Antipsychotic polypharmacy seems, it seems, will not just go away. Two papers in this issue re-emphasise that antipsychotic polypharmacy is widespread, poorly supported and very probably dangerous.1,2 Yet the practice continues and is resistant to all kinds of quality improvement interventions.3

Polypharmacy – is it ever justified?

In any area of medicine, the practice of polypharmacy arises as a result of the failure of single-drug regimens. Examples include the treatment of hypertension, Parkinson’s disease, tuberculosis and HIV infection. In multi-episode schizophrenia, treatment failure is commonplace, with only a small proportion of patients showing a marked response to a single antipsychotic – a placebo-adjusted response rate of less than 20% is not unusual.4,5 Adding another antipsychotic is one way of ‘doing something’ (or appearing to do something) to improve on this mediocre response. However, whereas the use of polypharmacy in, for example, tuberculosis has a rational pharmacological basis and a solid clinical evidence base, antipsychotic polypharmacy might be said to have neither.

Some facts about antipsychotics . . .

What do we know for certain about antipsychotics? They work acutely for some people and prevent relapse for others; they cause adverse effects for most people; they share an ability to modify central dopaminergic transmission; clozapine is more effective than other antipsychotics. This is the sum of what is certain. Everything else (e.g. other neurotransmitter involvement, negative symptom response, cognitive changes) is a subject of disagreement, conjecture and, one might say, some confusion.

. . . and popular beliefs

A good example of the type of thinking that arises from this confusion is the idea, often heard expressed on ward rounds, that increasing dopamine D2 blockade in people taking clozapine will in some way improve response. This well-meaning practice survives alongside a body of evidence which suggests that this is not the case. People on clozapine will already have been subjected to multiple antipsychotics and polypharmacy, often at high dose, and failed to respond.6 Also, individuals switched from a long-acting injectable antipsychotic to clozapine show no difference in response characteristics to those switched from oral antipsychotics.7 Clozapine saturates D2 receptors in extrastriatal areas at normal clinical doses8 and clinical trial evidence either suggests no advantage for clozapine-antipsychotic co-therapy,9 or a minute effect not in any way clearly linked to D2-related activity.10 A Cochrane review suggests that clozapine augmentation remains of uncertain value and that more research is needed.11 Thus adding another antipsychotic to clozapine is, at least at the moment, of dubious benefit and any benefit seen is unlikely to be a result of increased D2 blockade.

Research evidence

Antipsychotic polypharmacy in general has a similar dearth of cogent support from the literature, notwithstanding the numerous individual clinical observations of clear benefit. A large meta-analysis including a number of

Summary

Polypharmacy is usually employed where single drugs are considered insufficiently effective. Some polypharmacy is rational and evidence based, some neither. Antipsychotic polypharmacy remains stubbornly widespread despite condemnation of the practice by numerous bodies. The practice could not be said to be evidence based. Its persistence probably stems from a well-meaning desire to improve response and from confusion about the mechanism of action of antipsychotics. In particular, the concept that more antipsychotic(s) must always be, or might be, ‘better’ is virtually groundless. Nonetheless, some specific antipsychotic polypharmacy regimens have shown particular benefits on adverse effect profiles. Targeted, evidence-based polypharmacy may be the way forward.

Declaration of interest

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Antipsychotic dose–response: guidelines v. practice

Prescribers seem also to be confused about the dose–response relationship of antipsychotics, as practice seems to suggest that many believe that ‘more is better’. This is true up to a point, but the ‘point’ is not where you might think it is. The most efficacious dose of risperidone is 4 mg a day,24 for aripiprazole 10 mg a day,22 for haloperidol 5 mg a day,16 for quetiapine 300 mg a day27 and for haloperidol decanoate 100 mg a month.28 The largest fixed-dose study to date showed olanzapine 10 mg to be just as effective as 20 mg and 40 mg a day.29 These observations of a low ceiling of effect suggest that many believe that ‘more is better’. This is true up to a point, but the ‘point’ is not where you might think it is. The most efficacious dose of risperidone is 4 mg a day,24 for aripiprazole 10 mg a day,22 for haloperidol 5 mg a day,16 for quetiapine 300 mg a day27 and for haloperidol decanoate 100 mg a month.28 The largest fixed-dose study to date showed olanzapine 10 mg to be just as effective as 20 mg and 40 mg a day.29 These observations of a low ceiling of effect tie in nicely with receptor occupancy studies suggesting saturation of receptors at low doses.30 Thus, more is not better once a certain dose is reached, at least with antipsychotics used as single agents. To then assume that adding a second antipsychotic (very probably with an identical mode of action) will bring about improvement is, some might say, a leap of faith beyond reason and logic.

Nonetheless, antipsychotic polypharmacy can represent logical and advantageous prescribing practice in certain instances. For example, when switching from one antipsychotic to another, cross-tapering of antipsychotics seems entirely sensible. There is also a modicum of support for the use of as needed (p.r.n.) antipsychotics (in addition to regular antipsychotics) in rapid tranquillisation.38 Perhaps more intriguingly, although adding aripiprazole to clozapine does not improve efficacy, it does cause patients to lose weight and may also improve other metabolic parameters.32 Co-therapy with aripiprazole and haloperidol has been shown to normalise prolactin levels in those formerly treated with haloperidol alone.33 Aripiprazole’s very high affinity for D2 receptors34 provides a partial explanation for these effects and both practices represent rational prescribing likely to be of benefit to patients.

Need for more research

Clearly, confusion will continue to reign until robust clinical trials are conducted to establish the merits or otherwise of antipsychotic polypharmacy. However, there is almost no financial impetus for studies of this type to be undertaken and, in any case, proof that a particular combination has advantages over a single drug would tell us nothing about other combinations and other drugs. Moreover, the artificial clinical environment created for clinical trials might, as is always the case, tell us less than we might want to know about drug effects in the real clinical setting.

Conclusions

I was last asked to write an editorial for The Psychiatric Bulletin (the predecessor of The Psychiatrist) on antipsychotic polypharmacy in 2002.35 Since then, rates of antipsychotic polypharmacy seem not to have changed. What has changed is that evidence supporting antipsychotic polypharmacy has, if anything, diminished and evidence suggesting or demonstrating harm has grown. This mounting awareness of the probable futility of antipsychotic polypharmacy is reflected in the latest guidance issued by the National Institute for Health and Clinical Excellence (NICE).36 One has to hope that the audit processes demanded by NICE guidelines will at last go some way finally to reducing the extent of antipsychotic polypharmacy in UK mental health units.

About the author

David Taylor is Chief Pharmacist at the South London and Maudsley NHS Foundation Trust, and Professor of Psychopharmacology at King’s College London, UK.

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