose APM significantly decreased from 30.4 to 18.4% across the year, while the rate of high-dose APP significantly increased from 34.0 in 2005 to 45.3% in 2010, indicating a replacement of high doses APM treatment trends by a high-dose APP approach. In a regression analysis, APP was confirmed to be strongly associated with high doses of AP prescription compared to APM (odds ratio=18.6).38 Such high dose trends in APP (888.3 mg/d of CPZeq) compared to APM (445.1 mg/d of CPZeq) was also replicated in another Asian study.39

2. High doses of AP in APP is strongly associated with cognitive dysfunction

Currently, a large number of studies have shown possible differences in cognitive functions between FGA and
SGA where findings suggested that FGAs are mainly effective in controlling positive psychotic symptoms due to their affinity for and occupancy of D2 receptors, while SGAs are also effective to ameliorate or even improve some domains of cognitive dysfunction, not only positive symptoms. Such differences have also replicated in a number of well-designed, recent, systematic meta-analyses.40,41

A previous Japanese study (n=136) has investigated whether APP would have better a cognitive influence than APP irrespective of combinations of FGAs or SGAs. In the study, a significant negative correlation was found between cognitive function measured by composite scores of the Brief Assessment of Cognition in Schizophrenia (BACS) and the CPZeq dose of APs (BACS Z-score of difference was almost double favoring APP over APP). Such negative correlation between BACS scores and CPZeq doses was also significant in most individual cognitive components including verbal memory, motor speed, verbal fluency, attention, and speed of processing. Interestingly, the SGA APM group showed better cognitive function even when the APP was composed of SGA plus SGA. There were also no differences in cognitive function in the APP group, regardless of the classes of combined APs. Such data clearly indicates the high dose AP results in substantial deterioration of cognitive functions whether or not the APP regimen is SGAs or FGAs added to the first APs. In a previous study, it was also found that the average daily dose (ADD) of APs was significantly associated with the development of notorious cognitive impairment in schizophrenia patients. In fact, APP was strongly associated with high daily doses (12.1 mg/d of risperidone equivalent dose, RISeq) compared to APM (4.2 mg/d RISeq) as well as with poor cognitive functioning measured by BACS z-score. In the study, another important finding was that the ADD to create substantially detrimental cognitive dysfunction measured by the BACS score was found to be approximately RISeq 5 mg/d or more and additional increase of RISeq 2.3 mg/d would also cause a decrease of 0.5 standard deviations of the BACS score. Intriguingly, previous research found that dose-reduction of AP may lead to significant improvements in cognitive function measured by the Wisconsin card sorting test (WCST, 19.9% increase in total correct answers and 34.9% decrease in perseverative errors) in schizophrenia patients who were exposed to high-doses of APP, indicating the critical role of AP total dose in modification of cognitive dysfunction in schizophrenia patients.

3. Increase of metabolic syndrome (MS) risk with APP

MS is very important clinical issue in daily practice since we cannot avoid SE when we prescribe FGAs or SGAs for treating psychotic symptoms and it is highly associated with metabolic complications leading to increased cardiovascular mortality.44 MS is well-known to significantly increase the risks to developing Diabetes Mellitus (DM), stroke, coronary heart diseases and mortality.45 After 3 years of treatment with Aps, the trend of development of MS is gradually increasing, especially, SGAs are more associated with MS rather than FGAs.46

A recent meta-analysis investigated 126 analyses in 77 publications (n=25,692) regarding the association of APP and MS. According to the results, the overall rate of MS was 32.5%, giving only minor differences in accordance with different methodologies of studies included in the meta-analysis (i.e., criteria of MS definition, treatment setting and sample characteristics). According to another independent study using data from the records of 458 psychiatric inpatients to compare MS between APP and APM,47 the MS rate was significantly different between the two groups favoring APM (34.3%) over APP (50.0%). Some lipid markers such as HDL <40 mg/dL or Triglyceride/HDL-cholesterol >3.5 were also higher in APP group than in APM group. Such higher rates of MS in APP vs. APM have been consistently reported in a number of previous studies.42,48 A recent Japanese study has also proposed the association of APP (odds ratio=2.4) with the development of pre-metabolic syndrome, while APP was associated with neither pre-metabolic syndrome nor MS. This study strongly suggests that an adjustment of patients’ lifestyle and modulation of APP regimen could mitigate or prevent the development of MS. Another interesting point was that the visceral fat obesity group was associated with higher AP total daily dose.

However, there have been also mixed findings as to whether or not APP truly increase the risk of MS in comparison with APM, possibly proposing insufficient evidence to clearly answer the clinical question on this potential weak point regarding APP yet.49-51 In such studies,52 the significant baseline associations of increase in weight, body mass index, and other lipid parameters with APP compared to those with APM were not maintained at the end of the follow-up period, particularly, in schizophrenia patients following their first-episode, indicating that naturalistic clinical course and time effect, should be also considered in the development and attainment of MS associated with the use of APP during treatment period.

4. QTc prolongation risk in the use of APP

Traditionally AP use has been proposed to be associated with prolongation of the QT interval corrected for heart rate (QTc). Recently a cross-sectional survey (n=725) was conducted to investigate the relationship between APP and QTc interval. Among included patients, 186 (26%) were on APP and the mean cumulative AP dose was significantly higher in the APP group (PDD/DDD ratio=2.9) than APM (PDD/DDD ratio=0.8). As being expected, the mean QTc interval was significantly longer in the APP group (mean=420.9) than the APM group (mean=413.4). According to the large Italian network study based on routine practice (n=2,411), the APP treatment was positively associated with QTc prolongation despite heterogenous samples included in the study. A number of previous studies suggested that APP is potentially associated with the prolongation of QTc which is also relevant and speculative since the number of APs is a proxy of total AP dose, which...
is an established risk factor of QTc prolongation.\textsuperscript{57} Despite supporting evidence that APP may prolong the QTc interval, the previous meta-analysis\textsuperscript{60} failed to show clear evidence that APP significantly prolonged the QTc interval compared to APM due to a dearth of data regarding this research area and such research has been confined to the use of specific high-risk APs in the combinations such as ziprasidone, sertindole, or clozapine.

5. Poor treatment adherence in APP

Poor adherence is one of major barriers in achieving optimal clinical outcomes in schizophrenia patients and thus APP could pose a problem for maintenance of treatment since polypharmacy is clearly and consistently found to be associated with poor treatment adherence for many reasons.\textsuperscript{50} Poor adherence usually results in high rates of recurrence/ relapse within a few years of recovery from the first episode which is strongly associated with poor clinical outcomes, functional impairment, high medical costs, increased rehospitalization, and unnecessary antipsychotic prescription.\textsuperscript{60,61}

In a recent study,\textsuperscript{62} the non-adherence rate was 41.0% among schizophrenia patients, in which APP was found to increase the risk of non-adherence approximately twice as much compared to those with APM. There have been some debates about whether the number of APs are directly associated with poor adherence, however, a number of studies have clearly indicated that APP is strongly associated with increased risk of diverse SEs leading to poor adherence and persistence.\textsuperscript{63,64} APP was also found to be significantly associated with increased hospitalization.\textsuperscript{65} Hence it should be reasonable that APP could, at least indirectly, increase the risk of non-adherence and non-persistence, ultimately influences on the poor clinical course and treatment outcomes.

**DISCUSSION**

Based on currently available data from RCTs, small-scale open trials, and large cohort studies and findings from meta-analysis, APP has become closer to being a part of routine practice and not being a part of unacceptable practice in the last few decades. We have to consider practical points for proper use of APP in our routine practice.

According to intense analysis based on a large patient-level dataset of randomized multicenter trials, the rates of nonresponse and nonremission from APM for acute treatment for schizophrenia were notably high regardless of criteria for response and remission.\textsuperscript{15} APP can be actively considered if patients do not respond to adequate trials of APM with proper doses and durations of treatment or they cannot tolerate APM for any reasons (i.e., SE due to high-dose therapy). There have been no established adequate trial durations or numbers of APM. However, most treatment guidelines propose that at least two or more APMs should be tried before moving toward further treatment steps including APP. Furthermore, at least 8 weeks up to 4 months be used for evaluation of treatment effects coming from one APM trial based on treatment guidelines and existing literature.\textsuperscript{5,66}

According to the recent large cohort study that followed patients for 20-years,\textsuperscript{49} clozapine plus aripiprazole polypharmacy was associated with the best outcome regarding psychiatric rehospitalization among all the 29 different APM and APP types, giving 14% to 23% lower risk of rehospitalization than clozapine monotherapy which showed

**TABLE 2. Practical points in the use of antipsychotic polypharmacy in routine practice**

1. Make it clear why you use APP; target specific comorbid symptoms, improve overall psychopathology, or ameliorate side effects, etc.
2. Closely review all documents regarding past history of APs before starting APP
3. Measurement-based treatment would reduce the risk of unnecessary APP and it will give you proper assessment on the exact clinical status of your patients on when and how you go with APP treatment strategy, i.e., appropriate time for transition to APM, maintenance with APP, switch of APP agents
4. APP should be prudent for those who have failed to show response/remission despite of at least two or more adequate APM trials under correct diagnosis of schizophrenia
5. APP could be also appropriate for those with intolerance to APM
6. Consider different APs having differential pharmacological profile as APP to enhance efficacy and reduce side effects (D2 antagonist + partial D2 agonist)
7. Keep close monitoring the total AP doses to avoid unnecessary high dose exposure; minimal effective AP doses are also working is APP consider total AP doses occupying 65% to 80% of D2 receptors in APP
8. Do not reduce the total AP doses below chlorpromazine equivalent dose of 200 mg/d regardless of classes of APs when tapering off APs for transition to APM
9. Gradual tapering of AP should be always kept in mind upon transition to APM
10. Clozapine should be the most appropriate first agent for APP
11. Clozapine and aripiprazole were found to be the most beneficial APP combination based on clinical trial data
12. Discuss and hear your patients on their own choice on APP agents and family response if available
13. Reinstitute APP as maintenance treatment if relapse/recurrence is clearly documented and eminent upon APM
14. Consider long-acting injectable antipsychotics such as aripiprazole once-monthly or paliperidone palmitate in your APP regimen for acquisition of stable therapeutic level in one agent of APP and thereby easy monitoring and dose selection for the remaining AP regarding the total doses of APP

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the best outcome as APM. Interestingly, the clozapine doses were 426 mg/d and 399 mg/d in APM and APP, respectively, indicating that the reduced dose cannot be a main reason for better outcomes since the difference was slight, so diverse receptor activities from different APs (i.e., partial D2 receptor agonist effect of aripiprazole) may exert favorable treatment outcomes while ameliorating tolerability concerns. Any APP presented a 7% to 13% lower risk of psychiatric rehospitalization compared with any APM, indicating that rational APP should be considered proper and feasible particularly with the use of two different APs possessing different types of receptor profiles. The following could be also good examples: D2 receptor antagonist+ partial agonist; D2 receptor tight binding agent+loose D2 receptor binding agent, e.t.c.

The most recent, largest and longest cohort study has clearly stated the superiority of any APP over any APM, in terms of rehospitalization and mortality. Furthermore, improper switch to APM was vulnerable to increasing the risk of relapse and recurrence compared to staying on APP in chronic and stabilized patients with schizophrenia. Indeed Dr. Stahl suggested that the changing trend toward APP in the treatment of schizophrenia has already started and thereby treatment guidelines should properly incorporate the wise and wide usage of APP in certain clinical situations and for the subgroup of patients with schizophrenia, providing 12 sensible recommendations. Currently available data from large practical clinical trials, meta-analysis, and open trials, potentially demonstrated that APP appears to no longer be an eminence-based treatment approach but now it should be immersed as an evidence-based, acceptable treatment strategy in clinical practice. APP is no longer a dirty secret for clinicians as an option for treatment of schizophrenia in clinical practice since we cannot evenly apply APM for treating patients with schizophrenia. The naturalistic treatment settings are quite different compared to those of RCTs, more severe and selected patients are included in such controlled trials, while clinicians should meet very heterogeneous patients with diverse psychotic and comorbid conditions not easily responding to APM. Table 2 suggests practical points to consider in the use of APP in routine practice.

Best-match and properly-balanced APP may be one of the best-available treatment next steps and practical treatment options for patients with multiple treatment failures and tolerability issues today. It is the right time for intensive explorations and debates on the proper time and duration of APP, who should be the right patients for APP, wise shifting time/clinical situations from APP to APM, how to optimize the combination of APs in AP, clear benefits and risks of APP vs APM in individualized naturalistic treatment settings, and the appropriate revision of practice guidelines for APP which can promote secure APP and help policy makers to accept APP as one of routine treatment practices for the treatment of schizophrenia. Definitely, more adequately-powered and well-designed APP clinical trials should be attempted to help us determine the clinical benefits, best-available APP agents, disadvantages, pharmaco-economic aspects, and individualization of APP in routine practice.

CONFLICT OF INTEREST STATEMENT

None declared.

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