Recent reports have indicated that antipsychotic drugs exert their clinical effects earlier than previously thought, with the largest treatment effect occurring within 2-4 weeks, which has challenged the concept of delayed onset of action of antipsychotics. However, none of these studies investigated time to response in treatment-resistant schizophrenia. This is clinically important, since it is known that such patients constitute about a third of all cases. Moreover, these individuals are often subjected to multiple trials of antipsychotic drugs, including clozapine, without clear guidance as to the trial duration that could be considered adequate prior to moving to the subsequent strategies. For instance, time to response remains a controversial issue even with clozapine, a gold standard antipsychotic medication in treatment-resistant schizophrenia. At least one shorter-term investigation has demonstrated that a degree of improvement in Brief Psychiatric Rating Scale (BPRS) scores at 1 week can predict favourable outcome at 5 weeks. Further, some longer-term studies are not supportive of treatment response beyond 6-8 weeks. Nevertheless, another report showed that 55% of treatment response occurred after 6 weeks in individuals treated for a mean of 10 months (s.d. = 8). In addition, Meltzer et al reported that 20 of 36 participants classified as responders (response defined as more than 20% decrease in BPRS score) were identified after 6 weeks of treatment with clozapine. A majority of treatment response (70%; n = 14) was also attained later than 6 weeks if a threshold of response was set more stringently (i.e. a 50% or more decrease in BPRS score). Moreover, Zito et al reported that although 19% of patients exhibited improvement at 6 weeks, the rate response increased to 29% at 12 weeks. Indeed, one guideline recommends treatment to be continued for at least 8 weeks. However, clinicians have no guidance on when to consider a clozapine trial as failed, a question of major clinical relevance considering the adverse effects profile of this medication. We therefore conducted a systematic review of the literature in treatment-resistant schizophrenia to address response rates and time-line to response, focusing specifically on early improvement in this population.

Method

A literature search was conducted systematically to identify randomised, double-blind clinical trials of antipsychotics in adult...
patients with treatment-resistant schizophrenia. PubMed, Ovid Medline (since 1950), EMBASE (since 1980) and PsyCINFO (since 1967) through June 2010 were searched with key words 'treatment resistant schizophrenia' AND 'double blind'. A search was also conducted by replacing 'treatment resistant schizophrenia' with 'refractory schizophrenia'. Studies with fewer than ten adult patients in the treatment arms, studies for which response rate was not reported and dose-finding studies were excluded. Likewise, augmentation studies after run-in antipsychotic treatment were excluded.

Qualitative analysis
Each study was examined in terms of definition of treatment resistance. Definition of treatment response, rate of response, premature attrition rate and data reporting points were also investigated, to give an indication of time to antipsychotic response in treatment-resistant schizophrenia.

Statistical analysis
To investigate potentially early treatment improvement or onset of action, publications that reported at least three last observation carried forward (LOCF) data points in the Positive and Negative Syndrome Scale (PANSS; each item scored 1–7) or BPRS (each item scored 1–7) up to the first 6 weeks of treatment were extracted and mean item scores calculated by dividing the total score by 30 or 18 respectively. This method is in accordance with a previous investigation of early response to antipsychotics in non-resistant schizophrenia, and allows for the relative scarcity of available data points (see Results). The LOCF scores during the first 6 weeks of treatment were either reported as absolute scores in the original article (1 study), obtained by the authors on request (1 study), or determined from the accompanying figures detailing change over this period (3 studies) in the studies included in our meta-regression analyses. Data solely concerning clozapine were considered separately. Also, data on second-generation antipsychotics (other than clozapine) and study completer groups were investigated for descriptive purposes only.

A series of meta-regressions were carried out (via weighted mixed models analyses) using SAS version 9.1.3. for Windows to establish the time course of treatment response among participants with treatment-resistant schizophrenia. These models make less restrictive assumptions than many standard statistical tests (i.e. we need not to assume that our residuals are independent when using mixed models) and are appropriate for use when studying longitudinal data. Full maximum likelihood estimation was selected as the method of estimation for each of these analyses, and a variety of potential residual covariance structures (variance components, first-order autoregressive, compound symmetric, heterogeneous first-order autoregressive, heterogeneous compound symmetric, and unstructured) were investigated in each model. Although the structure and interpretation of the residual covariance matrix are of little clinical interest and do not relate directly to our research questions, we chose to investigate this series of modelling assumptions in order to identify the models and series of assumptions that best fit our data. This in turn improved the precision of our estimates (i.e. reduced the magnitude of standard errors in the estimates), which affects hypothesis testing related to our research questions.

To determine objectively which of these structures most accurately represents the true residual covariance structure within each of our investigations, we chose to examine the Bayesian information criteria (BIC) associated with each model – a goodness-of-fit statistic that penalises the model's likelihood function based on both model complexity (the number of parameters requiring estimation) and sample size. Use of this statistic allows us to identify the most parsimonious model (the model requiring estimation of the fewest parameters) which best fits our sample data. A smaller BIC indicates a better fit when comparing models, and the covariance structure ultimately adopted was the one that yielded the smallest BIC value for each of our analyses.

As one cannot reasonably assume independence of results across study arms within the same study, the findings for each of the two drug treatment groups within a given study were pooled to reflect a single set of results for each study, using a weighted average of the standardised scores across the two drug groups at each time point. Baseline sample sizes (pooled across treatment arms) for each study were used as sample weights in all analyses.

Percentage reduction in PANSS scores was obtained by taking into account the non-deductible value of the scale (i.e. 30), for instance, if the baseline mean item score was 4.0 (120 in total), a 20% reduction was defined as a post-treatment mean item score of 3.4, i.e. a total score of 102; note that (102 – 120)/(120 – 30) equals –0.2.

Results
The literature search using the key words 'treatment resistant schizophrenia' and 'double blind' located 14 studies and a replacement of the former search term with 'refractory schizophrenia' found five studies (Fig. 1). In total, 19 manuscripts reporting response rates following antipsychotic treatment of treatment-resistant/refractory schizophrenia in adult patients were identified (online Tables DS1 and DS2). The definition of treatment resistance varied among studies, although it invariably included past failure to respond to one or more antipsychotics at chlorpromazine equivalent doses of 400–1000 mg. Similarly, the definitions of treatment response were not necessarily consistent, although they universally adopted a specified degree of improvement in the representative rating scales (typically a 20–30% decrease in PANSS or BPRS score).

Study duration varied from 4 weeks to 1 year, with the majority (63%) lasting for 12 weeks or less (12 of 19 studies). In the light of these differences it is not surprising to note that both response rates and premature attrition rates were variable (0–76% and 2–72% respectively). A total of 11 antipsychotics were included in these studies, with the most frequently studied being clozapine (ten studies), haloperidol (seven), chlorpromazine (five), olanzapine (five) and risperidone (four). Only nine reports (47%) described data on key outcomes throughout available points; for the remainder, data were either unclear regarding assessment points, were selectively reported or were not fully provided throughout the treatment period (e.g. pre- and post-treatment presentation only). Only one study explicitly reported the rate of response through week 6 (excepting week 5). Results of this study showed a continuous increase in the percentage of responders (observed cases) for aripiprazole, whereas the slope of increase was more gradual for perphenazine after 3 weeks. In contrast, LOCF data from an intent-to-treat (ITT) sample implied the largest decrease in average PANSS scores in the first 3–4 weeks.

Meta-regression analysis
Studies included
One complex study that adopted a multiple transition among antipsychotics was excluded from the meta-regression analysis.
of these studies were not stratified by resistance or intolerance. With the exception of one study that described continuous data on BPRS four key items and an anergia item, positive v. negative symptoms data were either presented as pre- and post-treatment data, or not presented separately. Affecrive or cognitive symptoms were not addressed in these studies. All studies adopted a flexible dosing design, and the PANSS was used by all studies except for one that used the BPRS.

Analysis

Considering all antipsychotics, a first-order autoregressive covariance structure was found to yield the best fit among the various structures tested (five studies, n = 1019). Standardised mean item scores were found to change significantly over the course of the 6-week period (F = 34.68, d.f. = 6,22, P < 0.0001). Bonferroni-adjusted pairwise comparisons found that the baseline scores were significantly higher than the scores observed at all other time points (adjusted P < 0.0001). Week 1 standardised mean item scores were significantly higher than those observed at all subsequent time points (P < 0.0001 in each case). Week 2 scores were also significantly higher than those observed at weeks 3–6 (P < 0.0001). Week 3 scores were significantly higher than scores at weeks 4–6 (P = 0.00268, 0.0086 and 0.0004 respectively). Week 4 scores were significantly higher than scores at week 6 (P = 0.0172) but did not differ from week 5 scores (P = 0.6909). Scores at weeks 5 and 6 were not significantly different from each other (P = 0.7679). The mean standardised scores at each time point are summarised in Fig. 2.

For clozapine, a heterogeneous first-order autoregressive covariance structure was found to yield the best fit among the various structures tested (four studies, n = 356). Standardised mean item scores were found to change significantly over the course of the study (P = 90.91, d.f. = 6,18, P < 0.0001). Bonferroni-adjusted pairwise comparisons found that the baseline scores were significantly higher than the scores observed at all other time points (baseline v. week 1 P = 0.0177; baseline v. weeks 2–6 P < 0.0001). Week 1 scores were significantly higher than those observed at all subsequent time points (P < 0.0001 in each case). Week 2 scores were also significantly higher than those observed at weeks 3–6 (P < 0.0001). Week 3 scores were significantly higher than scores at weeks 4–6 (P = 0.0007, < 0.0001 and < 0.0001 respectively). Week 4 scores were significantly higher than scores at weeks 5 and 6 (P = 0.0002 and < 0.0001 respectively). Week 5 scores were also significantly higher than week 6 scores (P = 0.0003). The mean standardised scores at each time point are summarised in Fig. 3.

The mean standardised item scores were 3.4 for all antipsychotics and 3.5 for clozapine, corresponding to a PANSS score of 103. This baseline score was higher than that previously reported for patients with non-resistant schizophrenia by about 10 points. Overall, a 20% decrease was noted at week 4, and a 30% decrease was not achieved by the end of week 6 for all antipsychotics pooled. In contrast, for clozapine alone a 20% decrease was noted at week 3 and this increased further to 30% by week 6. An improvement of 0.17 for all antipsychotics (29% of the improvement from baseline through week 6) and 0.27 for clozapine (33%) was noted in weeks 4–6. Thus, approximately two-thirds of the improvement observed in 6 weeks occurred within the first 3 weeks. Nevertheless, a smaller improvement over the following 3 weeks corresponded to a further 5.1 point decrease in PANSS scores for all antipsychotics and 8.1 points for clozapine.

The results for second-generation antipsychotics other than clozapine (four studies, n = 380) and for completor-only samples (two studies, n = 233) are shown in online Figs DS1 and

<table>
<thead>
<tr>
<th>187 records identified through Ovid Medicine, EMBASE and PsycINFO (to June 2010)</th>
<th>152 records identified through PubMed (to June 2010)</th>
</tr>
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</table>
| | Duplicate records removed
| 169 records assessed for eligibility |
| 155 records excluded: 62 other focus/treatment/diagnosis 38 reviews 36 augmentation therapy 7 open studies 7 dose-finding studies 3 studies of children/adolescents 2 studies with no definition of response or n<10 |
| 14 studies included in qualitative analysis |
| Replacement of search term ‘treatment resistant schizophrenia’ with ‘refractory schizophrenia’ found 5 additional studies |
| 19 studies |
| 14 studies excluded from meta-regression analysis 1 complex study with a multiple transition among antipsychotics 13 studies without three or more data |
| 5 studies included in meta-regression analysis |

Fig. 1 Literature search.
We were interested in whether improvement with antipsychotic medication might be seen relatively early in treatment-resistant schizophrenia, similar to the early improvement found in patients with non-refractory disease. It would have been desirable if published antipsychotic trials in this population had adopted a uniform definition of treatment resistance and treatment response; however, study definitions as well as trial durations varied considerably, resulting in limited comparability across studies. Adopting the threshold recently proposed by Leucht et al (a 25% or more decrease in the BPRS or PANSS) would be expected to result in lower response rates compared with the more typical 20% cut-off, provided that other absolute criteria were not used (e.g. a post-treatment CGI-S score of 3 or less, or a BPRS score of 35 or less). This highlights the importance of researchers reaching a consensus on what constitutes ‘treatment-resistant schizophrenia’ and ‘treatment response’ thereafter. Moreover, although an evaluation of a third antipsychotic trial for those who have already failed to respond to two prospective antipsychotic trials appears to be ideal, such studies are few, and none has specifically reported on the timing issue. This noted, we evaluated five studies providing at least three LOCF data points within a 6-week treatment period. Preliminary results indicated that over this interval the greatest treatment effect (two-thirds of the improvement) may occur in the first 3 weeks, with more gradual and attenuated improvement in the following weeks. Therefore, prolonging an antipsychotic trial dosed appropriately in a patient with ‘treatment resistance’ in the absence of response beyond this period might not yield robust benefit, although even attenuated later improvements can still be of clinical significance. Determining the time at which the first trial should be abandoned and a subsequent trial initiated constitutes critical data to guide therapeutics, a question that only recently has begun to be addressed. This is especially pertinent to patients with treatment-resistant or refractory disorder who are often exposed to high doses and extended antipsychotic trials in an attempt to elicit a clinical response.

The results of our study may argue for caution on a ‘wait and watch’ approach. However, until we know more about possible differences in trajectory of response in this particular patient subgroup, it would be premature to argue for discontinuation in the face of poor response after the first 3–6 weeks, particularly for clozapine. Moreover, ‘treatment-resistant’ patients clearly constitute a heterogeneous group, including both patients truly ‘resistant’ to high doses of antipsychotics (e.g. CPZeq 1000 mg) and patients with ‘refractory’ disorder who may be less severely ill but are unable to tolerate such high doses. In fact, it is possible that the initial steeper decline in clinical scores is primarily driven by the latter group. In this respect, resistant and intolerant cases may need to be described separately, although most of the past studies have failed to do so.

**Limitations of the study**

Limitations of this investigation include the small number of studies available for analysis as well as the heterogeneous and non-standardised nature of the sample, although all patients were classified as having treatment-resistant/refractory schizophrenia according to each study. Specifically, a paucity of available data points precluded quantitative analysis and within-patient trajectory analysis of response, as well as examination of dose response for individual antipsychotics. The latter is particularly pertinent to clozapine with regard to initial titration, final dose and drug level at steady state. More consistent improvement with clozapine over 6 weeks may be a product of more gradual titration with this medication at an earlier treatment stage. Further, some data required extrapolation from figures accompanying the published reports, although a past work was obliged to use this strategy as well. We restricted the search to double-blind studies, which is consistent with previous work involving samples with non-refractory disorder, but limited the number of studies analysed. A distinction between completer and ITT results is important: LOCF data were available for five studies, whereas only two provided data for completers. The use of LOCF, a standard method in clinical trials to deal with participants leaving the study at various time points in the follow-up period of an intervention study, represents a compromise by including data from both completer and

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**Discussion**

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important issues in the management of patients who are treatment, whereas longer-term investigations can evaluate the delayed onset of antipsychotic action specifically for clozapine. Non-refractory patients, but does not necessarily rule out a than 6 weeks suggests that the greatest treatment effect is within the treatment period. Limited evidence from studies longer than 6 weeks suggests that the greatest treatment effect is within 3–4 weeks. This is in line with what has been reported for non-refractory patients, but does not necessarily rule out a delayed onset of antipsychotic action specifically for clozapine. Future studies need to monitor response earlier in the course of treatment, whereas longer-term investigations can evaluate the value of early response in predicting later outcome. It may well be that there are different trajectories of response, and distinguishing these will take us a step forward in defining subgroups, information that would have important clinical implications.

**Future research**

To better address the time-line issue it is desirable that future studies include longitudinal data on response rates as established by a priori criteria, e.g., 20%, 30%, 50% or 25% (in accordance with Leucht et al.) at each available assessment point, the average score of the primary outcome across time, median/mean time to response among responders, and time-line by which a certain percentage of patients are classified as responders, to allow for a better understanding of a reasonable antipsychotic trial for challenging patients in clinical practice. The data are better presented with ITT populations and observed cases at each assessment. Finally, such data can be interpreted in the context of the value of early improvement in predicting later response and the trajectory in response as well as identifying persistence on response status on the individual level, two important issues in the management of patients who are considered to have treatment-resistant disorder.

To conclude, definition of treatment-resistant schizophrenia and treatment response needs to be carefully delineated, in combination with more detailed data presentation throughout the treatment period. Limited evidence from studies longer than 6 weeks suggests that the greatest treatment effect is within 3–4 weeks. This is in line with what has been reported for non-refractory patients, but does not necessarily rule out a delayed onset of antipsychotic action specifically for clozapine. Future studies need to monitor response earlier in the course of treatment, whereas longer-term investigations can evaluate the value of early response in predicting later outcome. It may well be that there are different trajectories of response, and distinguishing these will take us a step forward in defining subgroups, information that would have important clinical implications.

**References**


10. Meltzer HY. Duration of a clozapine trial in neuroleptic-resistant schizophrenia. Arch Gen Psychiatry 1989; 46: 672.


