Long term studies of depression: what is relevant for the physician?

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ABSTRACT – Following a meticulous review of long term studies of depression, we conclude that prolonging antidepressant medication after full recovery is clearly beneficial. In this article, we examine recent long term studies that reveal substantial evidence in this sense. However, we advise physicians to keep in mind that this prophylactic effect has been proven for a restricted population, i.e., patients showing recurrent depressive disorders with low comorbidity on axis 1, and having revealed a satisfying response to antidepressant agents during the acute phase of the index episode. This does not mean that antidepressant agents are not useful for patients showing other characteristics, but more studies are needed to assert this possible advantage. We also examine the limited data on chronic forms of depressive disorders.

The effect of long term prescription is believed to be advantageous as long as the medication is taken; it has been demonstrated for up to 5 years.

Full dosage is indicated even if the effect of active drug over placebo persists at lower doses. The differences between antidepressant agents appear minor and physicians should be more concerned about the long term tolerance of these drugs than their efficacy when choosing the appropriate medication for maintenance treatment. Physicians should also be aware of the greater risk of recurrence during the 6 months following the discontinuation of medication. This risk occurs regardless of the total length of prescription. The possibility that recurrence may be mistaken for withdrawal symptoms cannot be ruled out.

Finally, the side-effects of antidepressant drugs are a major concern, particularly when extending the length of prescription. Even though the newer generation medications dis-
play a more favorable short-term side-effect profile, the effects of chronic use of these agents are still unclear. Therefore, the decision to extend treatment over several years requires comprehensive discussion with patients and cautious clinical monitoring to identify potential late-onset side effects.

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**Introduction**

During the past decades, catamnestic studies have contributed to change in our view of depressive disorders. Psychiatrists now admit that a single depressive episode over a lifetime constitutes an exception rather than a rule\(^1,2\). The high rates of persistent morbidity, recurrence and death among patients suffering from depression\(^3\) have highlighted the value of long term psychological and pharmacological treatment.

Surprisingly, there is a large gap between consensual recommendations and the physician’s routine practice in long term treatment of depression. Therefore, physicians may be concerned by several questions related to the management of this disease, particularly during the maintenance phase.

This article is a meticulous review of recent long term studies of depression in which we will try to answer to several questions:

What evidence do we have that prophylactic treatment is effective?

How should we choose between individual drugs?

What is the optimal duration of drug prescription?

Finally, what is the risk of lasting prescription of antidepressant agents?

**Definitions**

First of all, it should be noted that all long term studies on depression concern one type of depression: major depressive disorders. Major depressive disorders are major depressive episodes touching patients that are not suffering from schizophrenia or bipolar disorders. Therefore, other forms of depression such as minor depression or subsyndromic episodes are not covered in these studies. In this review, we also looked at chronic forms of depression (see Results section, Chronic Depression paragraph) which are not included in the definition of major depressive disorders. However chronic forms of depression can raise similar problems of management.

In order to describe the course of depressive episodes, consensual definitions have been proposed for the different stages of the illness and their corresponding treatment phases\(^4\).

<table>
<thead>
<tr>
<th>Illness stage</th>
<th>Definition</th>
<th>Treatment phase</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acute phase</td>
<td>DSM IV criteria</td>
<td>Acute treatment phase</td>
</tr>
<tr>
<td>Remission</td>
<td>Improvement of sufficient amplitude on psychometric scales (see table 1)</td>
<td>Continuation phase</td>
</tr>
<tr>
<td>Full recovery</td>
<td>Definitive ending of the depressive disorder (remission &gt; n months)</td>
<td>Prophylactic or maintenance phase</td>
</tr>
</tbody>
</table>
A relapse is a new depressive episode which occurs after remission, that is, during the continuation phase.

A recurrence is a new depressive episode which occurs after full recovery, that is, during the maintenance phase.

Long term drug treatment includes continuation and maintenance treatment.

The difference between continuation and maintenance treatment might appear arbitrary and may not replicate underlying biological processes, but most authors believe that a four to six month remission period should occur before a recurrence is diagnosed. This issue will be thoroughly discussed later.

Consequently, current guidelines regarding treatment recommend four to nine months of continuation antidepressant therapy following the remission of acute symptoms to allow total resolution of an episode.

Though the practice is fairly homogeneous concerning the prescription of continuation treatment, the situation is very different for maintenance treatment. The exact indication, dosage and total duration of treatment is not consensual and a great disparity exists between recommendations and usual practice. In this context, it is therefore essential to discern what information can be construed from long term studies.

Methodological issues

Before analysing the results yielded by long term studies, a number of methodological differences need to be pointed out as methodological disparities between trials make interpretation difficult.

All long term studies share a first stage where subjects, after selection, receive treatment during the acute phase of their depressive episode. Selection biases are a major issue, which will be discussed in detail in section 5.

One of the most crucial points is the time after which randomisation is undertaken. Though all studies proceed by this necessary first phase where every patient entering the trial must achieve remission under treatment, only a limited number of these trials continue on to a second requisite phase of maintenance treatment. Thus randomisation can occur directly after the acute phase or after maintenance treatment. In the former case, recurrence and relapse are grouped together.

It is certainly problematical to determine when the continuation phase of treatment ends and when the true prophylactic phase begins, but examination of the point at which symptoms reappear in studies of long-term treatment can offer some guidance. Data from these studies show a similar pattern of relapse and recurrence, that is a greater rate of relapse (under placebo but also, to a lesser degree, with antidepressant agents) during the first months following remission. The data of Mindham et al., one of the first long term studies, account for a rate of relapse superior to 40% during the first 16 weeks. This number is consistent with the results of Prien and Kupfer, who retrospectively examined relapse and recurrence rates with placebo in an earlier study that continued for 2 years. The authors discovered a high rate of relapse in the first 16 weeks (44%), the peak rate taking place in the first 8 weeks (38%). Additionally, if one looks closely at the rate of relapse and recurrence in function of time in all long term studies, it appears that the patterns of relapse and recurrence are highly similar:
there is a first phase with a high rate of relapse, during approximately 5 to 6 months after remission. Then, the rate of recurrence tends to be much more stable. The first phase corresponds to the length of time defining the continuation phase.

Another important issue is the length of the treatment phases. In the acute phase of treatment for instance, some studies will consider all patients who have reached remission, regardless of the time taken to achieve this condition, whereas other studies will assign a fixed period (6 or 8 weeks for example) after which the assessment is completed and patients who do not meet the remission criteria will be excluded from the trial. This introduces a large bias since, as we will see later, initial response to treatment is a major prognostic of relapse and recurrence. For the trials that comprise a maintenance phase, the length of this period may also be variable, fluctuating from 15 to 24 weeks.

Finally, the last and perhaps most important controversy concerns the definition of remission, which is of course fundamental as this key criteria constitutes a “filter” through which patients are selected for the trial. This definition, like the definition of recurrence, can differ significantly between studies (Table I).

Results of published studies

Pharmacological nature of maintenance treatment

The first and probably the most astonishing result is a remarkably stable advantage of treatment over placebo in the prevention of recurrence. We can see in table 2 that treatment prevents roughly 50% of the recurrences that occur under placebo, regardless of the duration of the study or the pharmacological nature of the antidepressant drug. It is noteworthy that this constant result is not influenced by the presence or absence of the different biases we discussed previously.

We might therefore think that all antidepressant drugs are equivalent in the prevention of recurrent episodes. But we must keep in mind that most of these trials used the same drug during the entire study, which means that a given drug is appropriate for prophylaxis only if it has shown sufficient effectiveness to enable remission during the acute phase and, in most cases, to sustain this remission during the consolidation phase.

The different prophylactic effects between antidepressant agents has been directly analysed in studies comparing several drugs in the same trial (see Table III)

Claghorn et al.\(^8\) proposed one of the first long-term studies with two antidepressant drugs. Paroxetine and imipramine were compared to placebo after a first acute phase of 6 weeks where patients obtained remission. The next stage did not include any continuation phase. The follow-up lasted one year. Results showed a superior efficacy of both antidepressant drugs over placebo. However, the number of drop outs due to side effects was twice superior in the imipramine group, suggesting that SSRI are better tolerated, especially in long term prophylaxis.

Franchini et al.\(^9\) compared in double bind two SSRI, sertraline and fluvoxamine, on long term efficacy during a 24 month follow-up. Patients presenting recurrent major depressive episodes were first treated during the acute index episode in an open phase. The majority of patients (77%) received tricyclics, others benefited from SSRI, IMAO
Table I
Definition variation between long term studies.

<table>
<thead>
<tr>
<th>Studies</th>
<th>Depressive antecedents</th>
<th>Inclusion criteria</th>
<th>Remission criteria</th>
<th>Recurrence criteria</th>
</tr>
</thead>
<tbody>
<tr>
<td>Montgomery (1988)</td>
<td>≥ 1 MDE in the previous 5 years</td>
<td>HDRS &gt; 18</td>
<td>HDRS &lt; 12</td>
<td>HDRS &gt; 18</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>HDRS &lt; 8 to enter prophylaxis period</td>
<td></td>
</tr>
<tr>
<td>Frank (1990)</td>
<td>≥ 3 previous MDE</td>
<td>HDRS &gt; 15</td>
<td>HDRS &lt; 7</td>
<td>HDRS &gt; 15</td>
</tr>
<tr>
<td>Robinson (1991)</td>
<td>Antecedent of minor or MDE</td>
<td>HDRS &gt; 18</td>
<td>HDRS &lt; 10</td>
<td>HDRS &gt; 18</td>
</tr>
<tr>
<td></td>
<td></td>
<td>RSD &gt; 7</td>
<td>GAS &gt; 70</td>
<td>RSD &gt; 7</td>
</tr>
<tr>
<td>Kupfer (1992)</td>
<td>≥ 3 previous MDE</td>
<td>HDRS &gt; 15</td>
<td>HDRS &lt; 7</td>
<td>HDRS &gt; 15</td>
</tr>
<tr>
<td></td>
<td></td>
<td>RSD &gt; 7</td>
<td>RSD &lt; 5</td>
<td>RSD &gt; 7</td>
</tr>
<tr>
<td>Terra (1998)</td>
<td>≥ 2 previous MDE in the last 5 years</td>
<td>MADRS &gt; 25</td>
<td>MADRS &lt; 10</td>
<td>5 criteria in DSM 3R</td>
</tr>
<tr>
<td></td>
<td></td>
<td>RSD &gt; 7</td>
<td>CGI &lt; 2</td>
<td></td>
</tr>
<tr>
<td>Rouillon (2000)</td>
<td>≥ 2 previous MDE in the last 3 years</td>
<td>HDRS &gt; 18</td>
<td>HSRD &lt; 12</td>
<td>HDRS &gt; 18</td>
</tr>
<tr>
<td>Gilaberte (2001)</td>
<td>≥ 1 previous MDE in the last 5 years</td>
<td>HDRS 17 &gt;18</td>
<td>HDRS 17 &lt; 8</td>
<td>HDRS 17 &gt;18</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CGI &gt; 4</td>
<td>CGI &lt; 2</td>
<td>CGI &gt; 4</td>
</tr>
<tr>
<td>Hochstrasser (2001)</td>
<td>≥ 2 previous MDE, last one in the last 5 years</td>
<td>MADRS &gt; 22</td>
<td>MADRS &lt; 11</td>
<td>MADRS &gt; 22</td>
</tr>
<tr>
<td>Klysner (2002)</td>
<td>Elderly patients Unipolar MD (DSM 4)</td>
<td>MADRS &gt; 22</td>
<td>MADRS &lt; 11</td>
<td>MADRS &gt; 22</td>
</tr>
<tr>
<td>Wilson (2003)</td>
<td>Elderly patients Major depressive disorder (DSM 3R)</td>
<td>HDRS 17 &gt;18</td>
<td>50 % reduction in baseline HRSD</td>
<td>HDRS 17 &gt;13</td>
</tr>
<tr>
<td></td>
<td></td>
<td>AGECAT &gt; 3</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>MMSE &gt; 11</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lepine (2004)</td>
<td>≥ 3 previous MDE, last one in the last 4 years Most recent MDE within 6 months of study</td>
<td>Based on items of DSMIV and MADRS (sadness, depressed mood)</td>
<td>Based on items of DSMIV and MADRS (sadness, depressed mood)</td>
<td>DSM IV criteria, Clinician’s opinion</td>
</tr>
<tr>
<td>Reynolds (2006)</td>
<td>Elderly patients Major depressive disorder (DSM 4)</td>
<td>HDRS 17 &gt;17</td>
<td>HDRS 17 &lt; 10</td>
<td>HDRS 17 &gt;15</td>
</tr>
</tbody>
</table>

Abbreviations:
MDE: Major depressive disorder
HDRS: Hamilton Depression Rating Scale score; HDRS 17: 17 items Hamilton Depression Rating Scale score
MADRS: Montgomery and Asberg Depression Rating Scale score
RSD: Raskin Severity of Depression score
CGI: Clinical Global Impression score
AGECAT: Geriatric Mental State

A or combined drug treatments. After a four-month continuation phase and a 3-week wash-out period, patients were randomly attributed to one of the two treatment groups. Both treatments showed similar tolerance and survival rates.

Montgomery et al. compared mirtazapine and amitriptyline in a double-blind placebo-controlled study. Here the long term study design was entirely different as it was an extension of a double-blind placebo-controlled study of acute treatment. After
remission, patients were given the possibility of continuing the same treatment for up to two years. Authors noted an advantage of mirtazapine over amitriptyline in preventing relapse in the survival analysis, but not in the number of relapses. It should be noted that there was an equivalent number of withdrawals in the two groups due to side effects, and that both mean dosages of amitriptyline and mirtazapine were relatively low (137.5 mg and 22.8 mg respectively), possibly sub-optimal.

Finally, Bump et al.\textsuperscript{11} conducted a study with elderly patients, comparing nortriptyline to paroxetine. Here, patients were randomly assigned to one of the two treatment groups during the acute phase. After this first stage, patients were offered the opportunity to continue in an open continuation and maintenance treatment for an additional 18 months open trial. Preliminary data showed that both paroxetine and nortriptyline had comparable efficacy in preventing relapse and recurrence, and time before relapse\textsuperscript{12}.

Table II
Recurrence rates in long term studies.

<table>
<thead>
<tr>
<th>Studies</th>
<th>Number of subjects</th>
<th>Length of treatment phase</th>
<th>Drug</th>
<th>Recurrence rate</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Acute</td>
<td>Contin</td>
<td>Mainten</td>
</tr>
<tr>
<td>Montgomery (1988)</td>
<td>456 initial 220 maint 182 complete</td>
<td>6 weeks</td>
<td>18 weeks</td>
<td>1 year</td>
</tr>
<tr>
<td>Frank (1990)</td>
<td>128 maint 106 complete</td>
<td>Until remission</td>
<td>17 weeks</td>
<td>3 years</td>
</tr>
<tr>
<td>Robinson (1991)</td>
<td>88 initial 47 maint 35 complete</td>
<td>Until remission</td>
<td>16 weeks</td>
<td>2 years</td>
</tr>
<tr>
<td>Kupfer (1992)</td>
<td>20 maint 19 complete</td>
<td>(3-)5 years 6 weeks</td>
<td>18 weeks</td>
<td>1 year</td>
</tr>
<tr>
<td>Terra (1998)</td>
<td>436 initial 204 maint</td>
<td>6 weeks</td>
<td>18 weeks</td>
<td>1 year</td>
</tr>
<tr>
<td>Rouillon (2000)</td>
<td>500 initial 214 maint 166 complete</td>
<td>Until remission</td>
<td>4 months</td>
<td>1 year</td>
</tr>
<tr>
<td>Gilaberte (2001)</td>
<td>253 initial 145 maint 121 complete</td>
<td>8 weeks</td>
<td>24 weeks</td>
<td>48 weeks</td>
</tr>
<tr>
<td>Hochstrasser (2001)</td>
<td>427 initial 269 maint 264 complete</td>
<td>6-9 weeks</td>
<td>Until remission</td>
<td>16 weeks</td>
</tr>
<tr>
<td>Lepine (2004)</td>
<td>371 initial 288 maint 165 complete</td>
<td>4-6 months followed by a 2 months placebo period</td>
<td>18 months</td>
<td>Sertraline</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Studies</th>
<th>Number of subjects</th>
<th>Length of treatment phase</th>
<th>Drug</th>
<th>Recurrence rate</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Acute</td>
<td>Contin</td>
<td>Maintain</td>
</tr>
<tr>
<td>Montgomery (1988)</td>
<td>172/141-135</td>
<td>8 weeks</td>
<td>1 year</td>
<td>Paroxetine</td>
</tr>
<tr>
<td>Versiani (1999)</td>
<td>283 initial 143 maint</td>
<td>6 weeks</td>
<td>23+23 weeks</td>
<td>Riboxetin</td>
</tr>
<tr>
<td>Studies</td>
<td>Number of subjects</td>
<td>Length of treatment phase</td>
<td>Drug</td>
<td>Recurrence rate</td>
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</tr>
<tr>
<td></td>
<td></td>
<td>Acute</td>
<td>Contin</td>
<td>Mainten</td>
</tr>
<tr>
<td>Reynolds (1999)</td>
<td>41 complete</td>
<td>Until remission</td>
<td>16 weeks</td>
<td>3 years</td>
</tr>
<tr>
<td>Klysner (2002)</td>
<td>230 initial</td>
<td>8 weeks</td>
<td>16 weeks</td>
<td>&gt; 48 weeks</td>
</tr>
<tr>
<td>Wilson (2003)</td>
<td>318 initial</td>
<td>8 weeks</td>
<td>16-20 weeks</td>
<td>100 weeks</td>
</tr>
<tr>
<td>Reynolds (2006)</td>
<td>195 initial</td>
<td>Until remission</td>
<td>16 weeks</td>
<td>2 years</td>
</tr>
</tbody>
</table>

Abbreviations:
- Initial: number of patients included in the study
- Maint: number of patients entering maintenance phase
- Complete: number of patients completing the study
- Conti: continuation phase
- Maint: maintenance phase

Table III
Long Term Studies Comparing the Effect of Different Antidepressants.

<table>
<thead>
<tr>
<th>Studies</th>
<th>Number of subjects</th>
<th>Length of treatment phase</th>
<th>Drugs</th>
<th>Relapse-Rec. rate</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Claghorn (1993)</td>
<td>717 219 conti</td>
<td>6 weeks</td>
<td>1 year double-bind phase</td>
<td>Paroxetine 15 %</td>
<td>Parox. and Imip. &gt; placebo 50% more drop-out with Imip. / Parox due to side effect.</td>
</tr>
<tr>
<td>Franchini (1997)</td>
<td>77 64 conti</td>
<td>4 months</td>
<td>24 months</td>
<td>Fluvoxamine 18.7%</td>
<td>No significant difference between two treatments</td>
</tr>
<tr>
<td>Montgomery (1998)</td>
<td>6 weeks up to 2 years Randomisation during all treatment phases</td>
<td>Fluo</td>
<td>Sertraline 21.9%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bump (2001)</td>
<td>116 59 conti Elderly patients</td>
<td>12 weeks</td>
<td>12 weeks 18 months</td>
<td>Randomisation during all treatment phases</td>
<td></td>
</tr>
</tbody>
</table>

The final results suggested that even if the efficacy of these two drugs was comparable, nortriptyline produced fewer residual depressive symptoms and side-effects.

All together, these results suggest that the prophylactic efficacy of all the antidepressant drugs tested is comparable; the main difference resides in the pattern of toler-
ance, which appears better with newer antidepressant agents.

**Dosage of maintenance treatment**

Another important issue is the determination of the drug dosage necessary to obtain an effective prophylactic effect. The current consensus is to employ the same medication dosage that allowed remission during the acute phase\textsuperscript{13,14,15}. The studies performed by Frank et al.\textsuperscript{13,14} aimed to assess the benefit of full dose maintenance therapy. In a first study (1990) the authors successfully demonstrated that full doses of antidepressant agents were more appropriate than the conventional low doses stated in previous studies\textsuperscript{16,17}. In a second study (1993) Frank et al. directly compared imipramine’s prophylactic effect at different dosages in a randomized study. Although the small size of the sample prohibited any definite conclusion, the authors suggested that the full dose of imipramine was more effective than the half dose in preventing recurrences.

Franchini et al.\textsuperscript{18} explicitly assessed this question by comparing the prophylactic effects of paroxetine at two different dosages, without any placebo group. Their results showed that paroxetine had a much better prophylactic effect at 40 mg than at 20 mg. Unfortunately, all the patients in this study were given the same dose of 40 mg to attain remission. A considerable bias was therefore introduced since some patients could perhaps have achieved remission with lower doses of paroxetine, but were randomised in the high dosage paroxetine group for the third stage analysis.

Also noteworthy are the data resulting from a large naturalistic study performed by the NIMH collaborative study of the psychobiology of depression\textsuperscript{19}. The results of this study suggest that the most effective treatment, not considering patients suffering from highly recurrent depressive disorders (i.e. more than 5 previous episodes), is a full dose treatment of limited duration, around 8 months. Of course the observational design of this study limits the extent to which cause and effect associations can be deduced. Nevertheless, these results also raise an important methodological issue: though the most important factor in the risk of recurrence is a history of previous depressive episodes, most long term trials do not distinguish patients based on this criterion. Therefore, patients having 5 or 6 previous episodes are assimilated with patients having only 2 previous episodes. Furthermore, the definition of recurrent depressive disorder may vary considerably between studies (Table I).

However, other studies showed contradictory results: Mindham et al.\textsuperscript{6} found that doses of amitriptyline lower than those used in the acute phase were effective in continuation treatment, but this was not the case for imipramine.

Montgomery et al.\textsuperscript{20} showed that doses of 20 or 40 mg of Citalopram were similarly effective in preventing relapse.

There are only a few randomized clinical trials in which the maintenance efficacy of different doses of antidepressant agents was compared (Table IV).

Rouillon et al.\textsuperscript{21} suggested that a lower than standard dose of maproptiline could be effective in preventing recurrences. Nevertheless, the lowest dose was less effective than the half-standard dose and the full antidepressant dose was not tested in this study, leaving aside the possibility that the latter could be the optimal dose.
Finally, the question of the total duration of the prophylactic treatment is still undecided. For many authors this treatment should be prescribed for life in the case of patients showing a high recurrence of depressive illness and/or severe characteristics. This issue cannot be directly corroborated by long-term studies as most of them last a maximum of two years. However, Kupfer et al.\textsuperscript{15} were able to obtain data on the prophylaxis of imipramine treatment after three years of maintenance treatment. This study was the extension of the Pittsburgh study of maintenance therapies in recurrent depression\textsuperscript{13}. Patients who completed this last three year trial by receiving active medication were asked to continue a two year additional randomized trial of active medication versus placebo. Despite the low number of subjects, results showed a clear advantage of imipramine over placebo in the prevention of recurrence, thus suggesting that medication can reduce the risk of recurrence as long as it is prescribed. It is also interesting to note that even after three years of active medication, patients who were randomised to placebo, thus discontinuing active medication, showed a maximum risk of recurrence in the first six months of the trial. This latter phenomenon can be explained in different ways. The first would be to consider that active medication provides effective protection for patients vulnerable to depression. Stopping the treatment thus raises the risk of early recurrence. We can also imagine that long term treatment also provides a “suggestive” effect, leading patients who stop medication to feel more vulnerable. Another possibility is that antidepressant drugs provide only a symptomatic action, which enables patients to overcome depressive episodes with less suffering (given that the standard duration of a depressive episode is about six months). For patients showing chronic depression, the discontinuation of long term treatment leads to the re-emergence of symptoms that had been contained by the drug. Finally, the possibility of withdrawal symptoms due to drug discontinuation should also be considered. Withdrawal symptoms have clearly been produced with tricyclic agents\textsuperscript{22} and more recently with SSRI\textsuperscript{23}.

### Chronic depression

Another complex clinical situation is the management of patients presenting chronic...
forms of depression. In the literature reviewed, three types of chronic depression can be distinguished:

- chronic major depressions concern patients presenting the criteria for major depression during at least two years,

- dysthymia refers to patients showing mild depressive characteristics (not fulfilling criteria for chronic major depression or major depression with incomplete remission) which last for at least two years,

- finally, double depression is an episode of major depression which occurs in patients who are suffering from dysthymia.

Three studies have explored these problematic conditions (Table V). Kocsis et al.24 focused on patients suffering from chronic forms of depression. In their studies, patients showing chronic major depression, “pure” dysthymia or dysthymia with current major depression (“double depression”), entered a long term trial including a ten week acute phase with desipramine followed by an open treatment continuation phase of 16 weeks. Patients sustaining remission were then randomised to placebo or active medication for up to two years. It should be noted that in contrast to other long term trials, patients showing only partial remission after the first phase (defined by a reduction of at least 50 % on the baseline HAM-D score) could also enter the second and third phases. The results of this study showed a much higher recurrence rate for partial responders (20% for patients under medication and 40 % for patients under placebo). Patients sustaining full remission after the maintenance phase displayed a 56 % risk of recurrence under placebo and only a 14 % risk when taking medication. Unfortunately, these risks were not singled out for the different types of chronic depression.

Table V
Recurrence Rates in Chronic Depression.

<table>
<thead>
<tr>
<th>Studies</th>
<th>Number of subjects</th>
<th>Length of study</th>
<th>Drug</th>
<th>Recurrence rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kocsis (1996)</td>
<td>129</td>
<td>2 years</td>
<td>Desipramine</td>
<td>11% 52%</td>
</tr>
<tr>
<td>Keller (1998)</td>
<td>161</td>
<td>1 year and a half</td>
<td>Sertraline</td>
<td>26% 50%</td>
</tr>
<tr>
<td>Gelenberg (2003)</td>
<td>165</td>
<td>1 year</td>
<td>Nefazodone</td>
<td>30.3% 47.5%</td>
</tr>
</tbody>
</table>

Keller et al.25 conducted a similar trial, this time selecting only patients suffering from a chronic form of major depression (chronic major depression or major depression with dysthymia (double depression). The trial consisted of three phases, but here the acute and continuation phases were also submitted to randomisation. A total of 161 patients successfully responded after a 12 week acute phase and a four month continuation phase under sertraline. Patients were then randomized to the same treatment or to placebo for 76 weeks. Sertraline allowed a significantly greater prophylaxis against placebo and the authors noted that depressive symptoms re-emerged twice as frequently with placebo than with sertraline. The astonishingly low recurrence rate of 23 % in the placebo group raised certain questions. The authors replied that chronic forms of major depression probably represent a lower risk factor of developing subsequent depressive episodes than a history of prior recurrent major depressive disorders. They also sug-
gest that SSRI could lead to fewer withdrawal symptoms than tertiary amine tricyclics; these withdrawal symptoms can be mistaken for a recurrent depressive disorder and explain the increased rates of recurrence in the placebo treatments of other studies. Lastly, it should be noted that the recurrence criteria used in this study were rather drastic compared to other long term studies.

Finally, Gelenberg et al. recently studied patients presenting chronic forms of major depression (chronic major depressive disorder, “double depression” or recurrent major depressive disorders with incomplete inter-episode recovery). Here again the acute phase was submitted to randomization. Only patients responding sufficiently to active medication were asked to prolong the same treatment during a continuation phase. Following this stage, and in the case of stable thymic response, patients were once again randomized in a 52 week double-blind maintenance phase of nefazodone or placebo. The authors found a significant difference between treatment and placebo in the probability of recurrence, but only when applying a competing-risk model. This model took into account, for instance, the unusual number of patients under placebo who interrupted the trial for reasons other than recurrence.

All of these studies suggest that maintenance treatments also play a significant role in the management of chronic depression. However, chronic forms of depression probably constitute a less important risk factor of recurrence than a history of multiple depressive disorders. The major issue in chronic depression is more probably determining how to obtain a satisfying initial response, rather than defining the prescription duration of antidepressant agents.

The links between major depressive disorders and chronic forms of depression raise an interesting questioning. There may be several ways of considering these links:

A first way is to consider chronic depression as the unfavorable outcome of a depressive episode. Comforting this view, interesting literature on residual symptoms can be quoted. Indeed, since the remission state corresponds to a precise score on psychometric scales, it is possible that some patients are considered as remitted according to these scores, even thought they still present a persistent level of symptoms. However these persistent symptoms may not justify a diagnosis of chronic depression or of dysthymia. Several studies have shown that the presence and variability of these residual symptoms were associated with a higher risk of recurrence. These findings thus suggest that there may be a continuum between “complete” remission and chronic depression.

Another approach is to consider that chronic depression constitutes a different form of illness than major depressive disorders. This would be apparent in the outcome and evolution of these different disorders but also in their respective etiopathology. For instance it appears that comorbid personality disorders and vulnerable cognitive processes are much more frequent in the case of chronic depressions. Along this line of thinking, it could be interesting to examine the influence of comorbid personality disorders on the rate of remission. Another major issue is to evaluate the benefit of psychotherapy in chronic depression according to the presence of personality disorders. For instance, a study performed by Nemeroff et al. shows that psychotherapy, compared to antidepressant agents, allows a higher rate of remission in chronic depression with childhood trauma. This is not the case for chronic depression without childhood trauma.
Are these results applicable to usual practice?

Value of the initial response

These results enable us to assert several points:

– patients suffering from recurrent depressive disorders (the patients usually selected in these studies) benefit from a prescription of prophylactic treatment;

– this benefit seems to be approximately the same for all pharmacological treatments;

– this benefit is present as long as the drug is prescribed.

However, due to the selection bias of these trials, these results are limited to two conditions. First, only patients with a high risk of depressive recurrence will benefit from prophylactic treatment since the main risk factor retained in the long term studies is a history of several depressive occurrences. Second, in order to benefit from the prophylactic effect, patients must also show a sufficient initial response to antidepressant drugs since patients not obtaining remission are excluded from these trials.

Initial response to antidepressant treatment is of considerable importance. Stewart et al. showed that only patients showing a “true drug” initial response (which is delayed and persistent) benefit from a continuation and maintenance treatment by active medication rather than placebo. Patients showing a “placebo” initial response (which is early and not persistent) had a comparable outcome whether continuing with placebo or active medication. Additionally, patients with a placebo pattern of response relapsed more often, regardless of the nature of treatment used (placebo or active drug). These findings have been confirmed by a recent study in this trial, 410 patients were treated with mirtazapine and were then randomized after remission to the same dose of mirtazapine or switched to placebo for the maintenance phase. The relapse rate for ‘true-drug in initial response pattern’ patients switched to placebo was significantly higher than for ‘placebo initial response pattern’ patients switched to placebo. Moreover, patients with a ‘true-drug initial response pattern’ relapsed significantly less when continuing mirtazapine rather than placebo. This was not the case for patients with a ‘placebo initial response pattern’.

In a study of elderly patients performed by Dew et al., the temporal pattern of initial response to acute treatment predicted the risk of recurrence. Thus, for patients showing a prolonged absence of response, maintenance treatment did not show any advantage whereas patients showing rapid initial response had a lower risk of recurrence when maintained under treatment rather than under placebo.

Comorbidity

The main criticism of long term trials is that “ideal” patients, who may not reflect the individual situations confronted by physicians, are often selected. This factor can be particularly relevant in the case of depressive disorders.

First of all, patients suffering from depressive disorders often show a high comorbidity with other psychiatric conditions. Sanderson et al., for example, found that about two-thirds of a panel of 260 patients with a principal diagnosis of depressive disorder had a least one additional coexisting axis 1 disorder. The most common comorbidity diagnosis is anxiety disorder. More recently, Zimmerman et al. observed that more than half
of a large group of depressed outpatients met the full criteria for a current anxiety disorder. Some long term studies include patients presenting a general anxiety disorder, but they usually exclude patients suffering from social phobia, obsessive disorders or other anxiety disorders.

Of course, anxiety disorders are not the only type of depressive comorbidity, and though some associated anxiety disorders may be included, addictive disorders are systematically ruled out. Nevertheless, substance abuse disorders play a significant role in the associated comorbidity of depression. Spaner et al.38 showed that nearly one third of individuals with an affective disorder also met the criteria for alcohol dependence. This proportion obviously increases when all types of substance use disorders are considered39.

Finally, it should be noted that medical comorbidity is also a frequent condition that is left out of trials. This association could lead to a more severe symptomatology and treatment resistance40. In a retrospective study of medically ill subjects treated with antidepressant medication for major depression, Popkin et al.31 obtained response rates as low as 40%.

Physicians should therefore keep in mind that the results of the long term trials are valid for a specific population, which includes only patients suffering from depressive disorder with low or no comorbidity.

What is the risk of long term treatment?

The issue of side-effects has dwindled from its position at the forefront of depression management since the introduction of new antidepressant agents. However, the consensual recommendations that tend to extend the indications and length of antidepressant prescriptions should also encourage a thorough inspection of short-term as well as late appearance side-effects. Numerous studies have confirmed the safer profile and the lower occurrence of side effects claimed for newer compounds42. Unlike TCAs, SSRIs induce no, or fewer, anticholinergic, hypotensive or sedating side-effects. The most common side-effects reported include nausea, headache, vomiting, nervousness, insomnia and sexual dysfunction. These effects are usually moderate and often diminish after the first weeks of treatment (for review see43). Dual action antidepressants are also better tolerated than tricyclic antidepressants44. As with SSRIs, they do not present the autonomic inconvenience of tricyclics. But here again, sedation, nausea, sexual dysfunction and weight gain are sometimes reported when using these treatments.

Anderson & Tomenson45 conducted a meta-analysis comparing the discontinuation rates with SSRIs and TCA. This work showed that the drop-out rate due to side-effects was 25% lower with SSRI than with TCA, whereas the total discontinuation rate was only 10% better with SSRIs. More recent meta-analyses46,47 tend to nuance this finding and seem favourable to a more modest advantage for tolerance of SSRIs compared to TCA.

Managing these side effects is all the more important since they play a major role in adherence to pharmacotherapy. Several studies have shown that adequate information concerning side-effects and good control could allow considerable compliance increase48. In addition, respect of guidelines for depression treatment has been shown to
play a major role in reducing the risk of relapse and recurrence\textsuperscript{49}.

In addition to these commonly described side-effects, physicians should be aware of the more dangerous consequences of these pharmacological treatments on the course of the depressive illness. The induction of maniac/hypomaniac symptoms for instance, is a classic risk that is poorly documented. It has been estimated at less than 1\% for strictly unipolar depression, but other authors suggest this risk could attain 10\%.\textsuperscript{50} Additionally, some manifestations usually attributed to antidepressants, such as insomnia, nervousness or irritability, could represent some subclinical form of hypomania. Finally, antidepressant agents have also been incriminated in various behavioral changes; though these findings are not substantiated, it has been suggested that these drugs could play a role in inducing suicidal behavior\textsuperscript{21}.

It is true that little data exists on the long-term consequences of antidepressants, especially concerning the newer compounds. Apart from the possible modifications of sleep structure, biorhythm and hepatic metabolism, neurobiological changes after chronic use of such agents largely challenge our knowledge. The possibility that distinct patterns of pharmacologic modifications may appear after several years of treatment can not be ruled out and some authors have even suggested that opposing effects could occur in some patients after chronic use of these drugs, leading to an aggravation of the course of depressive illness\textsuperscript{51}.

**Conclusion**

Long-term studies provide substantial aid to physicians in the management of recurrent depressive disorders, especially since no consensual guidelines exist on this specific topic.

Following a meticulous review of long term studies of depression, we conclude that prolonging antidepressant medication after recovery is clearly beneficial. The effects of long term prescription are advantageous as long as the medication is taken; it has been demonstrated for up to 5 years. The dosage used in maintenance treatment is the same as that which was effective during the acute phase. Even if the dosage of the antidepressant drug is diminished, it is still more effective than placebo treatment, but to a lesser extent. Physicians should therefore pay particular attention to the long term tolerance of these drugs, probably more so than to their long term efficacy, since the difference in efficacy between drugs seems to be minor.

Considerable efforts to standardize the methods applied in recent trials allow finer comparisons and lead to more reliable conclusions. Nevertheless, these studies concern a limited population and their value is diminished by a lack of data concerning people with comorbidity axis 1 and axis 3 disorders, children and adolescents. The focus of future research should be expanded to include the potential late onset adverse events and withdrawal phenomena that can result from the chronic use of antidepressant drugs.

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