Most Antidepressant Use in Primary Care Is Justified; Results of the Netherlands Study of Depression and Anxiety

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Abstract

Background: Depression is a common illness, often treated in primary care. Many studies have reported undertreatment with antidepressants in primary care. Recently, some studies also reported overtreatment with antidepressants. The present study was designed to assess whether treatment with antidepressants in primary care is in accordance with current guidelines, with a special focus on overtreatment.

Methodology: We used baseline data of primary care respondents from the Netherlands Study of Depression and Anxiety (NESDA) (n = 1610). Seventy-nine patients with treatment in secondary care were excluded. We assessed justification for treatment with antidepressant according to the Dutch primary care guidelines for depression and for anxiety disorders. Use of antidepressants was based on drug-container inspection or, if unavailable, on self-report. Results were recalculated to the original population of primary care patients from which the participants in NESDA were selected (n = 10,677).

Principal Findings: Of 1531 included primary care patients, 199 (13%) used an antidepressant, of whom 188 (94.5%) (possibly) justified. After recalculating these numbers to the original population (n = 10,677), we found 908 (95% CI 823 to 994) antidepressant users. Forty-nine (95% CI 20 to 78) of them (5.4%) had no current justification for an antidepressant, but 27 of them (54.5%) had a justified reason for an antidepressant at some earlier point in their life.

Conclusions: We found that overtreatment with antidepressants in primary care is not a frequent problem. Too long continuation of treatment seems to explain the largest proportion of overtreatment as opposed to inappropriate initiation of treatment.

Introduction

Depression is a common disorder which is associated with a great amount of morbidity because of its highly recurrent and chronic nature.[1] Most patients with depression are treated in primary care.[2,3] Guidelines on the treatment of depression in primary and secondary care consider treatment with antidepressants and/or psychotherapy indicated for all patients with major depressive disorder (MDD).[4–10] According to these guidelines the treatment should be continued for 6 months after remission (continuation treatment) of a first episode, while it should be continued for one or more years (maintenance treatment) in patients with a recurrent MDD or chronic depression.[4–10]

Various studies reported that treatment of depression in primary care is not according to guideline recommendations.[11–16] Most studies reported undertreatment, especially with antidepressants of patients with MDD.[11–16] However, in recent years there has also been a lot of attention for overtreatment with antidepressants.[17–19] The fact that in the last 6 months of 2005 760,000 people in the Netherlands (population 16,500,000) were prescribed an antidepressant, according to the Dutch Foundation for Pharmaceutical Statistics (SFK, www.sfk.nl), led to discussions in
The 79 respondents already receiving treatment for psychiatric conditions in secondary care (defined as more than one contact with either an institute for mental/psychiatric health care or an independent psychiatrist) were excluded from our study sample, yielding a total sample of 1531 respondents for the present analyses.

### Description of Procedures or Investigations undertaken

#### Measures.
As part of the screening procedure, all respondents filled out the K-10 plus.

Demographic data (age, gender, ethnicity, education) were assessed during the baseline interview.

Current and lifetime diagnoses of MDD, Dysth, current diagnosis of miD, comorbid anxiety disorders (social phobia, panic disorder, agoraphobia, generalized anxiety disorder) based on DSM-IV were assessed with a structured interview, the World Health Organization Composite International Diagnostic Interview – lifetime version 2.1 (CIDI), which is considered the gold standard for diagnosing depressive and anxiety disorders in large epidemiological studies.[35–37] Specifically trained research staff (mainly psychologists and research nurses) conducted the baseline interview including the CIDI.

From the data of the CIDI interview, in which all depressive symptoms were listed separately, we created a variable for depressive symptoms, defined as having had one or more DSM-IV symptoms of depression during at least two weeks lifetime with at least either anhedonia or depressed mood, without fulfilling the criteria for diagnosis of MDD or dysthymia. From this data we also created a variable “chronic MDD”, defined as having had a lifetime diagnosis of MDD and 24 months of (probably uninterrupted) symptoms of depression in the past five years as recorded with the life-chart method. The life chart is a method for recalling depressive symptomatology, the respondent was asked during the interview to mention several important (personal) events from the last several years and was subsequently asked to recall if there was some depressive symptomatology at that point. The life chart has been proven useful to assess the course of illness in patients with mood disorders.[38–40]

#### Outcome variables

Whether respondents used antidepressants was based on drug container inspection of all drugs used in the past month at baseline and classified according to the World Health Organization Anatomical Therapeutic Chemical (ATC) classification. If respondents had forgotten to take the medication to the interview, their use was based on self-report (done for 35.3% of all subjects). The use of two different methods for assessing antidepressant drugs was not a problem in the current study, as we were not interested in patient compliance, for which self report and drug container inspection can give very different results, but only in physician prescription behaviour. We therefore used the drug container inspection only to assess which medications were used and not for pill counts. Use of antidepressants included selective serotonin reuptake inhibitors (ATC-code N06AB), tricyclic antidepressants (N06AA) and other antidepressants (N06AF/N06AX). St. John’s wort was not considered an antidepressant.

### Justification for treatment with an antidepressant

To determine the justification for treatment with an antidepressant, we followed the recommendations from the guidelines for depression and for anxiety disorders of the Dutch General Practitioner’s Association (NHG).[7,41] Treatment was considered justified when it was mentioned in the guideline as (one of the) first step option(s) and considered possibly justified when it was
mentioned as (one of the) second step option(s). For depressive disorders the depression guideline recommends the use of an antidepressant during six months after response for a first episode of MDD as one of the first step treatment options, although dependent on the degree of suffering or dysfunction. As dysfunction is a criterion for the diagnosis of MDD and patients consulted their physician, we assumed that probably most had at least some degree of suffering or dysfunction. Therefore, we considered treatment with antidepressants justified when a respondent had suffered an episode of MDD in the past year. In case of recurrent or chronic MDD the guideline recommends one to five years of maintenance treatment, with the option for longer in patients with previous recurrences after withdrawal of antidepressants. Therefore, treatment of chronic or recurrent MDD for up to two years was considered justified, all treatment longer than two years was considered possibly justified. In case of dysthymia an antidepressant is mentioned in the depression guideline as (one of the) second step option(s) and therefore considered possibly justified. Antidepressants were not considered justified for depressive states not fulfilling criteria for MDD or dysthymia.

As antidepressants are also registered for the treatment of anxiety disorders, we also considered treatment with antidepressants justified in case the guideline recommendations from the anxiety disorder guideline were followed. This guideline recommends treatment with an antidepressant in case of the presence of an anxiety disorder in the last year, with the option (i.e. possibly justified) to continue the treatment for a longer period.

Over-treatment was considered present when a respondent received an antidepressant without justification or possible justification, i.e. without a non-recurrent (i.e. single) episode of MDD or dysthymia in the past year, or an anxiety disorder or recurrent MDD or chronic depression in lifetime.

**Ethics**

The study protocol of NESDA was approved centrally by the Ethical Review Board of the VU University Medical Center and subsequently by local review boards of each participating center. After full verbal and written information about the study, written informed consent was obtained from all participants at the start of baseline assessment. A full ethics statement of NESDA is found elsewhere. [30]

**Statistical methods.** Descriptive statistics and frequencies were used to describe the use of AD and psychological treatment. We recalculated the found numbers and percentages of justified and unjustified treatment with antidepressants in our sample to the original population of 10,677 persons who returned a completed K-10 plus screener questionnaire. This backward projection was done in several steps, which can be derived by reading Figure 1 from the bottom up, or from table 1.

In the first step, we split our sample into four groups; no use of an antidepressant, justified use, possibly justified use and unjustified use. We will refer to these groups as “justification groups”. After that, we registered the number of screen-positives and screen-negatives in each of the justification groups. These numbers were then multiplied by a correction factor (respectively

![Figure 1. Recruitment flow respondents.](doi:10.1371/journal.pone.0014784.g001)
Table 1. Results of recalculation of justified and unjustified antidepressant use to screener population.

<table>
<thead>
<tr>
<th>Study sample (n = 1531)</th>
<th>Screen positive (n = 4592)</th>
<th>Screen negative (n = 6085)</th>
<th>Screener population (n = 10,677)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Definitely justified AD use</td>
<td>95 (47.7%)</td>
<td>94 * 4.50 = 422</td>
<td>1* 12.00 = 12</td>
</tr>
<tr>
<td>Possibly justified AD use</td>
<td>93 (46.7%)</td>
<td>92 * 4.50 = 413</td>
<td>1* 12.00 = 12</td>
</tr>
<tr>
<td>Unjustified AD use</td>
<td>11 (5.5%)</td>
<td>11 * 4.50 = 49</td>
<td>0 * 12.00 = 0</td>
</tr>
<tr>
<td>Total</td>
<td>199 (100%)</td>
<td>197 * 4.50 = 884</td>
<td>2 * 12.00 = 24</td>
</tr>
</tbody>
</table>

1) Correction factor screen-positive group: total number of screen-positives divided by number of screen-positives in the study sample (4592/1024 = 4.5).
2) Correction factor screen-negative group: total number of screen-negatives divided by number of screen-negatives in the study sample (6085/506 = 12.0).
3) 95% Confidence Interval.

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Results

Study population

The average age of the population was 45.8 years, 1054 (68.8%) of respondents were female and 477 (31.2%) male.

The diagnoses of the respondents with lifetime depression/depressive symptoms (n = 1064, of whom 651 with a comorbid anxiety disorder) were as follows: 807 respondents had a lifetime MDD of whom 428 in the past year; 23 had a lifetime Dysth without history of MDD of whom 16 in the past year; and 234 had a lifetime mD or depressive symptoms in lifetime without a history of MDD or Dysth, of whom 44 had a mD in the past month (incidence in last year unknown). The diagnoses of the respondents with a lifetime anxiety disorder (n = 762) were: 345 patients with social phobia; 344 patients with a panic disorder (with or without agoraphobia); 131 patients with agoraphobia; and 330 patients with a generalized anxiety disorder (324 patients had more than one anxiety disorder).

Justified and non-justified treatment

Of the respondents with a depression/depressive symptoms (MDD, dysthymia, mD or depressive symptoms, n = 1064), 189 (17.8%) used an antidepressant, of whom 75 (39.7%) had a justification, 68 (36.0%) a possible justification and 46 (24.3%) did not have a justification.

However, of the 46 antidepressants users with a depression/depressive symptoms without a justification for treatment with an antidepressant (non-recurrent MDD, mD or depressive symptoms) nine had only a depression/depressive symptoms and 37 had a lifetime anxiety disorder (26 in the past year), which means that the antidepressant may have been prescribed for the anxiety disorder rather than the depression. This means that in only nine patients (4.5%) with a depression/depressive symptoms treatment was not justified.

Of the 111 respondents with a lifetime anxiety disorder without a depression/depressive symptoms eight patients (10.1%) used an antidepressant, in three this was justified and in five possibly justified.

Only 2 (0.6%) of the 356 respondents without a lifetime depression or anxiety disorder used an antidepressant. This treatment obviously was not justified.

Table 2 shows a summary of justified and not justified treatment in our sample. In total 11 respondents used an antidepressant without a definite or possible justified reason. Eight of them used a SSRI, one a TCA (at low dose) and two another antidepressant; the duration of antidepressant use varied from 0 to 120 months, with a median of 48 months. Nine of these respondents had a depression/depressive symptoms (six a single episode of MDD more than one year ago, three mD or depressive symptoms). They represent 0.77% of the 1175 respondents with a lifetime depression or anxiety disorder. The other two represent 0.56% of the 356 respondents without a lifetime depression/depressive symptoms or anxiety disorder. Thus, 6 of the 11 respondents (54.5%) had a justified reason for treatment with an antidepressant at some earlier point in their life because of an episode of MDD.

Table 2. Treatment with antidepressants and justification for treatment.

<table>
<thead>
<tr>
<th>Justification for treatment</th>
<th>No AD</th>
<th>AD</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unjustified</td>
<td>566 (98.1%)</td>
<td>11 (1.9%)</td>
<td>577</td>
</tr>
<tr>
<td>Possibly justified</td>
<td>225 (70.8%)</td>
<td>93 (29.3%)</td>
<td>318</td>
</tr>
<tr>
<td>Definitely justified</td>
<td>541 (85.1%)</td>
<td>95 (14.9%)</td>
<td>636</td>
</tr>
<tr>
<td>Total</td>
<td>1332 (87.0%)</td>
<td>199 (12.9%)</td>
<td>1531</td>
</tr>
</tbody>
</table>

Treatment was considered definitely justified in case of a MDD in the past year, or recurrent or chronic MDD with antidepressant treatment ≥24 months, or an anxiety disorder in the past year. Treatment was considered possibly justified in case of dysthymia in the past year, or a recurrent or chronic MDD with antidepressant treatment for more than 24 months, or an anxiety disorder over one year ago. All other antidepressant treatment was considered unjustified. All numbers are absolute number of respondents (percentage of total respondents in row).

AD = antidepressant.

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Use of antidepressants in the primary care population

With the selection procedure as described in the method section, we can calculate what our findings mean for the total population of respondents (n = 10,677) who returned the K-10 plus (table 1).

In our sample there were 188 antidepressant users with justified antidepressant use of whom 95 (50.5%) with a definite justified reason, and 93 (49.5%) with a possible justified reason: 2 among the respondents with a negative K-10 plus (n = 6885, of whom 506 participated in the study) and 186 among the respondents with a positive K-10 plus (n = 4592, of whom 1023 participated). Recalculated to the respondents who completed the K-10, there are 24 (95% CI 0 to 57) antidepressant users with a definite or possibly justified reason in the screen-negative group and 835 (95% CI 726 to 943) in the screen-positive group.

All 11 respondents who used an antidepressant without a definite or possible justified reason had a positive K-10 plus. The total number of antidepressant users without a justified reason among all respondents with a positive K-10 plus can be recalculated at 49 (95% CI 20 to 78). As there were no antidepressant users without indication with a negative K-10 plus, the total number of respondents who used an antidepressant without indication among all 10,677 people who completed the K-10 plus, is also 49 (95% CI 20 to 78). K-10 plus results were unknown for one respondent without antidepressant use.

Combining the results of antidepressant users with and without justification, recalculated to the sample of 10,677 GP patients from 63 GPs who had consulted their GP in the past four months irrespective of reason for consultation and did return a completed screening questionnaire, 908 (8.3%), (95% CI 823 to 994), used an antidepressant of whom 859 (94.6% of the antidepressant users) with a definite (n = 494, 47.8%) or possible (n = 425, 46.8%) justification in accordance with the guideline and 49 patients (5.4% of the antidepressant users) without justification.

Table 3 shows the relation between severity of depression at baseline and indication for treatment with an antidepressant, based only on depression diagnosis.

<table>
<thead>
<tr>
<th>Severity of depression at baseline</th>
<th>Definitely justified AD users (n,%),</th>
<th>Possibly justified AD users (n,%),</th>
<th>Unjustified AD users (n,%),</th>
<th>Total (n)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MDD Single Mild</td>
<td>7 (24%)</td>
<td>6 (21%)</td>
<td>16 (55%)</td>
<td>29</td>
</tr>
<tr>
<td>MDD Recurrent Mild</td>
<td>12 (39%)</td>
<td>19 (61%)</td>
<td>N/A</td>
<td>31</td>
</tr>
<tr>
<td>MDD Single Moderate</td>
<td>19 (54%)</td>
<td>7 (20%)</td>
<td>9 (26%)</td>
<td>35</td>
</tr>
<tr>
<td>MDD Recurrent Moderate</td>
<td>9 (64%)</td>
<td>5 (36%)</td>
<td>N/A</td>
<td>14</td>
</tr>
<tr>
<td>MDD Single Severe</td>
<td>11 (33%)</td>
<td>8 (24%)</td>
<td>14 (42%)</td>
<td>33</td>
</tr>
<tr>
<td>MDD Recurrent Severe</td>
<td>17 (53%)</td>
<td>15 (47%)</td>
<td>N/A</td>
<td>32</td>
</tr>
<tr>
<td>Dysthymia</td>
<td>N/A</td>
<td>8 (100%)</td>
<td>0 (0%)</td>
<td>8</td>
</tr>
<tr>
<td>mID/depressive symptoms</td>
<td>N/A</td>
<td>N/A</td>
<td>7 (100%)</td>
<td>7</td>
</tr>
<tr>
<td>Total</td>
<td>75</td>
<td>68</td>
<td>46</td>
<td>189</td>
</tr>
</tbody>
</table>

N/A = not applicable.

Respondents with a recurrent MDD and possibly justified AD use had a possible justification because they had used an antidepressant for more than 2 years.

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Discussion

Summary of main findings

The main finding of our study is that overtreatment with antidepressants did not appear to be a very frequent problem in Dutch primary care. Of all GP patients (and after excluding patients who were treated in secondary care including hospitals, institutes for mental health care and psychiatrists with private practices) 8.5% received an antidepressant. When compared with the guidelines of the Dutch General Practitioner’s Association (NHG), 94.6% of the antidepressant use was in accordance with the guideline (47.8% definitely and 46.8% possibly) while only 5.4% was not. The latter was for the large part not due to treatment of mild forms of depression or patients without psychiatric diagnoses, but to non-justified long continuation of antidepressant treatment in patients who at some earlier point in their life had a justified reason for treatment with an antidepressant.

Strengths and limitations of the study

The current study has several very strong points. First, we used a screening method to recruit participants which did not affect the awareness of patient’s psychiatric status for GPs in our study. This means that the GPs could only rely on their own diagnostic judgments also for their prescription of antidepressants. The second strength of this study is its large sample size, which is rather rare in a primary care study. The third strength is that all patients were diagnosed based on a structured interview and not on the GPs’ records.

However there are also limitations. First, the last mentioned strength is also a weakness, as the structured interview we used (the CIDI) does not assess the degree of suffering and dysfunction, which should be part of the GPs’ consideration for antidepressant treatment according to the guideline recommendations. Second, the representativeness of the population may be limited. The GPs in this study, and thus their patients, may not be representative for all Dutch primary care practices, as these practices/GPs agreed to...
Comparison with existing literature

Several previous studies also looked at rates of overtreatment, with various outcomes. The major contrast with our study is that these studies only looked at relative small groups of GP patients and did not allow recalculation of the results to the total population of GP patients. Two of the previous studies found high rates of overtreatment: 25% (Sihvo et al.) and even 35% (Berardi et al.).[17,18] Berardi et al. described a group of 361 primary care patients of whom 82 used an antidepressant. They only considered treatment with antidepressants indicated for current depression, ignoring possible continuation or maintenance treatment and other indications for the use of antidepressants like anxiety disorders. This is a rather limited definition, as continuation treatment is a well-established part of depression treatment and many antidepressants are also registered for anxiety disorders. Sihvo et al. described a group of 526 patients who used an antidepressant. They adopted a definition of overtreatment slightly broader than ours. Treatment was considered “non-psychiatric” in case there was no CIDI diagnosis of MDD, Dysth, anxiety disorder (generalised anxiety, social phobia, panic disorder or agoraphobia), bipolar disorder or alcohol dependence in the last 12 months.

The third previous study (Cameron et al) reported a small percentage (exact percentage not mentioned) of overtreatment in a Scottish primary care sample of 120 antidepressant users.[19] They did not have a diagnosis based on a structured interview, but used a cut-off score of the Hospital Anxiety and Depression Scale (HADS). Of their sample 45 had “no depression”, 34 had a “possible depression” and 41 had “probable depression.” In the respondents with “no depression”, they reported that 32 had a history of depression (as recorded by the GP), 5 had anxiety disorder, 5 had neuropathic pain and for only 3 it remained unclear why they received an antidepressant.

It can be argued that our definition of non-justified antidepressant use is rather small, and that in reality more patients in our sample did not actually need an antidepressant. First, we defined lifelong treatment non-justified only for patients without a definite or possible justification. This created the problem of how to classify the category ‘possibly justified’ for treatment beyond two years of patients with a recurrent depression, while most guidelines recommend several years and only for a (non-specified) subgroup lifelong treatment. Second, also in case of anxiety disorders it could be argued that lifelong treatment is unnecessary in many if not most cases. Third, the diagnosis chronic MDD was based on the self-report of 24 months (of probably uninterrupted) symptoms of depression in the past five years according to the life chart and a lifetime diagnosis of MDD. It could very well be that these patients had no chronic MDD, but just symptoms of depression during two or more years. This would mean that in more patients antidepressants would be unnecessary. When we would have classified all possible justified cases as overtreatment the percentage would indeed rise substantially: from 5.4% to 52.2%, which is much higher than Berardi et al. and Sihvo et al. and clearly illustrates the importance of a clear definition of overtreatment.

We think however that our definition including lifelong treatment for patients with recurrent depressive episodes is justifiable, considering the high recurrence rate of depression, especially after multiple episodes.[45,46] As discussed above, at least part of the respondents with a possible justification will have a very good reason for the use of an antidepressant. Moreover, there could be patients without a justified reason for the use antidepressants, for whom the physician (or the patient himself) found an antidepressant needed, e.g. because of residual symptoms after a single episode of MDD. One could also argue that, although not indicated according to the guideline, this is justified treatment.

If indeed the percentage of overtreatment is lower in the Netherlands than in other countries: what would be the explanation? A possible explanation is the difference in primary care systems. In the Netherlands, the GP is the gatekeeper to secondary and mental health care; patients need a reference from the GP before they can consult mental health care. Secondly,
Dutch GPs have been trained during the last years in how to diagnose and treat depression and anxiety disorders, as part of the implementation of the Dutch primary care guidelines. This also implies that our results probably cannot be generalized globally, as primary healthcare systems vary across countries.

We sought for the difference or be that antidepressant use in our study was based on drug-container inspection in most patients and on self-report in a minority of cases, while the Dutch Foundation for Pharmaceutical Statistics (SFK, www.sfk.nl) data are based on pharmacy prescription data. From previous studies it is known that many patients do not pick up their prescription or do not take their medication.[47,48]

In this study we focused on overtreatment with antidepressants, and therefore we looked at whether patients received an antidepressant without a justified reason. However, “justified” does not mean “needed”. Patients with a mild or even moderately severe episode of MDD do not necessarily need treatment (either an antidepressant or psychotherapy), although they do have a justified reason for treatment with an antidepressant.

New guidelines like the recent 2009 update of the Dutch multidisciplinary guideline for depression recommend to reserve antidepressants for patients with moderate to severe depression.[42] Therefore, an alternative interesting question is: who needs an antidepressant, and how many of the patients who need an antidepressant do actually receive an antidepressant. This however, was not the focus of our study as undertreatment of depression has already been the focus of many studies in the past.[11–16] Moreover, the NESDA study is not suitable for answering this question. It is a naturalistic study and part of the study population did not seek any help. It is therefore impossible to determine which patients are “undertreated” by their GP and which did not seek help for their psychological complaints.

Implications for future research and clinical practice

In conclusion, the current study provides a unique insight into the justification of the prescription of antidepressants in Dutch primary care. In contrast to the scarce literature, the rate of overtreatment with antidepressants in the present study was low.

Another interesting finding is that overtreatment is not so much due to treatment of mild forms of depression or patients without psychiatric diagnoses, but rather to an excessive duration of antidepressant treatment in patients with remitted (recurrent) MDD. This latter finding presents several implications for clinical practice. First, projects on optimizing treatment with antidepressants in primary care, should not focus on reducing overtreatment but on identifying patients who do not need antidepressants anymore. Second, GPs should be aware that apparently many patients tend to continue the antidepressant over many years. Which of these patients might be able to stop the antidepressant is unclear. Therefore, further studies addressing this question are warranted before starting campaigns to reduce the use of antidepressants in primary care.

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Author Contributions

Conceived and designed the experiments: EP KvM WAN. Performed the experiments: EP. Analyzed the data: EP KvM WAN. Wrote the paper: EP KvM WAN. Critically revised the design and critically revised the manuscript: WJGH BWJHP.


