Adverse Effects of Antidepressants Reported by a Large International Cohort: Emotional Blunting, Suicidality, and Withdrawal Effects.

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Abstract:

Background: Studies of the adverse effects of antidepressants tend to focus on biological symptoms. The prevalence of suicidality and withdrawal effects are currently a source of controversy.

Objective: To directly ascertain the experiences of an international sample of antidepressant users.

Method: An online survey asked adult antidepressant users whether they had experienced 20 adverse effects ‘as a result of taking the antidepressant’, and if so, to what degree of severity. 1,431 people from 38 countries responded.

Results: 61% of the respondents reported at least ten of the 20 effects, most commonly: ‘Feeling emotionally numb’ (reported by 71%), ‘Feeling foggy or detached’ (70%); ‘Feeling not like myself’ (66%), ‘Sexual difficulties’ (66%), ‘Drowsiness’ (63%), and ‘Reduction in positive feelings’ (60%). ‘Suicidality’ as a result of the drugs was reported by 50%. Withdrawal effects were reported by 59%, and ‘Addiction’ by 40%. Rates of adverse effects were higher for those prescribed multiple antidepressants and those who also took antipsychotics. Younger age and longer use of ADs were positively related to total adverse effects. One third did not recall being told about any side effects by the prescriber. Less than 5% were told about suicidality, emotional numbing, withdrawal effects or addiction.

Conclusions: Asking people directly reveals far higher rates of adverse responses to antidepressants than previously understood, especially in the emotional, psychological and interpersonal domains. Given recent findings that antidepressants are only marginally more effective than placebo, the findings of the current study imply a cost-benefit analysis that cannot justify the extremely high prescription rates for these drugs.
Keywords: Adverse effects, antidepressants, depression, emotional numbing, side effects, suicidality, withdrawal effects.
1. INTRODUCTION

Prescription rates for antidepressants (ADs) are high and increasing [1]. One in eight adults in the USA were prescribed ADs by 2012 [2]. In 2016, England had 64.7 million prescriptions for a population of 55.3 million. This was double the prescriptions in 2006 and means that about one in seven women and one in 13 men are currently receiving antidepressants in England [3]. These extraordinary prescription rates are difficult to understand in terms of evidence-based cost-benefit analyses. Less than half of trials find ADs superior to placebo [4], with properly blinded studies particularly unlikely to find any difference to placebo [5, 6]. An early meta-analysis found that ‘the overall effect of new-generation antidepressant medications is below recommended criteria for clinical significance’ [7]. A recent meta-analysis [8] reached much more favorable conclusions but the median duration of treatment in the included studies was just eight weeks and studies were excluded if 20% or more of the participants were ‘treatment-resistant’ (i.e. it was known that the drugs did not work for them) were excluded. Furthermore the meta-analysis paid no attention to adverse effects. Another recent, more rigorous, meta-analysis [9], of 131 placebo-controlled trials, which did include adverse effects, again found that the overall effect size does not reach the threshold for ‘clinical significance’ and also found that ADs produce significantly more ‘serious’ and non-serious’ adverse effects than placebos. It was concluded that ‘The potential small beneficial effects seem to be outweighed harmful effects of SSRIs versus placebo for major depressive disorder seem to outweighed by harmful effects’ [9]. Of particular relevance to the current study was the finding that ‘There was almost no data on suicidal behavior, quality of life and long-term effects’.

Studies, and checklists, of these harmful effects have indeed focussed on biological or medical effects. Of the 84 adverse events reported by the 2017 meta-analysis to have been studied at least once, almost all were biological/medical reactions. The eight effects most
often assessed in the 131 drug trials were: nausea, headache, dry mouth, insomnia, somnolence, diarrhea, dizziness and constipation effects’ [9]. Both the ‘Antidepressant Side-Effect Checklist’ [10] and the ‘Patient-Rated Inventory of Side Effects (PRISE)’ [11] focus almost exclusively on these bio-medical phenomena and fail to address the psychological or interpersonal domains (although some, such as sexual impairment, can be seen as both biological and interpersonal).

A 2013 review [12] of the few, relatively small scale, studies of the actual experiences of AD users [e.g. 13-17] identified multiple adverse effects relating to quality of life, in the psychological, emotional and interpersonal domains. These included emotional detachment, a belief that ADs prevent natural sadness, harmful effects on relationships, caring less about self and others, fear of addiction, and suicidality. For example of 468 descriptions of adverse effects by AD users on a website the most frequent were sedation, impaired cognition, reduced libido, emotional blunting, activation (arousal, insomnia, agitation) and emotional instability [14].

The largest survey of AD users to date [18] identified surprisingly high rates of adverse effects among 1,829 New Zealanders, especially in the personal and interpersonal domains. These included: Sexual difficulties (62%), Feeling emotionally numb (60%), Feeling not like oneself (52%), Agitation (47%), Reduction in positive feelings (42%), Suicidality (39%), and Caring less about others (39%). It also identified high rates of Withdrawal effects (55%) and Addiction (27%).

The New Zealand survey reflects the growing interest in the views of ‘experts-by-experience’ in general and of Patient-Reported Outcome Measures (PROMs) in particular. These tools are increasingly recognized as an essential component of the governance of healthcare systems and the assessment of drugs, with a particular focus on understanding whether health care [19] and mental health care [20] are beneficial and safe. The current
study aimed, therefore, to expand our knowledge of the first-hand experience of people who take ADs by replicating the New Zealand study with a similarly large, but international, sample.

2. METHODS

2.1 Instrument

‘The Experiences of Anti-depressant and Anti-psychotic Medication Survey’ was developed for this study (https://swinuw.au1.qualtrics.com/jfe/form/SV_0jPOxpXhPLNZjmZ). It was in English, with no translations made available. The anti-depressant sections of the questionnaire were based on the New Zealand ‘Views on Antidepressants’ questionnaire [18, 21-23]. The questionnaire generated quantitative (yes/no and multiple-choice questions) and qualitative data (open-ended questions), about: the prescribing experience, the positive and negative effects of medications, causal beliefs about psychosis/depression, alternative treatments, experiences of withdrawing from the medications, and demographics,

The current paper reports the adverse outcomes of the AD section. Adverse effects were assessed with the question ‘Please rate the following side effects you may have experienced as a result of taking the anti-depressant’, followed by a list of 20 effects and the options to respond to each with ‘Not at all’, ‘Mild’, ‘Moderate’ or ‘Severe’ (see Table 2). Three of the 20 side effects in the original ‘Views on Antidepressants’ questionnaire were not included; two because they were the least frequently endorsed (‘Diarrhoea’ - 20% and ‘Weight loss’ - 15%), and one, ‘Failure to reach orgasm’, because it was covered by ‘Sexual difficulties’. These three were replaced with three new items that had been spontaneously reported by numerous participants: ‘Insomnia’, ‘Feeling foggy or detached’, and ‘Distorted dreams’ [18]. Participants were also invited to report ‘Other’ side effects. They were further asked (in a
Section titled ‘When you were first prescribed anti-depressant’) ‘Did the doctor inform you of any possible side effects?’ (Yes/No) and ‘If Yes, what side effects were mentioned?’

2.2 Participants

Of the 2,346 people who responded 668 were recruited via an Australian online research company, and 1,678 people via advertisements on social media and snowball sampling. Of the 2,133 who had taken ADs, 42 were deleted because they ticked ‘no’ when asked to confirm that they met the following criteria: ‘I have been taking or have previously taken antidepressant medication continuously for at least one month for any reason’; ‘I am aged 18 or older’; and ‘I am not currently compulsorily detained in a psychiatric hospital’.

Among the remaining 2,091 responses, the 44 that responded to ‘What is the name of your current or most recent anti-depressant medication?’ with a drug that is not an anti-depressant were deleted (most commonly valium - 16, xanax - 7 and lithium - 5). There were 39 responses emanating from the same Internet Protocol (IP) address as another response, indicating use of the same computer. Of these 39, 33 were deemed a repeat response by the same person (based on identical demographics or very similar responses) and were deleted. Of the remaining 2,014, 497 completed the demographics section but very few or no questions in the AD section, leaving 1,517. The questions about adverse effects were not answered by 86 people. This left 1,431 for the analyses for this paper (see Figure 1).
Figure 1. Flowchart of participants.

- 2,346 responded
- 213 had not taken ADs
- 2,133
- 42 did not confirm the three inclusion criteria (see text)
- 2,091
- 44 named drug other than AD
- 2,047
- 33 repeat responses by same person
- 2,014
- 497 insufficient data
- 1,517
- 86 did not answer questions about adverse effects
- 1,431 analysed
2.3. Data Analysis

A Total Adverse Effects (TAE) score was calculated by combining the scores (0 - 3) for the 20 side effects, producing a potential range of 0 to 60. (There were 57 responses with between one and four empty boxes in the adverse effects section and these were scored as ‘0’- ‘not at all’). The TAE thereby incorporates both the frequency of different adverse effects and their severity. Spearman Rank Order Correlation Coefficients (rho) were used to test for relationships between dimensional variables, e.g. age and likert scale measures of adverse effects. Independent sample, two-tailed t-tests were used to explore differences between mean scores in relation to gender, duration of treatment (≤ 3 years vs > 3 years) and types of drug. The level of significance was set at the p < .01 level because of the high number of tests and the consequent risk of false positives.

3. RESULTS

3.1. Sample Characteristics

Of the 1,517 participants, 1,018 (67.1%) had taken only ADs, and 499 (32.9%) had also taken APs. When completing the survey 813 (53.6%) were still taking ADs. Of the 1,497 who reported how long they had taken APs, 6.2% responded less than three months, 14.7% said 3 to 12 months, 16.5% one to three years, and 62.5% more than three years. The initial prescriber was a GP in 61.7% of cases and a psychiatrist in the other 38.3%.

The majority of respondents (70.5%) were women. Respondents’ ages ranged from 18 to 78 and averaged 45.1 (s.d. 13.5), with no gender difference. Half (50.7%) were employed; 14.8% were unemployed; 7.3% were students; and 27.2% ticked ‘other’. The most common responses to the open question ‘What ethnicity or cultural group do you most identify with?’ were: ‘White’ or ‘Caucasian’ (47.9%), ‘Australian’ (21.8%), ‘British’/’English’/’Anglo’ (12.3%), ‘European’ (8.0%), ‘New Zealander’ (3.0%) and ‘Asian’ (1.5%).
IP addresses showed that the 1,517 respondents lived in 38 countries, most commonly Australia (667; 44.0%) (from where the survey was conducted), the UK (277; 18.3%), and the USA (247; 16.3%). The nine other countries with more than 10 respondents were: New Zealand (61), Germany (48), Denmark (31), Canada (30), Ireland (29), the Netherlands (24), Norway (19), South Africa (18) and Sweden (14). The following 26 countries contributed between one and seven respondents: Albania, Algeria, Austria, Belgium, Bosnia, Bulgaria, Croatia, Czechia, Estonia, Faroe Islands, Finland, France, Greece, Iceland, India, Italy, Israel, Lithuania, Poland, Portugal, Romania, Slovakia, Slovenia, Spain, Switzerland, Ukraine.

Most participants (1283; 84.6%) replied ‘Yes’ to ‘Do you think you were prescribed antidepressant medication to treat depression?’ Of the 485 who reported their primary diagnosis, the most commonly reported diagnoses were: Depression - 30.1%; Bi-Polar Disorder - 21.6%; Schizophrenia spectrum disorders - 19.9%; Personality Disorders - 8.7%; Post Traumatic Stress Disorder - 3.5%, and Anxiety Disorders - 2.6%.

Although the focus of this article is the adverse effects of ADs, the respondents’ views on whether the drugs were helpful are important characteristics of the sample. Positive outcome was assessed with two questions: (i) ‘As a result of taking antidepressants, my depression was . . . .’ and (ii) ‘When taking antidepressants my quality of life is . . . .’ each followed by a five point multiple choice scale. Table 1 shows that about two thirds 65.2%) thought the drugs reduced their depression (‘greatly’ or slightly’), while 14.5% reported that they made the depression worse (‘slightly’ or ‘a lot’). Similarly 60.9% reported that their Quality of Life was ‘improved’, while 21.2% thought it was made ‘worse’.

Table 1. Perceived efficacy of antidepressant medication.

<table>
<thead>
<tr>
<th>‘As a result of ADs my depression was.....’</th>
<th>Greatly reduced</th>
<th>Slightly reduced</th>
<th>Unchanged</th>
<th>Slightly worse</th>
<th>A lot worse</th>
</tr>
</thead>
</table>

10
| (n = 1475) | 29.9% | 35.3% | 20.3% | 4.9% | 9.6% |
| 'As a result of ADs my quality of life was.....' | Greatly improved | Slightly improved | Unchanged | Slightly worse | A lot worse |
| (n = 1488) | 24.7% | 36.2% | 17.9% | 7.7% | 13.5% |

### 3.2. Total Adverse Effects

The side effects section was completed by 1,431 respondents. Nearly all (97.3%) reported at least one of the 20 side effects; with 83.3% reporting five or more, and 61.2% reporting ten or more. The mean number of adverse effects per respondent was 11.22 (s.d. 5.85).

Participants reported an average of 3.02 (s.d. 4.02) of the effects at the ‘severe’ ‘level. Table 2 shows that 16 of the 20 side effects listed in the questionnaire were reported by more than half the respondents, most commonly: ‘Feeling emotionally numb’ (70.6%), ‘Feeling foggy or detached’ (70.0%); ‘Feeling not like myself’ (66.2%), ‘Sexual difficulties’ (66.1%), ‘Drowsiness’ (62.7%), and ‘Reduction in positive feelings’ (60.4%).

The mean Total Adverse Effects (TAE) score was 21.02 (s.d.14.12).

### 3.3. The Antidepressants and Antipsychotics Group

The 499 respondents who had taken both ADs and APs (AD+AP) were younger (X = 42.8 years) than those who had only taken ADs (46.3 years), (t = 4.84. df = 1049.8, p < .0001); but this was unrelated to gender.

Of the 499, 478 completed the side effects section of the questionnaire. This AD+AP group reported a significantly greater mean number of symptoms (12.87) than the AD only group (10.39) (t = 4.55, df 934.1, p < .0001). The TAE mean of the AD+AP group (25.84, s.d.14.56) was also significantly greater than the AD only group (18.60, s.d.13.29) (t = 9.16, df 885.4, p < .0001). Table 2 shows that the AD+AP respondents had experienced each of the
20 effects to a significantly greater degree than the AD only group. The largest difference was for Suicidality, which was reported by 65.9% of the AD+AP group (24.7% at the severe level) compared to 42.5% of the AD only group (10.3% severe).

Table 2. Percentages reporting each of 20 adverse effects, with and without antipsychotics.

<table>
<thead>
<tr>
<th>Adverse Effect</th>
<th>Total sample (n = 1431)</th>
<th>ADs only (n = 953)</th>
<th>ADs + APs (n = 478)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Feeling emotionally numb</td>
<td>70.6</td>
<td>66.1</td>
<td>79.7 ***</td>
</tr>
<tr>
<td>Feeling foggy or detached</td>
<td>70.0</td>
<td>65.1</td>
<td>79.7 ***</td>
</tr>
<tr>
<td>Feeling not like myself</td>
<td>66.2</td>
<td>61.4</td>
<td>75.9 ***</td>
</tr>
<tr>
<td>Sexual difficulties</td>
<td>66.1</td>
<td>64.2</td>
<td>69.9*</td>
</tr>
<tr>
<td>Drowsiness</td>
<td>62.7</td>
<td>59.6</td>
<td>68.8 ***</td>
</tr>
<tr>
<td>Reduction in positive feelings</td>
<td>60.4</td>
<td>55.2</td>
<td>70.9 ***</td>
</tr>
<tr>
<td>Weight gain</td>
<td>60.1</td>
<td>55.8</td>
<td>68.6 ***</td>
</tr>
<tr>
<td>Dry mouth</td>
<td>59.3</td>
<td>54.6</td>
<td>68.6 ***</td>
</tr>
<tr>
<td>Distorted dreams</td>
<td>59.2</td>
<td>55.7</td>
<td>66.1 ***</td>
</tr>
<tr>
<td>Withdrawal effects</td>
<td>58.9</td>
<td>55.4</td>
<td>65.9 ***</td>
</tr>
<tr>
<td>Agitation</td>
<td>58.0</td>
<td>52.9</td>
<td>68.2 ***</td>
</tr>
<tr>
<td>Insomnia</td>
<td>57.7</td>
<td>55.7</td>
<td>61.5 ***</td>
</tr>
<tr>
<td>Caring less about others</td>
<td>54.5</td>
<td>49.9</td>
<td>63.6 ***</td>
</tr>
<tr>
<td>Dizziness</td>
<td>51.6</td>
<td>47.5</td>
<td>59.8 ***</td>
</tr>
<tr>
<td>Headaches</td>
<td>50.4</td>
<td>47.1</td>
<td>56.9 ***</td>
</tr>
<tr>
<td>Suicidality</td>
<td>50.3</td>
<td>42.5</td>
<td>65.9 ***</td>
</tr>
<tr>
<td>Nausea</td>
<td>46.8</td>
<td>43.3</td>
<td>53.8 ***</td>
</tr>
<tr>
<td>Feeling aggressive</td>
<td>41.0</td>
<td>37.6</td>
<td>47.9 ***</td>
</tr>
</tbody>
</table>
### 3.4 The Antidepressant Only Group

In order to reduce potential contamination of the rates of the side effects by the influence of APs, and so as to facilitate comparisons with the largest similar survey to date, in New Zealand [18], the findings for the 1,018 people who had taken ADs but not APs are presented separately. They are then analysed, like the New Zealand survey, in terms of gender, age and duration of treatment.

Of these 1,018 people, 953 completed the side effects section of the survey (see Tables 2 and 3). The mean number of adverse effects in this group was 10.39 (s.d. 5.83), and the mean TAE score was 18.60 (s.d. 13.29). Most (97.0%) reported at least one of the 20 side effects; with 80.3% reporting five or more, and 54.7% reporting ten or more. The mean number of adverse effects per respondent was 10.39 (s.d. 5.83). Table 2 shows that 12 of the 20 side effects listed in the questionnaire were reported by more than half the respondents, most commonly: ‘Feeling emotionally numb’ (66.1%), ‘Feeling foggy or detached’ (65.1%), ‘Sexual difficulties’ (64.2%) and ‘Feeling not like myself’ (61.4%). The mean Total Adverse Effects (TAE) score was 18.60 (s.d. 13.29).

#### 3.4.1 Gender

<p>| | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Addiction</td>
<td>40.5</td>
<td>36.8</td>
<td>47.7 **</td>
</tr>
<tr>
<td>Tremors</td>
<td>37.4</td>
<td>32.3</td>
<td>47.5 ***</td>
</tr>
<tr>
<td>Total Adverse Effects (TAE) a</td>
<td>21.02</td>
<td>18.60</td>
<td>25.84 ***</td>
</tr>
<tr>
<td>Mean number of adverse effects b</td>
<td>11.22</td>
<td>10.39</td>
<td>12.87 ***</td>
</tr>
</tbody>
</table>

a = range 0-60  
b = range 0-20  
* < .01; ** < .001; *** < .001 (greater than ADs only group)
Gender was related to neither mean number of adverse effects nor TAE mean scores. Table 3 shows, however, that two of the 20 specific adverse effects were reported to a greater degree by men: Sexual difficulties and Caring less about others (both at the p < .01 level). For example 58.2% of the men reported Caring less about others, compared to 46.6% of the women.

3.4.2. Age

Age was negatively correlated with number of adverse events (\(\rho = .18\), p < .0001) and to TAE (\(\rho = .10\), p < .0001). Age was significantly negatively correlated to eleven individual adverse effects, (nine at the p < .0001 level; see Table 3), most strongly with Nausea and Feeling emotionally numb. Age was positively related only to Dry mouth (p < .001).

3.4.3. Treatment duration

The 549 who had taken ADs for more than three years reported a significantly higher number of adverse effects (10.80) than the 404 who had taken them for three years or less (9.81) (t = 2.60, df 950, p < .01), and a higher TAE mean (19.75 vs. 17.01) (t = 3.16, df 950, p < .01). Table 3 shows that five adverse effects were reported at a higher level by the group on ADs for more than three years (all at the p < .0001 level), with the biggest differences being for Addiction, Withdrawal effects and Weight Gain. For example, 26.5% of those on ADs for three years or less reported Addiction to ADs (10.3% severe), compared to 44.5% of those on them for over three years (19.9% severe).

Table 3. Percentages of respondents who took only antidepressants reporting each of 20 adverse effects, analysed by gender, age and treatment duration (more than three years vs. three years or less); with findings of the 2014 New Zealand survey in brackets.
<table>
<thead>
<tr>
<th>Symptom</th>
<th>any</th>
<th>severe</th>
<th>gender</th>
<th>age</th>
<th>duration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Feeling emotionally numb</td>
<td>66.1</td>
<td>18.2</td>
<td>Y***</td>
<td>y***</td>
<td></td>
</tr>
<tr>
<td>Feeling foggy or detached</td>
<td>65.1</td>
<td>16.3</td>
<td>*</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sexual difficulties</td>
<td>64.2</td>
<td>20.0</td>
<td>M*</td>
<td>&gt;3yrs*</td>
<td></td>
</tr>
<tr>
<td>Feeling not like myself</td>
<td>61.4</td>
<td>16.7</td>
<td>Y***</td>
<td>y***</td>
<td>≤3yrs*</td>
</tr>
<tr>
<td>Drowsiness</td>
<td>59.6</td>
<td>10.3</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Weight gain</td>
<td>55.8</td>
<td>14.0</td>
<td>F*</td>
<td>&gt;3yrs**</td>
<td></td>
</tr>
<tr>
<td>Insomnia</td>
<td>55.7</td>
<td>11.4</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Distorted dreams</td>
<td>55.7</td>
<td>11.0</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Withdrawal effects</td>
<td>55.4</td>
<td>23.8</td>
<td>Y***</td>
<td>&gt;3yrs**</td>
<td></td>
</tr>
<tr>
<td>Reduction in positive feelings</td>
<td>55.2</td>
<td>13.2</td>
<td>M*</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Agitation</td>
<td>52.9</td>
<td>9.2</td>
<td>Y**</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dry mouth</td>
<td>54.6</td>
<td>10.7</td>
<td>O***</td>
<td>&gt;3yrs**</td>
<td></td>
</tr>
<tr>
<td>Caring less about others</td>
<td>49.9</td>
<td>11.0</td>
<td>M*</td>
<td>&gt;3yrs**</td>
<td></td>
</tr>
<tr>
<td>Dizziness</td>
<td>47.5</td>
<td>6.3</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Headaches</td>
<td>47.1</td>
<td>6.4</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nausea</td>
<td>43.3</td>
<td>5.1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Suicidality</td>
<td>42.5</td>
<td>10.5</td>
<td>Y***</td>
<td>&gt;3yrs*</td>
<td></td>
</tr>
<tr>
<td>Feeling aggressive</td>
<td>37.6</td>
<td>7.1</td>
<td>M*</td>
<td>y***</td>
<td></td>
</tr>
<tr>
<td>Addiction</td>
<td>36.8</td>
<td>13.4</td>
<td></td>
<td>&gt;3yrs**</td>
<td></td>
</tr>
<tr>
<td>Tremors</td>
<td>32.3</td>
<td>4.8</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* < 01; ** < .001; *** .0001  
Age: spearman rank correlations (Y = younger age; O = older age)  
Gender and Treatment Duration: t-test independent means  
(-) = this variable not included in New Zealand survey
### 3.5. Medication Types

Of the 1151 in the total sample who identified their AD, the most frequently named were: fluoxetine 223, 19.4%; sertraline 175, 15.2%; citalopram 157, 13.6%; paroxetine 112, 9.7%; venlafaxine 112, 9.7%; and escitalopram 93, 8.1%. There were also 126 cases of tricyclic ADs (10.9%), mostly amitriptyline. Fifty one (4.4%) responded in ways that indicated use of multiple ADs (e.g. ‘lots’, ‘various’, ‘Lost count of how many different ones’).

Table 4 shows that there were no significant differences between the types of drugs in terms of the mean number of adverse effects. Citalopram, however, produced a lower mean TAE score than paroxetine, fluoxetine and tricyclics, all at the p < .01 level. For example: the mean TAE for citalopram (18.14) was significantly lower than the mean for paroxetine (23.44); t = 3.15, df = 250, p < .01. Citalopram’s TAE mean was also lower than the mean for the other six drug types combined (21.21); t = 2.85, df = 233.1, p < .01. More specifically citalopram produced significantly lower means for Suicidality, Caring less about others, Reduction in positive feelings and Feeling aggressive (all at the p < .01 level or beyond), with the biggest difference being for Caring less about others; t = 3.63, df = 231.3, p < .0001.

Table 4 also shows that usage of multiple ADs produced both a significantly higher TAE mean (p < .0001) and a higher mean number of adverse effects (p < .01) than all seven individual drug types combined. (Use of multiple ADs was unrelated to age and gender).

### Table 4. Total Adverse Effects scores, and number of adverse effects, by medication type.

<table>
<thead>
<tr>
<th>Medication Type</th>
<th>n</th>
<th>Mean ‘Total Adverse Effects’ score (and S.D.)</th>
<th>Mean number of adverse effects (and S.D.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Paroxetine</td>
<td>102</td>
<td>23.44 (14.88)</td>
<td>11.83 (5.90)</td>
</tr>
<tr>
<td>Fluoxetine</td>
<td>205</td>
<td>22.25 (14.38)</td>
<td>11.55 (5.74)</td>
</tr>
<tr>
<td>Drug Type</td>
<td>Count</td>
<td>Mean (SD)</td>
<td>SE (SD)</td>
</tr>
<tr>
<td>------------</td>
<td>-------</td>
<td>-----------</td>
<td>---------</td>
</tr>
<tr>
<td>Tricyclics</td>
<td>15</td>
<td>22.79 (15.17)</td>
<td>11.53 (5.91)</td>
</tr>
<tr>
<td>Escitalopram</td>
<td>90</td>
<td>20.32 (14.80)</td>
<td>11.09 (6.05)</td>
</tr>
<tr>
<td>Venlafaxine</td>
<td>104</td>
<td>21.59 (15.07)</td>
<td>11.02 (5.95)</td>
</tr>
<tr>
<td>Sertraline</td>
<td>171</td>
<td>19.18 (12.90)</td>
<td>10.51 (5.31)</td>
</tr>
<tr>
<td>Citalopram</td>
<td>150</td>
<td>18.14 a,b (11.73)</td>
<td>10.27 (5.52)</td>
</tr>
<tr>
<td>Multiple ADs</td>
<td>51</td>
<td>28.71 c (14.80)</td>
<td>13.49 d (5.48)</td>
</tr>
</tbody>
</table>

a. Lower than Paroxetine, Fluoxetine, and Tricyclics (p < .01)
b. Lower than the six other individual drug types combined (p < .01)
c. Higher than the seven drug types combined (p < .0001)
d. Higher than the seven drug types combined (p < .01)

3.6. Suicidality

Suicidality was reported to be a side effect of taking ADs by 50.3% of the total sample, 42.5% of the AD only group, and 65.9% of the AD+AP group. The percentages reporting ‘severe’ suicidality were 15.1%, 10.3%, and 24.7% respectively. Suicidality was negatively correlated with age (rho = .12, p < .0001), but was not related to gender. The specific adverse effects most strongly correlated with Suicidality were: Feeling not like self (rho = .58), Reduction in positive feelings (rho = .58) and Feeling aggressive (rho = .54) (all p < .0001). (The specific tests reported here are for the AD only group, but the findings were very similar for the AD+AP group and the sample as a whole).

Citalopram was significantly less likely than the other drugs combined to lead to suicidality (t = 2.53, df = 236.3, p < .0001). Both forms of polypharmacy significantly increased the rate of suicidality: multiple ADs (t = 3.38, df 1072, p < .001), and APs in addition to ADs (t = 9.92, df 837.3, p < .0001).
3.7. Information about Adverse Effects

In response to ‘Did the doctor inform you of any possible side effects?’ 520 (34.3%) replied Yes, 973 (64.1%) said No, and 24 (1.6%) did not reply. Being informed was unrelated to age or gender, or to whether the prescriber was a GP or a Psychiatrist.

Those who had been informed about side effects reported fewer adverse effects on average (9.75) than those not informed (12.01) \(t = 7.07; df = 1422, p < .0001\); and also produced a lower TAE mean (36.71 vs 43.36; \(t = 8.69, df = 11180.9, p < .0001\)). Every one of the 20 side effects was reported to a significantly greater extent by people who had not been informed about side effects (18 at the \(p < .0001\) level).

Of the 520 who recalled being told about side effects, 371 (71.3%) recalled one or more specific side effects they had been told about (see Table 5), most frequently Nausea (77) and Weight changes (75). A further 21 referred to a written list of effects being given, or read, to them.

Table 5. Frequency of adverse effects about which participants had been informed (6 or more).

<table>
<thead>
<tr>
<th>Adverse Effect</th>
<th>Frequency</th>
<th>Description</th>
<th>Count</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nausea/vomiting</td>
<td>77</td>
<td>Increased depression</td>
<td>22</td>
</tr>
<tr>
<td>Weight/appetite changes</td>
<td>75</td>
<td>Anxiety</td>
<td>17</td>
</tr>
<tr>
<td>Sleepiness/drowsiness/sedation</td>
<td>56</td>
<td>Mania/mood swings</td>
<td>10</td>
</tr>
<tr>
<td>Dry mouth</td>
<td>51</td>
<td>Withdrawal symptoms/addiction</td>
<td>9</td>
</tr>
<tr>
<td>Sexual dysfunction</td>
<td>42</td>
<td>Constipation</td>
<td>8</td>
</tr>
<tr>
<td>Suicidality</td>
<td>35</td>
<td>Tremors</td>
<td>8</td>
</tr>
<tr>
<td>Insomnia</td>
<td>22</td>
<td>Agitation/restlessness</td>
<td>8</td>
</tr>
<tr>
<td>Dizziness</td>
<td>26</td>
<td>Loss of interest, emotions/lethargic</td>
<td>8</td>
</tr>
<tr>
<td>Headaches</td>
<td>22</td>
<td>Blurred vision</td>
<td>6</td>
</tr>
</tbody>
</table>
4. DISCUSSION

This study confirms previous studies that have found that when AD users are asked directly they report an extensive range of adverse effects at very high frequencies. Thus this kind of survey can be a valuable adjunct to more traditional methodologies for assessing adverse drug effects, as can spontaneous reporting systems [24] and real-world data gathered from clinical practice [25].

Sixteen effects were experienced by more than half of the total sample. Amongst those who had only taken ADs the rates are slightly higher than those found in in the earlier New Zealand survey [18], but are quite similar overall. For example Feeling emotionally numb was reported by 66.1% in the current study and 60.4% in the New Zealand study. Furthermore, the three side effects introduced to the current study because of spontaneous reporting in the New Zealand study all produced very high response rates: Feeling foggy or detached (65.1%), Insomnia (55.7%) and Distorted dreams (55.7%) (see Table 3).

The second major finding is the confirmation of very high rates of adverse effects in the emotional and interpersonal domains [12-17]. In both the current international study and the earlier New Zealand survey [18] Emotional numbing, Sexual difficulties, Feeling not like myself, Reduction in positive feelings and Caring less about others were extremely common. Similarly, a UK online survey of over 1,000 AD users [26] found that 60% reported negative effects in at least one of the five interpersonal domains assessed (most commonly ‘Sex life’ and ‘Work or study’). A recent examination of over 3,000 online comments by AD users found that ‘Emotional and behavioral’ adverse effects were reported far more than biological/medical effects such as Cardiovascular, Gastrointestinal or Metabolic [27]. All
these findings contrast with more traditional studies of adverse effects, such as drug trials which have focussed, like most officially sanctioned checklists, on the adverse biological/medical effects [9-11].

There was some consistency between the New Zealand and current study in terms of gender and age. In both studies men were more likely than women to report Sexual difficulties and Caring less about others. In both studies younger people reported more adverse effects overall, particularly for Feeling emotionally numb, Feeling not like myself, Suicidality, Dizziness, and Nausea.

4.1. Polypharmacy

Presenting the rates of adverse effects for those taking only ADs is a sensible way to try to ensure that the reported effects are caused specifically by ADs. They underestimate, however, rates in the real world, where it is commonplace for people to be on multiple types of psychiatric medication [28, 29]. Therefore the rates for the total sample in the current study may be the most accurate account of the adverse effects being experienced by AD users. The finding that every one of the 20 adverse effects was significantly more common amongst those who took both ADs and APs than among those who took only ADs is important. The interactive negative effects of different types of psychiatric drugs is rarely studied in traditional drug trials.

In the UK survey of over 1,000 AD users [26] more than half were taking one or more additional psychiatric drugs. The number of types of medication correlated with overall severity of side effects. The number of medication types was also related to each of the six specific types of adverse effect. For example, while 44% of those on antidepressants alone reported an adverse effect on their sex life, this was the case for 54% of those on
antidepressants plus one other psychiatric medication, and 63% of those on three additional medications. A 2013 review [28] concluded:

While evidence for the added benefit of psychiatric polypharmacy is limited, there is growing evidence regarding the increased adverse effects associated with such combinations.

In the current study, three specific adverse effects were reported by three out of every four of the participants who took both ADs and APs: Feeling emotionally numb, Feeling foggy or detached, and Feeling not like myself.

4.2. Suicidality

Since the 1990s studies have documented that ADs increase suicidality for a small number of people [30, 31]. Asking AD users directly, however, suggests that the problem may be far greater than originally thought. In the current study ‘suicidality’… ‘as a result of taking the antidepressant’ was reported by 42.5% of the AD only group. This corroborates the 38.9% finding in the large New Zealand survey [18]. The rates at the ‘severe’ level of suicidality were 10.5% and 7.8% respectively. In both studies the likelihood of experiencing suicidality increased the longer the person stayed on the ADs. Furthermore, nearly two in every three people (65.9%) taking both ADs and APs reported suicidality (24.7% at the ‘severe’ level). The other form of polypharmacy, multiple ADs, also increased risk of suicide compared to taking just one AD.

Increased suicidality as a result of ADs is especially strong in children, adolescents and young adults [32, 33], leading the F.D.A. to place a ‘Black Box’ warning, in 2006, on all ADs for people under the age of 26 [32]. The current study excluded children but, like the New Zealand survey, confirmed that younger adults are indeed even more likely to report suicidality than older adults.
4.3. Withdrawal Effects and Addiction

More than half of participants taking only ADs (55.4%) reported Withdrawal effects (23.8% at the ‘severe’ level). This is consistent with the 54.9% in the New Zealand survey [18]. Furthermore, about a third reported Addiction in both studies (36.8% and 27.4% respectively). In both studies both Addiction and Withdrawal effects were, unsurprisingly perhaps, reported significantly more by people who had been on ADs for more than three years. For example, in the New Zealand study 74% of those on ADs for more than three years reported Withdrawal symptoms, and 46% of those reporting the symptoms described them as ‘severe’ [34]. A recent Dutch study found that of 692 users of tapering strips 671 (97%) had experienced some level of withdrawal effects when they had previously tried to stop their ADs; and that 339 of those 671 (51%) reported the severity of the withdrawal as 7 on a 7 point scale [35].

The issue of whether ADs should be considered addictive has polarised opinion. In a survey of over 2,000 members of the British public, many of whom of course were either taking ADs or knew people who were taking them, 78% described ADs as addictive [36]. The Diagnostic and Statistical Manual for Mental Disorders [37] requires that two of eleven criteria are met for ‘substance dependence’. The current study and the large New Zealand survey [18] indicate that ADs may meet at least four of these criteria for many people: ‘Withdrawal manifesting as a characteristic syndrome’, ‘Wanting to cut down or quit but not being able to do it’, ‘Continued use despite persistent or recurring social or interpersonal problems caused or made worse by substance use’ and ‘Stopping or reducing important social, occupational, or recreational activities due to substance use.’

Nevertheless, the only reference to the issue in the American Psychiatric Association Practice Guideline for depression is ‘Common misperceptions about antidepressants (e.g.
they are addictive) should be clarified’ [38]. (The authors listed, between them, over 80 instances of receiving payments from drug companies). In the UK the Royal College of Psychiatrists (R.C.P.) states: ‘We would like to reassure readers that despite some people having symptoms of withdrawal when stopping antidepressants, antidepressants are not addictive’ [39]. In February, 2018 the President of the R.C.P. and the Chair of its Psychopharmacology Committee jointly informed the public that ‘We know that in the vast majority of patients, any unpleasant symptoms experienced on discontinuing antidepressants have resolved within two weeks of stopping treatment’ [40].

The term ‘Discontinuation Syndrome’ was introduced in 1996, at a ‘closed symposium’ in Phoenix, Arizona [41]. The gathering was funded by Eli Lilly. The ‘Discontinuation Consensus Panel’ consisted of industry funded professionals who argued that AD withdrawal is very different from withdrawal from benzodiazepines.

Researchers at the Nordic Cochrane Centre disagree with the A.P.A, the R.C.P, and Eli Lilly spokespersons. Their 2012 review [42] of 45 papers on benzodiazepines and 31 papers on SSRIs concluded that ‘Withdrawal reactions to SSRIs appear to be similar to those for benzodiazepines; referring to these reactions as part of a dependence syndrome in the case of benzodiazepines, but not selective serotonin re-uptake inhibitors, does not seem rational’.

The disputes seem to rest primarily on disagreements about the definition of ‘addiction’. An approach which circumvents these disagreements is to allow people who have taken ADs to interpret the term ‘addiction’ as they wish, thereby respecting their actual personal experience. A 2004 review of studies of ‘Patients’ preferences in the treatment of depressive disorder in primary care’ found that ‘Antidepressants were often regarded as addictive’ [43]. A 2014 review [12], of studies of ‘Patient-centred perspectives on antidepressant use’, concurred, reporting that ‘the most frequently mentioned reason for a negative opinion of antidepressants is that they may be addictive’. High rates of addiction and withdrawal effects
among AD users, when asked directly, have been found in Denmark [44], the Netherlands [35, 45], Scotland [46] and the USA [13, 47]. A 2012 survey by the Royal College of Psychiatrists in the UK found that of 817 people who had stopped taking ADs, 63% experienced withdrawal symptoms [48].

High rates of addiction and difficulties withdrawing are consistent with findings that the ever increasing prescription rates of ADs are not explicable in terms of increased incidence of depression, or changes in help-seeking behavior [49], but rather in terms of increases in repeat prescriptions. A study of 189,851 GP patients in the UK found that a doubling of prescribing over eight years was explained not by increases in new prescriptions but by a doubling of the number of prescriptions per patient [50].

4.4. Information

In the current study only one third of AD users (34%) recalled being told about any side effects. This is lower than the 64% in the New Zealand survey and the findings of two British studies, 59% [51] and 55% [52]. Together, these four studies suggest that between one and two thirds of AD recipients are told nothing about side effects. What information that is imparted seems to focus on biological side effects. In both the current study and the New Zealand study [53] the side effects that people were most commonly warned about were nausea and weight gain. Less than five percent were warned about increased suicidality in either study. None were told about feeling less like themselves or about possible effects on their relationships with others (other than sexual dysfunction). In both studies fewer than two percent were told anything about withdrawal effects [53]. This breaches the fundamental ethical principle of ‘informed consent’ which is binding on all health professionals.

4.5. Strengths and Limitations
This is a self selected, convenience sample, despite being the second largest sample ever directly surveyed. Although it is more international than the original New Zealand based survey, almost the entire sample lived in Europe, North America or Australasia. The over representation of women (70.5\%) is less of an issue as women are prescribed ADs approximately twice as often as men. The very high rates of adverse effects raises the possibility that the sample is biased towards people who had found ADs ineffective and therefore may have had ‘an axe to grind’. However, two thirds (65.2\%) reported that the ADs had reduced their depression. The study relies on self-report, but so do most traditional studies of the effects of drugs, positive and negative. The fact that some of the reports are for events many years previously is more of a problem. For example, some participants may actually have been given more information about adverse effects than they could recall years later.

The failure to gather data about the duration of withdrawal effects is also a limitation. A recent review [54] found that they typically ‘last a few weeks’ but identified ‘many variations… including longer persistence of disturbances’ and two studies documenting withdrawal symptoms up to one year following paroxetine discontinuation. More research is urgently needed in this area.

It is possible that some respondents may not have taken ADs and may be responding on behalf of a loved one or may even have, for some unknown reason, fabricated a set of responses. Safeguards were in place, however, against repeat responders and people who mistakenly thought their drug was an AD when it was not (see Methods). Finally, the study did not allow for possible confounding variables such as substance abuse and other medicines besides APs.
5. CONCLUSION

As noted earlier, the experience of patients is finally receiving the attention it so obviously warrants. Directly surveying patients, and using PROMs, helps to ‘ensure that research is both robust and of maximum value for the use of medicinal products, therapy, or health services’. . . . and ‘reflects the ongoing health service commitment of involving patients and the public within the wider context of the development and evaluation of health care service delivery and quality improvement.’ [19, p.61]

Asking people directly about their experience of ADs reveals that their adverse effects may be far more varied and prevalent and damaging than previously realised when relying on more traditional research methodologies. Together with the consistent findings that ADs are no more effective than placebo for many recipients, the findings of these recent direct-to-consumer surveys produce a cost-benefit analysis that cannot justify the current extraordinarily high prescription rates of ADs. A first step towards reducing inappropriate prescribing would be for all prescribers to fully inform all potential recipients of all potential side effects, including suicidality, emotional blunting and withdrawal effects. Psychiatric polypharmacy should be a last resort rather than standard practice. It seems that health services need to challenge the denial and minimization of the drug companies and some professional bodies and urgently develop drug withdrawal services for the millions already trying to come off these drugs but experiencing withdrawal effects which are sometimes severe and protracted.

CONFLICT OF INTEREST

Both authors declare no financial, personal or potential conflicts of interest.
REFERENCES


[26] Read J, Gee A, Diggle J, Butler H. The interpersonal adverse effects reported by 1,008 users of antidepressants; and the incremental impact of polypharmacy. Psychiatry Res 2017; 256: 423-7.


