Commentary

How Many Treatments Before Clozapine?
Medication Choices Across the Spectrum of Treatment Resistance in Schizophrenia

Peter J. Weiden, MD

For some patients with schizophrenia, antipsychotic medications will completely eradicate all psychotic (positive) symptoms. But most patients are not that fortunate and continue to experience psychotic symptoms despite taking medication. This reality brings us to some critical questions that come up frequently in practice. When are persistent psychotic symptoms acceptable? If the present symptoms are not acceptable, and a medication change is warranted, then which medication comes next?

We often use response categories to guide medication choices. If a patient's symptoms are within the boundary of what we believe to be a good response, then the patient is a “responder.” Patients who seem to have no improvement with medication are categorized as “nonresponders” or called “treatment refractory.” What about those who show some improvement with medication but are too symptomatic to be categorized as a responder? A common example is when antipsychotics are able to prevent acute relapse but the person is still burdened by psychotic symptoms every day. These are patients whose antipsychotic response is somewhere between full response and complete treatment resistance.

In this issue of the Journal, the study by Schooler et al carefully selected outpatients with schizophrenia who represented this middle group. Outpatients who were categorized as “moderately refractory” agreed to a 6-month course of treatment of either clozapine or risperidone. The study was randomized, and medications were given under double-blind conditions. The study was done at a time when less than half of the patients had any prior experience with risperidone. While the question of relative effectiveness of clozapine and risperidone is important on its own, the broader question of the role of using clozapine or any 1 of the 12 post-clozapine antipsychotic options across the spectrum of treatment-resistant schizophrenia is also very important.

Before discussing the results, let’s put this study into the larger context of treatment-resistant schizophrenia. The modern history of “treatment resistance” comes from the 1988 publication of the breakthrough study (“Clozapine for the Treatment-Resistant Schizophrenic,” which is also known as “Study 30”). The definition of treatment-resistant schizophrenia was, by today’s standards, severe. The study took place in long-term inpatient facilities in which the average duration of current hospitalization was 2 years. All had been treated and failed at least 2 of the antipsychotics available at that time, including a prospective treatment of 6 weeks of high-dose haloperidol. If there was no improvement, only then were patients randomly assigned to clozapine or chlorpromazine. Clozapine-treated patients had a 30% response compared to only a 4% response for those assigned to chlorpromazine. Clozapine was approved in 1990 on the basis of this study, and the US Food and Drug Administration kept “failing 2 antipsychotics of a different class” as the main criterion for starting clozapine treatment. This criterion of “failing 2 or more antipsychotics” remains the standard definition of treatment-resistant schizophrenia to this day (see Figure 1A for a visual depiction of treatment resistance circa 1990).

For the next 4 years after its approval in 1990, clozapine was the only “atypical” antipsychotic available. Given the dearth of other options at the time, a pressing question was whether clozapine was more effective than the other available medications for less gravely ill patients with schizophrenia. A predecessor to the current study was done when clozapine was the sole atypical antipsychotic, so haloperidol was chosen as a standard-of-care comparison to clozapine. Under double-blind conditions, “moderately refractory” patients were randomly assigned to a 6-month course of clozapine or haloperidol, the prevailing standard of care. Clozapine was indeed much more effective than haloperidol, with over 50% of clozapine-treated subjects meeting response criteria, which was twice the response seen with haloperidol. Clozapine also seemed more acceptable than haloperidol, with about two-thirds of clozapine-treated patients completing 6 months compared to about one-third for haloperidol-treated patients.

If clozapine had remained the only atypical antipsychotic between then and now, clozapine would be the clear treatment of choice for the wide range of patients encompassing the “moderately refractory” category. Of course, many other antipsychotics have been approved since clozapine. All of the post-clozapine antipsychotics were influenced by clozapine. We have at least 12 other atypical antipsychotics approved after clozapine. Now, let’s turn to the clozapine versus risperidone study published in this issue. As mentioned, this was the second of 2 studies of clozapine in “moderately refractory” schizophrenia, and followed the clozapine versus haloperidol study just discussed. The major difference from

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A. Circa 1990 treatment resistance as defined by clozapine selection and response criteria

<table>
<thead>
<tr>
<th>Response category</th>
<th>Medication by category</th>
</tr>
</thead>
<tbody>
<tr>
<td>“Responder”</td>
<td>Any other antipsychotic</td>
</tr>
<tr>
<td>Treatment-resistant</td>
<td>Clozapine</td>
</tr>
</tbody>
</table>

The non-clozapine antipsychotic options were all by definition from the class of conventional antipsychotics. Original study was for the most severely resistant patient group.

B. Circa 1995 revised subcategories of treatment resistance and relative antipsychotic response

<table>
<thead>
<tr>
<th>Revised response category</th>
<th>Treatment-resistant</th>
</tr>
</thead>
<tbody>
<tr>
<td>“Responder”</td>
<td>Moderately resistant</td>
</tr>
<tr>
<td>Clozapine</td>
<td></td>
</tr>
</tbody>
</table>

In this moderately resistant subgroup, clozapine was more effective than haloperidol, but risperidone was almost as effective as clozapine. Therefore, alternatives to clozapine can be effective in the middle group of “treatment resistance.”

C. Circa 2016 treatment-resistance concept as a gradient of relative pharmacologic nonresponse

The likelihood of unique response to clozapine depends on the patient’s position along a gradient. The clinical implication is that patients should be informed of clozapine as soon as some treatment resistance is observed, but alternatives may be as effective as clozapine for part of the treatment-resistant gradient, yet clozapine is uniquely effective at the far end of the nonresponse gradient.

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the prior study was the choice of risperidone rather than haloperidol to address the urgent question of whether another atypical antipsychotic besides clozapine can also be useful for the “moderately refractory” patient. Why risperidone rather than any other post-clozapine antipsychotic? The simple answer is that risperidone was approved in 1994 and was the first to arrive after clozapine and was available when this study started. I prefer to think of risperidone as a proxy for the other post-clozapine antipsychotics as well. If the results for risperidone were as grim as those for haloperidol, then clozapine was likely to remain the only realistic option across the entire spectrum of treatment-resistant schizophrenia.

The results of the current clozapine versus risperidone study by Schooler et al. tell a very different story than its predecessor study of clozapine versus haloperidol. Over the 6 months of treatment, patients taking clozapine were somewhat more likely to respond than patients taking risperidone (71% for clozapine and 57% for risperidone), but the difference was not statistically significant. Although the overall likelihood of finishing the entire 29 weeks was about the same for both medication conditions, the reasons differed. Patients were more likely to stop risperidone early due to insufficient response (38% for risperidone vs 15% for clozapine; *P* < .05); however, patients taking clozapine were...
more likely to stop early because of tolerability problems. The finding that risperidone-treated patients had over a 50% chance of improving was a very encouraging result.² Compared to haloperidol-treated patients from the prior study,³ risperidone-treated patients were twice as likely to achieve the same response criteria as those treated with haloperidol.² So, in the present study of clozapine versus risperidone,² clozapine retains (just barely) its title as being more effective for positive symptoms for “moderately refractory” patients, but risperidone was a close second. Put another way, clozapine is no longer the clear winner for the “moderately refractory” category. The results of the 2 studies together provide very valuable information on the sequencing and relative benefits and risks of clozapine across at least 2 categories of treatment resistance. Given the challenges of using clozapine, most patients in this category would probably choose risperidone before moving on to clozapine. However, some patients are so distressed by their symptoms that they would choose a clozapine trial sooner rather than later. Figure 1B illustrates the combined findings. When clozapine is chosen, it is important to be patient. The time course for response from clozapine is considerably longer than it was for risperidone. Risperidone response came earlier, so six weeks into treatment, the response rates were about the same. After 29 weeks, when the study ended, response to clozapine surpassed that of risperidone.

Failing 2 Antipsychotics Is No Longer Useful for Defining Treatment Resistance

The study by Schooler and colleagues² shows that risperidone can sometimes match clozapine for some patients along a nonresponse gradient. This finding makes the current way treatment resistance is categorized harder to use as the best method to select appropriate clozapine patients. There is reason to think that at least some of the 12 post-clozapine atypical antipsychotics can also be effective across a gradient of treatment resistance. But our definition of treatment resistance as a way to decide on clozapine selection has not kept up. It is time to abandon the criterion of failing 2 antipsychotic trials as a proxy for treatment-resistant schizophrenia.

The evolving nature of “failing” antipsychotic trials would not matter if the relative efficacy of clozapine and other non-clozapine options did not depend on the degree of treatment resistance. If clozapine always was the best choice irrespective of the degree of treatment resistance, then changing the threshold would not change the relative risks and benefits of choosing between clozapine and other antipsychotics. While clozapine is effective for moderately resistant patients, it is not unique in this segment of treatment resistance. Other post-clozapine antipsychotics may be as effective but less risky. However, clozapine continues to be the clear winner when treating the most severe end of treatment-resistant schizophrenia.⁵ Therefore, the relative benefits and safety of clozapine depend on where the patient is along a treatment-resistance gradient. Figure 1C illustrates the relative usefulness of clozapine across the gradient of treatment resistance.

It has been almost 30 years since the publication of the major clozapine treatment-resistant study.³ In 1988, there were no atypical antipsychotics. At the time of Study 30,⁴ failing 2 antipsychotics was a proxy for the extreme end of the treatment-resistance spectrum. Today, by comparison, the usual patient who is considered to have failed 2 antipsychotics is not as ill as that and is more likely to fall under the “moderately refractory” umbrella. The relative risks and benefits of subsequent medication choices will depend on where the patient is along the nonresponse gradient. Therefore, clozapine criteria should be updated to account for the relative degree of nonresponse across a gradient. Patients who are somewhere in the middle range of treatment resistance have different response patterns to other, non-clozapine atypical antipsychotics.⁶ Clozapine will retain its position at the end of the gradient as the single most effective medication for persistent positive symptoms (as well as for suicide risk). In the middle of the nonresponse gradient, offering clozapine is still important, but there is at least a reasonable chance that other non-clozapine options would also be helpful.

Submitted: March 8, 2016; accepted March 8, 2016.

Potential conflicts of interest: Dr Weiden has received grants and support from multiple pharmaceutical companies but has determined that they pose no conflicts relevant to this commentary on clozapine.

Funding/support: None.

REFERENCES