Self-poisoning is a common method of suicide, especially in women. Antidepressants are frequently used for self-poisoning, being involved in around 20% of all poisoning suicides in the UK and in 20–30% of non-fatal overdoses. This reflects the facts that depression is the most frequent psychiatric disorder in people dying by suicide, the method used for suicidal acts is often determined by availability, and self-poisoning in individuals with depression often involves antidepressants prescribed for them. Relative toxicity is an important factor likely to determine the outcome of an antidepressant overdose. Studies using different approaches have shown wide variation in the relative toxicity of antidepressants, with the older tricyclic antidepressants (TCAs) generally being more toxic than the newer selective serotonin reuptake inhibitors (SSRIs). We have used two approaches to assessing the relative toxicity of classes of antidepressants and individual antidepressants. The first approach was to relate drug-specific poisoning mortality rates to prescription rates—the ‘fatal toxicity index’. The second, and generally less used approach, is to compare the rate of death with the rate of non-fatal self-poisoning, which generates a ‘case fatality’ index. The fatal toxicity index approach is probably less accurate because it is more heavily influenced by prescribing policies, including use of some antidepressants for conditions other than depression, and selective prescribing, for example, based on the clinician’s assessment of suicide risk i.e. ‘confounding by indication’. The specific aims of the study were to provide up-to-date information on the relative toxicity of individual antidepressants that may assist clinicians in making decisions about prescribing and inform interventions by regulatory authorities.

**Results**

Fatal toxicity and case fatality indices provided very similar results (rho for relative ranking of indices 0.99). Case fatality rate ratios showed greater toxicity for TCAs (13.8, 95% CI 13.0–14.7) than the SNRI venlafaxine (2.5, 95% CI 2.0–3.1) and the NaSSA mirtazapine (1.9, 95% CI 1.1–2.9), both of which had greater toxicity than the SSRIs (0.5, 95% CI 0.4–0.7). Within the TCAs, compared with amitriptyline both dosulepin (relative toxicity index 2.7) and doxepin (2.6) were more toxic. Within the SSRIs, citalopram had a higher case fatality than the other SSRIs (1.1, 95% CI 0.8–1.4 v. 0.3, 95% CI 0.2–0.4).

**Conclusions**

There are wide differences in toxicity not only between classes of antidepressants, but also within classes. The findings are relevant to prescribing decisions, especially in individuals at risk, and to regulatory policy.

**Declaration of interest**

D.G. is a member of the Medicines and Healthcare products Regulatory Agency’s Pharmacovigilance Expert Advisory Group.

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**Background**

Self-poisoning is a common method of suicide and often involves ingestion of antidepressants. Information on the relative toxicity of antidepressants is therefore extremely important.

**Aims**

To assess the relative toxicity of specific tricyclic antidepressants (TCAs), a serotonin and noradrenaline reuptake inhibitor (SNRI), a noradrenergic and specific serotonergic antidepressant (NaSSA), and selective serotonin reuptake inhibitors (SSRIs).

**Method**

Observational study of prescriptions (UK), poisoning deaths involving single antidepressants receiving coroners’ verdicts of suicide or undetermined intent (England and Wales) and non-fatal self-poisoning episodes presenting to six general hospitals (in Oxford, Manchester and Derby) between 2000 and 2006. Calculation of fatal toxicity index based on ratio of rates of deaths to prescriptions, and case fatality based on ratio of rates of deaths to non-fatal self-poisonings.
2000–2006 in England and Wales. In England and Wales it has
been customary to assume that the majority of injuries and
poisonings of undetermined intent are cases where the harm
was self-inflicted but there was insufficient evidence to prove that
the deceased intended to kill themselves.12,13 We have restricted
our analyses to deaths involving single drugs or single drugs and
alcohol. Data were obtained for males and females separately for
all drugs. Mortality rates were calculated as the number of deaths
per 100 000 population in England and Wales, for people aged 10
years and over.

Self-poisoning
Self-poisoning data came from three centres currently involved
in the Multicentre Monitoring of Self-harm project (see Hawton
et al6 for a description of the first phase of this project). Data were
collected on all people who presented with self-harm to emergency
departments at general hospitals in Oxford (one hospital),
Manchester (three hospitals) and Derby (two hospitals)
during the study period. Self-harm is defined as intentional
self-poisoning or self-injury, irrespective of motivation.14 Self-
poisoning includes the intentional ingestion of more than the
prescribed amount of any drug, or other non-ingestible substance,
whether or not there is evidence that the act was intended to result
in death. Data were collected on gender, age, date of self-harm and
method of self-harm, including which drug(s) were ingested in
overdoses. The analyses for non-fatal self-poisoning were based
on all non-fatal overdoses of the antidepressants, including those
with other drugs (with or without alcohol).

Episodes of self-poisoning involving the antidepressants for
people aged 10 years and over for the defined population areas
of Oxford City (mean annual population 128 411), Manchester
City (381 236) and Derby Unitary Area (204 032) were included in
the study. Mid-year population estimates for these areas for
2000–2006 were obtained from the Office for National
Statistics.15 Rates of self-poisoning per 100 000 population were
calculated for these areas combined.

Statistical analysis
Fatal toxicity index
Rate ratios for drug-specific poisoning mortality relative to
prescribing rates were calculated from the mortality rate
(numerator) and the prescription rate (denominator).

Case fatality
Rate ratios for poisoning mortality relative to non-fatal self-
poisonings for specific drugs were calculated from the mortality
rate (numerator) and the self-poisoning rate (denominator).

Relative toxicity indices were calculated by standardising the
rate ratios to amitriptyline. This drug is used as the reference
preparation in many studies.

Rate ratios and 95% confidence intervals, Spearman’s rho and
heterogeneity were calculated using Stata version 10.0 for
Windows.

Ethical approval
Oxford and Derby both have approval from local health/
psychiatric research ethics committees to collect data on self-harm
for local monitoring and multicentre projects. Self-harm
monitoring in Manchester is part of a clinical audit system, and
has been ratified by the local research ethics committees. All
monitoring systems are fully compliant with the provisions of the
Data Protection Act of 1998. All centres also have approval
under Section 251 of the National Health Service Act 2006
(formerly section 60 of the Health and Social Care Act 2001)
regarding the use of patient identifiable information.

Results
The total numbers of deaths, prescriptions and episodes of self-
poisoning are given in Table 1 (see online Table DS1 for data
stratified by gender). The antidepressants most frequently involved
in suicide deaths were (in order) dosulepin, amitriptyline,
venlafaxine and citalopram. For non-fatal poisonings the most
frequently involved antidepressants were (in order) fluoxetine,
citalopram, amitriptyline, paroxetine, venlafaxine, dosulepin,
sertaline and mirtazapine (Table 1). These patterns were similar
in both genders.

Fatal toxicity index (mortality to prescriptions ratio)
There was significant heterogeneity in fatal toxicity within the
TCA group ($\chi^2 = 365.02$, d.f. = 6, $P < 0.001$). The mortality to
prescriptions rate ratio was considerably higher for dosulepin
and doxepin than for amitriptyline (Table 2). This is reflected in
the toxicity ratios relative to amitriptyline for these drugs
dosulepin 3.2, doxepin 2.5). The fatal toxicity ratios were
generally far higher in males than females but the relative toxicity
indices were mostly similar in males and females.

The relative toxicity index for venlafaxine (0.46) was
approximately half that for amitriptyline but over five times
higher than that for the SSRIs (0.08). The relative toxicity index
for mirtazapine (0.32) was slightly lower than that of venlafaxine.

There was significant heterogeneity in fatal toxicity within the
SSRI group ($\chi^2 = 28.76$, d.f. = 4, $P < 0.001$). The fatal toxicity index
citalopram was three times higher than that for the other SSRIs
(1.7, 95% CI 1.3–2.3 v. 0.6, 95% CI 0.4–0.8), with a threefold
higher relative toxicity index (0.15 v. 0.05).

Case fatality (mortality to self-poisonings ratio)
There was significant heterogeneity in the relative toxicity within
the TCA group ($\chi^2 = 123.01$, d.f. = 6, $P < 0.001$). The mortality
to self-poisonings rate ratio was significantly higher for dosulepin
doxepin than amitriptyline (Table 3). The relative toxicity
indices for both dosulepin and doxepin suggest that these are
two to three times more toxic than amitriptyline. There was a
particularly high relative toxicity index for doxepin in females
and for trimipramine in males (although the 95% CIs for the rate
ratios were wide).

The relative toxicity index for venlafaxine (0.29) was much
lower than for the TCAs (1.6). However, it was considerably
greater than for the SSRIs (0.06). The relative toxicity index for
mirtazapine (0.22) was slightly lower than that of venlafaxine,
although no different in females.

There was significant heterogeneity in relative toxicity in the
SSRI drug group ($\chi^2 = 33.88$, d.f. = 4, $P < 0.001$). Citalopram had
a higher case fatality than the other four SSRIs (1.1, 95% CI
0.8–1.4 v. 0.3, 95% CI 0.2–0.4), which was reflected in a threefold
higher relative toxicity index (0.12 v. 0.04).

Comparison of fatal toxicity and case fatality
There was a very high correlation between the rankings of the
results of the fatal toxicity and case fatality approaches to
estimating relative toxicity of the specific antidepressants
(Spearman’s rho = 0.99, $P < 0.001$).
We have used two methods for assessing relative toxicity of antidepressants, the fatal toxicity index and the case fatality index. Both are subject to limitations. However, the case fatality index is probably a more accurate indicator of relative toxicity, being less influenced by selective prescribing. The findings based on both approaches show that dosulepin and doxepin are considerably more toxic than amitriptyline. Because of extensive prescribing of dosulepin relative to doxepin (Table 1), attention regarding toxicity has mainly focused on the former drug. In the UK in December 2007 the Medicines and Healthcare products Regulatory Agency issued advice regarding dosulepin and measures to reduce the risk of fatal overdose. Since November 2007 pack sizes have been limited and packaging made safer.15 Our findings support the need for such measures but also highlight the need to extend these to doxepin.

There were some gender differences, especially in relation to doxepin where relative toxicity based on case fatality was greater in females than males, and trimipramine where relative toxicity was greater in males. These differences may be the result of chance...
because of low numbers. They could also reflect differences in prescribing patterns (e.g. dosages) for the two genders.

Venlafaxine is clearly far less toxic in overdose than the TCAs. Its relative toxicity index based on both assessment approaches was intermediate between TCAs and SSRIs. In the UK the Medicines and Healthcare products Regulatory Agency issued a warning about the relatively high toxicity of venlafaxine in 2006, recognising that selective prescribing to individuals at risk of suicide could be a contributory factor to this finding. The regulatory body also issued prescribing advice, including restricting pack sizes and initial supplies for patients. In our study the relative toxicity of mirtazapine was slightly lower than that of venlafaxine, but considerably greater than for the SSRIs. It was intermediate between TCAs and SSRIs. In the UK the Medicines and Healthcare products Regulatory Agency issued a warning about the relatively high toxicity of venlafaxine in 2006, recognising that selective prescribing to individuals at risk of suicide could be a contributory factor to this finding. The regulatory body also issued prescribing advice, including restricting pack sizes and initial supplies for patients. In our study the relative toxicity of mirtazapine was slightly lower than that of venlafaxine, but considerably greater than for the SSRIs. Its relative toxicity index based on both assessment approaches, the findings of which were comparable with the SSRIs, may reflect known cardiotoxic and proconvulsant effects of citalopram in overdose. Although the absolute fatal toxicity is low, this finding should nevertheless be considered when making risk–benefit decisions regarding prescribing for individual patients.

Limitations and strengths of the study

One limitation is that data for different factors were derived from different areas: prescribing for the UK, deaths for England and Wales, and non-fatal poisonings for three local centres. We have made the assumption that they are all reasonably representative of the UK. This may be a particular limitation for the data on non-fatal self-poisonings, especially as the catchment areas used in this study are cities. We have commented elsewhere about differences in self-poisoning rates relative to prescribing patterns in the three centres. These were limited and unlikely to substantially bias our overall findings. The populations of all three centres include people from a wide range of socioeconomic backgrounds.

For fatal self-poisoning we have focused on single antidepressant overdoses (with or without alcohol), in order to avoid cross-contamination of the results with other drugs. However, certain antidepressants may be more toxic in overdose when combined with other drugs (e.g. doxepin and amitriptyline). Also, we have not been able to take account of the size of overdoses, which may be influenced by local prescribing practices and use of antidepressants for indications other than depression (especially when lower doses may be used). For non-fatal self-poisonings we included both episodes of single drugs and those involving other drugs. The non-fatal self-poisonings are based on episodes that resulted in presentation to hospital. There are likely to be other cases where self-poisoning does not result in hospital presentation, so that the case fatality rates may be overestimated. The prescription data we used were only available for all ages.

One factor that could influence the findings is differential use of specific antidepressants in overdoses, which is likely to reflect differences in indications used when they are prescribed. Elsewhere we have confirmed that this is the case for venlafaxine; which tends to be more often prescribed for, and taken in overdose by, people with longer-term psychiatric disorders and a history of self-harm.

The international applicability of the findings may be limited because of differences in regulatory advice, availability of specific antidepressants and prescribing practices between countries. Therefore country-specific and international comparative studies are warranted.

Implications

Of the TCAs, dosulepin and doxepin have the greatest toxicity when combined with other drugs (e.g. doxepin and amitriptyline). Also, we have not been able to take account of the size of overdoses, which may be influenced by local prescribing practices and use of antidepressants for indications other than depression (especially when lower doses may be used). For non-fatal self-poisonings we included both episodes of single drugs and those involving other drugs. The non-fatal self-poisonings are based on episodes that resulted in presentation to hospital. There are likely to be other cases where self-poisoning does not result in hospital presentation, so that the case fatality rates may be overestimated. The prescription data we used were only available for all ages.

One factor that could influence the findings is differential use of specific antidepressants in overdoses, which is likely to reflect differences in indications used when they are prescribed. Elsewhere we have confirmed that this is the case for venlafaxine; which tends to be more often prescribed for, and taken in overdose by, people with longer-term psychiatric disorders and a history of self-harm.

The international applicability of the findings may be limited because of differences in regulatory advice, availability of specific antidepressants and prescribing practices between countries. Therefore country-specific and international comparative studies are warranted.

A major strength of the study is that we have used two approaches to assessing relative toxicity, the findings of which are remarkably similar. The data on non-fatal self-poisoning have come from three well-established monitoring systems. Also, by using 7 years of data the study has considerable statistical power.
slightly more toxic than mirtazapine. Of the five SSRIs that we examined, citalopram appears to be more toxic than the other four. When prescribing antidepressants the clinician should take account of the risk that may be associated with an overdose, especially in someone judged to be at risk of self-poisoning, as well as relative efficacy, acceptability and possible interactions with other medication and alcohol, and concurrent physical morbidity. We suggest that the assessment of the relative toxicity of antidepressants should continue as new antidepressants are marketed and that international comparisons are warranted in view of differences between countries in prescribing practices.

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Keith Hawton, DSc, Helen Bergen, PhD, Sue Simkin, BA, Centre for Suicide Research, University of Oxford, Department of Psychiatry, Warneford Hospital, Oxford; Jayne Cooper, PhD, Centre for Suicide Prevention, University of Manchester, Manchester; Keith Waters, RMN, Derbyshire Mental Health Service NHS Trust, Mental Health Resource Centre, Rehabilitation Centre, Royal Derby Hospital, Derby; David Gunnell, PhD, Department of Social Medicine, University of Bristol, Bristol; Navneet Kapur, MD, Centre for Suicide Prevention, University of Manchester, Manchester, UK.

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References

**Data supplement**

**Table DS1** Deaths by suicide and undetermined intent in England and Wales, prescriptions in the UK and self-poisoning episodes in three centres in England involving each antidepressant, stratified by gender for people aged 10 years and over, for 2000–2006

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TCAs, tricyclic antidepressants; SNRIs, serotonin and noradrenaline reuptake inhibitor; NaSSA, noradrenergic and specific serotonergic antidepressant; SSRI, selective serotonin reuptake inhibitors.

a. Prescriptions include those for under 10 years old.
Toxicity of antidepressants: rates of suicide relative to prescribing and non-fatal overdose
Keith Hawton, Helen Bergen, Sue Simkin, Jayne Cooper, Keith Waters, David Gunnell and Navneet Kapur
Access the most recent version at DOI: 10.1192/bjp.bp.109.070219

Supplementary Material
Supplementary material can be found at:
http://bjp.rcpsych.org/content/suppl/2010/05/04/196.5.354.DC1

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