Short report

Ethnicity and quality of antipsychotic prescribing among in-patients in south London

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Summary

Ethnicity may influence treatment decisions in mental disorders. We undertook a survey of the prescribing of antipsychotics for in-patients in three south London mental health trusts. A total of 255 patients (152 White, 103 Black) were included. Median dose of antipsychotic (% of licensed dose) was 58.5% for White and 50.0% for Black patients (adjusted effect size=0.14, 95% CI –0.34 to 0.63). High-dose antipsychotics were prescribed to 15.1% of White and 11.7% of Black patients (adjusted odds ratio (OR)=0.5, 95% CI 0.19–1.33), and antipsychotic polypharmacy was recorded for 25.7% and 31.1% respectively (adjusted OR=3.05, 95% CI 1.44–6.46). Prescribing quality was similar for Black and White patients.

Declaration of interest

None. Funding detailed in Acknowledgements.

There have been suggestions of institutional racism in UK mental health services.1 Several, mainly American, studies indicate that Black patients are more likely than White patients to receive high doses of antipsychotics and depot formulations2–4 and less likely to be treated with atypical antipsychotics.5 Many of the studies are limited by the failure to collect and correct for other factors likely to affect prescribing practice. We published a single-centre study6 of antipsychotic prescribing which took into account over 20 potentially confounding factors, but found no difference in corrected odds of receiving high doses of antipsychotics or atypical polypharmacy. We wanted to know whether our results would differ in a larger study with greater power to detect smaller differences and which included participants from other trusts.

Method

This study was conducted at the South London and Maudsley, South West London and St George’s, and Oxleas National Health Service (NHS) Trusts during late 2006 and early 2007. We sought and obtained individual approvals for the study through local clinical audit channels.

Patients included were in-patients, designated Black or White, and prescribed and taking one or more regular antipsychotics. Patients were classed as Black if both parents were also Black (that is, Africans, African Americans and African–Caribbeans). Mixed-race patients were excluded. All suitable patients on all acute general psychiatry wards in every hospital within each trust were approached over a 3-month period in 2006/2007: none was excluded except for the reason above. The outcomes of dose (expressed as a percentage of licensed maximum),7 being prescribed antipsychotic medication above maximum dose, polypharmacy and costs were determined by reference to each patient’s medication chart and to standard reference texts for dose8 and cost.9

Potential confounding factors (23 in total) were predetermined and details were obtained from case notes, self-report, or by measurement or calculation and confirmed by nursing or medical staff where appropriate. Clinical Global Impression–Severity10 (CGI–S) was rated on the day of data collection (nurse or medical staff assessment).

A sample size of 298 was calculated to be required to give an 80% chance of detecting a 5% absolute difference for our main outcome (dose) (z=0.05, p=0.8). We aimed to compare four outcomes (dose, rate of polypharmacy, high-dose prescribing and use of atypical antipsychotics) between our two groups and to adjust comparisons for the effect of confounding (predictive) variables. For the outcome of dose we used a linear regression model to provide an estimate of unadjusted effect of ethnicity on dose. Potential confounding variables were then tested to identify predictive factors (significance level of P<0.1). Predictive factors were then included in a rerun regression model, producing adjusted effect size for ethnicity. Transformations were used when necessary. A similar approach was used for the binary outcomes of high dose, polypharmacy and prescribing of atypical drugs, but with logistic regression modelling used.

Results

We approached 300 patients and 255 gave informal informed consent to be interviewed. Of the 45 patients who declined to take part, 21 (46.7%) were Black, 23 (51.1%) were male and the mean age was 41.7 years. Details of included patients are given in the online Table DS1.

Median dose was 58.3% for White patients and 50.0% for Black patients (adjusted effect size=0.14, 95% CI –0.34 to 0.63; P=0.56). High-dose antipsychotics (>100% licensed maximum) were prescribed to 15.1% of White and 11.7% of Black patients (adjusted OR=0.5, 95% CI 0.19–1.33; P=0.16) and polypharmacy to 25.7% and 31.1% respectively (adjusted OR=3.05, 95% CI 1.44–6.46; P=0.004). With polypharmacy, the adjusted odds ratio was largely driven by centre differences: one centre showed an exceptionally high rate of polypharmacy in Black patients (74% v. 37% in White patients; other centres: 13% v. 17% and 16% v. 10% respectively). There was no difference in the prescribing of atypical antipsychotics (White 77.6%, Black 68.9%; adjusted OR=0.57, 95% CI 0.21–1.5; P=0.25).

Discussion

In this study, ethnicity was not significantly associated with dose of antipsychotic, the prescribing of high-dose antipsychotics or the use of atypical antipsychotics. Prescribing quality was thus no worse for Black patients than for White patients. Only the outcome of adjusted odds ratio for antipsychotic polypharmacy showed any association with ethnicity. This is an important observation but it should be noted that absolute rates of polypharmacy differed markedly in only one centre and the overall difference in prevalence was small (25.7% for White v. 31.1% for Black patients).
Our findings are therefore in some contrast to studies which suggest a higher likelihood of higher-dose prescribing in Black patients²–⁵ and a lower use of atypical drugs. This may reflect true differences in practice at different times (there is evidence of ethnic differences in prescribing in the 1990s in south London)¹¹ or in different locations (many of the studies examine US prescribing) but may also be linked to the relatively limited extent of adjustment for confounding variables in previous studies. Adjustment for these confounders is essential to the process of establishing or otherwise ethnicity as having an association with prescribing quality.

There were several limitations in our study design. We did not meet our recruitment target of 298 patients, although confidence intervals ultimately excluded major differences in outcome, particularly with respect to a possibility of a lower quality of prescribing for Black people. In addition, some of our data collection relied on patient self-report – a notoriously unreliable source, especially, perhaps, in psychiatric in-patients. Also, our assessment of clinic status was approximate (using the CGI–S scale) and may not of have been relevant to patients’ condition at the time of the initial prescription. Lastly, our sample was exclusively in-patients and so our results may not generalise to the majority of patients now treated in the community.

Notwithstanding these limitations, it is reasonable to conclude that in this study prescribing quality did not differ substantially between Black and White patients. Black patients were not prescribed higher doses than White patients. Black patients are more likely to receive antipsychotic polypharmacy, but this difference was only noticeably higher in one centre. Black patients were just as likely as White patients to receive atypical antipsychotics.

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References