Antipsychotics: is it time to introduce patient choice?
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Summary
Evidence regarding overestimation of the efficacy of antipsychotics and underestimation of their toxicity, as well as emerging data regarding alternative treatment options, suggests it may be time to introduce patient choice and reconsider whether everyone who meets the criteria for a schizophrenia spectrum diagnosis requires antipsychotics in order to recover.

Effectiveness of antipsychotics
Recent evidence from systematic reviews and meta-analyses suggests that the efficacy and effectiveness of antipsychotics to produce clinically meaningful benefits for people with psychotic disorders have been overestimated. A meta-analysis showed that although there may be demonstrable effects of antipsychotics in comparison with placebo, the improvements over placebo are not as great as previously thought: the average change in symptoms rated with the Positive and Negative Syndrome Scale (PANSS) attributable to antipsychotics did not meet an empirically derived threshold for minimal clinical improvement, and only 17–22% experienced an important benefit (significant improvement or prevention of relapse) which could be attributed to the drugs rather than to placebo effects or natural recovery. A subsequent systematic review concluded that the improvements claimed for antipsychotics, old and new, are of questionable clinical relevance, with most trials failing to demonstrate even minimal improvement measured using the PANSS. There is also growing recognition that there is no discernible difference in effectiveness between first- and second-generation antipsychotics, supported by evidence from a recent meta-analysis. It is also relevant that there is evidence from double-blind trials in healthy volunteers that antipsychotic medication can result in secondary negative symptoms.

Adverse effects of antipsychotics
There is also evidence, again from systematic reviews and meta-analyses as well as from large controlled studies, to suggest that the adverse effects of antipsychotics have been underestimated. For example, a recent systematic review concluded that some of the structural abnormalities in brain volume previously attributed to the syndrome of schizophrenia may be the result of antipsychotic medication. There is also considerable evidence that antipsychotics are associated with an increased risk of sudden cardiac death, and that some of the increased mortality observed in people with a diagnosis of schizophrenia is attributable to antipsychotic medication.

Risk–benefit ratios, informed choice and collaborative decisions
Given that mental health services appear to have overestimated the strength of the evidence base for antipsychotic medication, while underestimating the seriousness of the adverse effects, it seems sensible to re-evaluate the risk–benefit ratio of such drugs. This risk–benefit profile may be a factor in the high rates of non-adherence and discontinuation of medication found in patients with psychosis; thus, some decisions to refuse or discontinue antipsychotic medication may represent a rational informed choice rather than an irrational decision due to lack of insight or symptoms such as suspiciousness. Given accurate and honest assessments of both risks and benefits, it should be possible to prescribe antipsychotics in a more thoughtful and collaborative way, and these considerations should involve explicit discussion of the possibility of not prescribing at all. Provision of such choices may help to engage people who might otherwise...
To facilitate informed choice and decision-making, we require a much better evidence base to help address questions such as how and when medication might be required, who is most likely to respond and what alternatives exist. There is some evidence for different trajectories of response, with a small proportion of patients demonstrating a rapid and dramatic favourable response to antipsychotics, but more research is clearly required to inform our ability to predict those most (and least) likely to respond to antipsychotics. Shorter duration of untreated psychosis has been shown to be a predictor of response to antipsychotics, which could be employed as an argument against offering no medication as a choice. However, any additional benefits of early treatment would still need to be evaluated against the long-term risks, and the traditional assumption that ‘untreated psychosis’ can only be treated by prescribing antipsychotics (and therefore not by psychosocial therapies) has yet to be comprehensively tested. It is relevant to this assumption that 20-year outcome data from the Chicago Follow-Up Study suggest that service users who decide not to take antipsychotics (often against medical advice) do relatively well, if not better, in comparison with service users who take such medication continuously.

In addition to research regarding predictors of response to antipsychotics, research is also required to inform evidence-based alternatives to antipsychotic medication, since the most likely candidates (such as psychosocial treatments including cognitive therapy and family interventions) have almost exclusively been evaluated as an adjunct to medication. There are a few exceptions, such as a recent trial of cognitive therapy for people who chose not to take antipsychotics; however, more clinical trials with greater methodological rigour are clearly needed.

It may be time to reappraise the assumption that antipsychotics must always be the first line of treatment for people with psychosis; rather, this should be a collaborative decision that is balanced with provision of informed choices and the offer of evidence-based alternatives. These decisions should be negotiated with service users on the basis of the likely positive and negative consequences and the prioritisation of their goals and values; such a collaborative approach might also result in better response for those who choose to take antipsychotics, since the quality of relationship with the prescribing clinician is associated with attitudes and adherence to medication.

**References**


