Remission and relapse in the ambulatory treatment of patients with schizophrenia. Outcomes at 3 years

Introduction. Three year data collected in the frame of the SOHO study within Spain were used to evaluate antipsychotic treatment outcomes by analyzing remission and relapse as well as the factors influencing them.

Methods. The SOHO was a prospective, long-term, observational study of the outcomes of schizophrenia treatment in ambulatory who initiated therapy or who changed to a new antipsychotic drug performed in 10 European countries, with a focus on olanzapine. This article reports the attainment of international schizophrenia clinical remission and relapse criteria and the associated correlates in these patients.

Results and conclusions. A total of 2,020 patients were recruited in Spain. Almost 2/3 (60.1%) of the patients met the criteria for clinical remission. Factors that influence the likelihood of remission were identified, such as gender, baseline clinical and/or functional status, time since treatment initiation, treatment with olanzapine versus oral typical antipsychotics, duration of treatment, gender or the need for concomitant anxiolytics. Relapse occurred in 18.7% of patients. Treatment with quetiapine or the prescription of anticholinergics was associated with a greater risk of relapse.

Conclusions. These results highlight some prognostic factors of the course of schizophrenia and underscore the importance of the antipsychotic choice and its maintenance to achieve favorable long-term clinical outcomes in routine practice.

Key words:

Originals

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Remisión y recaída en el tratamiento ambulatorio de los pacientes con esquizofrenia. Resultados a 3 años

Introducción. Los datos del seguimiento de 3 años realizado para el estudio SOHO en España se han empleado para evaluar los resultados del tratamiento antipsicótico en términos de incidencia y factores asociados a la remisión y la recaída.

Métodos. El SOHO fue un estudio observacional, prospectivo, longitudinal realizado en 10 países europeos sobre los resultados del tratamiento de la esquizofrenia en pacientes ambulatorios que iniciaron o modificaron su farmacoterapia antipsicótica. El presente artículo presenta la incidencia y los factores asociados a la remisión y recaída clínicas de la esquizofrenia (definidas según los criterios internacionales al uso) en la muestra española.

Resultados. En España se reclutaron 2,020 pacientes. Casi dos tercios (60,1%) cumplieron los criterios de remisión clínica. Se identificaron varios factores relacionados con la probabilidad de remisión, tales como el sexo, el estado clínico y/o funcional, la edad de inicio del tratamiento, el tipo de antipsicótico, el uso de concomitantes ansiolíticos. La recaída se produjo en un 18,7% de los pacientes. El tratamiento con quetiapina y la asociación de anticolinérgicos se asociaron con un mayor riesgo de recaída.

Conclusions. Los resultados de este estudio señalan algunos factores asociados a la evaluación de la esquizofrenia y subrayan la importancia de una correcta elección del fármaco antipsicótico y su mantenimiento para lograr un resultado clínico favorable a largo plazo en la práctica clínica habitual.

Palabras clave:
INTRODUCTION

The efficacy and safety of atypical antipsychotics for the treatment of schizophrenia have been well established thanks to the results of several experimental studies and these have been considered for the up-dating of the international treatment guidelines. However, there is still no final consensus on the role that this group of drugs has for the clinician in the actual clinical practice. Translating the results of the randomized clinical trials into routine clinical practice is hindered by the inherent limitations of these studies, such as strict selection criteria, small sample sizes or short follow-up periods. Given the complexity of schizophrenia, the gap between experimental clinical research and routine practice is even wider than in other chronic disorders; but, to date, the efforts made to perform observational studies aiding to close such a gap have been scarce. In general, the naturalistic studies conducted have methodological limitations that affect their external validity, for example, cross-sectional or retrospective designs, reduced sample sizes or lack of appropriate control groups.

The improvement of therapeutic (pharmacologic and non-pharmacologic) options has helped to better characterize and control the disease course of schizophrenia. The classical view of a progressive evolution towards a chronic illness with substantial morbidity and persistent deficits in cognition and psychosocial function has given way to the more optimistic notion based on the long-term symptom stability as an attainable goal. Furthermore, long-term symptomatic stability is not only a possible outcome but may also be a foundation for functional improvement. In other words, maximum efficacy is achieved with psychosocial therapies and rehabilitation only when there is an adequate control of the symptoms.

The aims of this article were to assess in a large sample of outpatients with schizophrenia in Spain who initiated treatment with a new antipsychotic: a) the clinical outcomes of antipsychotic treatment in terms of remission and relapse, and b) the factors that influence these outcomes.

METHOD

Study design

The SOHO is a prospective, 3 year, observational study of the treatment of schizophrenia in Europe. It pays special attention to the outcomes obtained when the patient is treated with olanzapine. In Spain, 2020 patients were recruited between September 2000 and December 2001 and were followed-up by 86 psychiatrists from different settings (public or private, urban or rural) with a wide geographical distribution.

Those patients who initiated treatment with an antipsychotic for the treatment of schizophrenia, regardless of whether this substituted another previous one, was added to a previous treatment or if they had not been taking any antipsychotic medication previously were invited to participate. The patients should have been receiving ambulatory treatment or, if hospitalized, should not have been hospitalized for more than 15 days and it only should have been indicated for treatment change. The patients should be at least 18 years of age and not be participating in any clinical trial.

As the primary purpose of the study was to compare olanzapine with the other antipsychotics, two patient cohorts of similar size were recruited: a) those who initiated or changed to olanzapine therapy as single drug therapy and b) those who initiated or changed to or added a non-olanzapine antipsychotic therapy. Thus, the olanzapine group was over-sampled to obtain a sample in which approximately 50% of patients had initiated treatment with this drug.

To avoid interference with routine practice, the psychiatrists were instructed to make treatment decisions before and independently from assessing patients for enrolment, the recruitment period was purposely long, and the investigators were not committed to a minimum recruitment.

The ethics committees of the regional communities involved approved the protocol beforehand and obtained written informed consent from each participant to gather information before their recruitment.

The study protocol did not include any restriction regarding the patients’ treatment, neither before nor after their inclusion, this being left up to the discretion of the participating psychiatrists.

Study assessments

The data was collected during the routine visits occurring along the patients’ treatment. The intention was to obtain data at three and six months after the initiation/change of therapy that motivated the enrolment in the study and then every six months until the completion of the three-year follow-up period. In order not to reduce the impact on the usual practice, a period allowing for a range of one month before and one month after the foreseen dates to gather the information was established. Those patients who did not come to the routine visit within any of these periods were not withdrawn from the study, however, in such case, the corresponding assessment was left blank.

Three types of outcomes, clinical severity, quality of life and social functioning, were assessed. For the clinical severity, the Clinical Global Impression-Schizophrenia (CGI-SCH) scale was used. This instrument is an adaptation from the CGI scale to assess overall, positive, negative, depressive and cognitive symptoms at the time of the visit with physician-
rated scores ranging from 1 (not ill) to 7 (among the most severely ill patients)\(^4\). It requires only a few minutes to administer, thus having a minimal impact on the observational nature of the study.

The advisory board of the SOHO study agreed on the criteria of definitions of clinical remission and relapse based on these scores prior to knowing study results.

In agreement with the Andreasen et al.\(^3\,13\) criteria, remission was defined as an overall, positive, negative and cognitive score lower than or equal to 3 in 3 or 2 consecutive visits separated by at least six months in the absence of any in-between hospitalization.

Relapse was defined as the need for hospitalization or increase of at least 2 points in the overall CGI-SCH score that ends in a moderately severe or worse score. Those patients who did not achieve an overall CGI score equal to or lower than 3 throughout the 3 year period were excluded from the analysis of the relapses since patients who did not achieve clinical improvement could not relapse.

Given the importance of hospitalizations in cost-effectiveness evaluations, the occurrence of hospitalizations and their associated factors were analyzed separately.

Other data were collected on quality of life using the EuroQol-5 Dimensions (EQ5D), featuring the descriptive part and the visual analogue scale (VAS); social functioning with the social functioning scale, concomitant medications, compliance with treatment, violence/arrest episodes, health resource utilization, social and employment statuses, and appraisal of global activity. The results of these outcomes are outside of the scope of this article.

**Statistical analyses**

Treatment cohorts were defined based on the antipsychotic treatment initiated in the first visit, considering the following categories: olanzapine, risperidone, quetiapine, clozapine, other atypical antipsychotics, and any oral typical antipsychotic, depot typical antipsychotic, two or more antipsychotics. Incidences of remission and relapse and the factors influencing them were compared among study cohorts by means of multivariate analyses. Backward stepwise selection of variables with significant predictive ability was performed to abridge the models. Furthermore, time to each of these events was analyzed using the Kaplan-Meier method. Time to relapse was imputed by the mid-point between two consecutive visits for those patients who fulfilled the hospitalization criteria.

Analysis of the factors associated to achieving clinical remission during the follow-up period was made with a GEE (generalized estimating equation) logistic model. Presence of remission in each one of the intervals ranging from 6-12, 12-18, 18-24, 24-30 and 30-36 months was analyzed as dependent variables of the model. The medication that the patient was receiving at the first visit of each interval, taking olanzapine as the reference category, was considered to analyze the outcome. The model was also adjusted for baseline differences between the cohorts, taking the baseline status of each patient into account. The following clinical and sociodemographic clinical variables were included in the model as independent variables —demographic: gender; clinical: age at first treatment for schizophrenia, time since first treatment, alcohol or substance dependency and/or abuse, suicide attempts, overall positive, negative, depressive and cognitive CGI-SCH, hostility, treatment compliance, body mass index; social functioning: marital status, living independently, having paid employment, being socially active; side effects: extrapyramidal symptoms, amenorrhea; gynecomastia; impotence; galactorrhea, tardive dyskinesia; medication: use of concomitant medication (anticholinergics, antidepressants, anxiolytics, and mood stabilizers), and the visit in question— and the medication that the patient was taking at the beginning of each of the periods, defined as the last medication that had been prescribed to the patient.

Cox proportional hazards method was used to analyze the risk of relapse. The independent variables used were the antipsychotic (cohort) that the patient was taking in the last visit prior to the relapse, using the olanzapine cohort of reference and those that had been described in the logistic regression model on remission.

In addition, an independent Cox hazards risk model was prepared to investigate separately the factors associated to the hospitalization.

No a priori calculation was made of the sample sizes used in each one of these analyses because they were done using subgroups of patients from the principal study. Adjustment for multiple comparisons was also not considered.

**RESULTS**

**Sample characteristics**

The 2,020 patients included and followed-up in Spain account for 18.4% of the total sample of the SOHO study (10,972 patients). Of these, 27 were excluded from all analyses either because they could not be ascribed to any treatment cohort or did not meet the selection criteria. Most of the patients were receiving typical antipsychotics prior to enrolment, were male, received treatment in the public healthcare system, lived in an urban setting, and changed their treatment because of lack of efficacy. The olanzapine cohort accounted for 54.8% of the sample (1,092 patients). Among the remaining patients (901 patients), 51.8% (467 patients; 23.4% of the total sample) started treatment with risperidone and 20.3% (183 patients; 9.2% of the total)
with quetiapine. A total of 2.4% of the patients (22; 1.1% of the total) initiated polytherapy (two or more antipsychotics), this always being with typical antipsychotics. In Table 1 there are more details about the antipsychotic drugs started by the patients at the time of enrolment. Sociodemographic and clinical characteristics were generally homogeneous among study cohorts with the exception of the proportion of patients who initiated treatment for the first time, that were greater in the olanzapine and risperidone cohorts; and the body mass index, that was lower in the olanzapine cohort. A complete description of this sample has been reported in a previous article.\textsuperscript{15}

**Analysis of remission**

A total of 1,366 patients (68.5% of the total of the evaluable Spanish sample). Those patients in the other atypical antipsychotic cohort, those who did not complete the 3 year follow-up, those who had missed more than one visit or had incomplete data in CGI-SCH evaluations or hospitalization assessments were excluded. In all, 821 (60.1%) patients met the remission criteria. By cohorts (fig. 1), patients treated with olanzapine and clozapine showed the highest remission rates (65.5% and 60%, respectively), while the lowest ones were observed in those treated with typical depot or with polytherapy (28% and 45.5%, respectively). The proportion of females, of patients in the first treatment for schizophrenia, with stable partner, living independently, with paid employment, engaged in social activities, or starting treatment with olanzapine, risperidone, clozapine or oral typical antipsychotics were greater among patients who showed remission at some time than those who did not. Furthermore, the duration of the illness was shorter and the scores of the CGI-SCH scales lower in the group of patients that showed remission (Table 2 and fig. 1).

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Absolute and relative frequencies of the treatments initiated by patients recruited in the SOHO study in Spain</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of patients (%)</td>
<td></td>
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<tr>
<td>Patients in the olanzapine group</td>
<td>1,092 (54.1)</td>
</tr>
<tr>
<td>Patients in control group</td>
<td>901 (45.9)</td>
</tr>
<tr>
<td>Risperidone</td>
<td>467 (51.8)</td>
</tr>
<tr>
<td>Quetiapine</td>
<td>183 (20.3)</td>
</tr>
<tr>
<td>Any typical depot antipsychotic</td>
<td>98 (10.9)</td>
</tr>
<tr>
<td>Any typical oral antipsychotic</td>
<td>94 (10.4)</td>
</tr>
<tr>
<td>Clozapine</td>
<td>34 (3.8)</td>
</tr>
<tr>
<td>Two or more antipsychotics</td>
<td>22 (2.5)</td>
</tr>
<tr>
<td>Any other atypical antipsychotic</td>
<td>3 (0.3)</td>
</tr>
</tbody>
</table>

![Figure 1](image-url)
The adjusted analyses (fig. 2) revealed that the likelihood of remission was significantly lower in patients who started treatment with an oral typical antipsychotic drug or with two or more antipsychotics than those who began treatment with olanzapine (respective p-values: 0.0104 and <0.0001). Similarly, male patients and patients who were prescribed concomitant anxiolytics had less likelihood of remission (p-values: 0.0255 and 0.0060, respectively); while those who had participated in any social activity in the 4 weeks prior to their inclusion, had a paid employment, or who initiated treatment for their schizophrenia for the first time had greater likelihood of achieving remission during the follow-up (p-values: 0.0069, 0.0254 and 0.0069, respectively). Remission was also less likely between months 6 and 12, 12 and 18, or 18 and 24 than between months 30 and 36 (p-values: < 0.0001, < 0.0001 and 0.0030, respectively); as well as and for patients with higher (worse) cognitive, negative or positive CGI-SCH scores or greater BMI at the initial visit (p-values: < 0.0001, < 0.0001, 0.0036 and <0.0001, respectively).

The survival distribution functions of the time to remission by cohort (fig. 3 A) shows that it was shorter with olanzapine (solid line, below the remaining lines) than with other antipsychotics, followed by clozapine and oral typical antipsychotics. Patients initially prescribed multiple antipsychotic prescriptions had the longest time to remission.

Analysis of relapse

A total of 1,018 patients were included in this analysis. The patients were excluded from this analysis for the same reasons as described for remission. Outcomes at 3 years

### Table 2 Sociodemographical and clinical characteristics used in the analyses of remission, relapse and treatment maintenance. The values correspond to the baseline visit relevant to each analysis

<table>
<thead>
<tr>
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<th>Remission (n = 1,366)</th>
<th>Relapse (n = 1,018)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Sociodemographical characteristics</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gender: women (n [%])*</td>
<td>338 (41.22) 175 (32.11)</td>
<td>78 (41.05) 326 (39.42)</td>
</tr>
<tr>
<td>Age at initial visit in years [mean (SD)]</td>
<td>25.83 (4.41) 27.39 (5.31)</td>
<td>26.61 (4.41) 26.83 (4.54)</td>
</tr>
<tr>
<td></td>
<td></td>
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<tr>
<td><strong>Clinical variables</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Time from first contact in years [mean (SD)]</td>
<td>9.97 (10.78) 13.81 (12.33)</td>
<td>9.03 (10.58) 10.72 (11.17)</td>
</tr>
<tr>
<td>Age at first contact in years [mean (SD)]</td>
<td>27.74 (9.87) 26.59 (9.34)</td>
<td>26.74 (9.59) 27.74 (9.86)</td>
</tr>
<tr>
<td>In the first episode in the initial visit [n [%]]*</td>
<td>91 (11.08) 22 (4.04)</td>
<td>16 (8.42) 84 (10.14)</td>
</tr>
<tr>
<td>Patients without stable partner [n [%]]*</td>
<td>213 (26.13) 103 (19.04)</td>
<td>52 (27.66) 222 (26.94)</td>
</tr>
<tr>
<td>Independent patients [n [%]]*</td>
<td>364 (44.50) 187 (34.38)</td>
<td>83 (44.15) 367 (44.59)</td>
</tr>
<tr>
<td>Paid employment [n [%]]*</td>
<td>178 (21.68) 51 (9.36)</td>
<td>45 (23.68) 196 (23.82)</td>
</tr>
<tr>
<td>Participation in social activities [n [%]]*</td>
<td>617 (75.52) 330 (60.66)</td>
<td>158 (83.16) 711 (86.39)</td>
</tr>
<tr>
<td>Extrapyramidal symptoms [n [%]]*</td>
<td>281 (34.39) 250 (46.13)</td>
<td>35 (18.42) 116 (14.11)</td>
</tr>
<tr>
<td>History of alcohol abuse [n [%]]*</td>
<td>23 (2.80) 26 (4.77)</td>
<td>6 (3.17) 19 (2.29)</td>
</tr>
<tr>
<td>History of substance abuse [n [%]]*</td>
<td>31 (3.78) 18 (3.30)</td>
<td>12 (6.35) 22 (2.66)</td>
</tr>
<tr>
<td>Concomitant anticholinergics [n [%]]*</td>
<td>111 (13.52) 99 (18.17)</td>
<td>36 (18.95) 74 (8.94)</td>
</tr>
<tr>
<td>Concomitant mood stabilizers [n [%]]*</td>
<td>44 (5.36) 45 (8.26)</td>
<td></td>
</tr>
<tr>
<td>Decrease of libido observed [n [%]]*</td>
<td>319 (42.43) 237 (51.52)</td>
<td></td>
</tr>
<tr>
<td>History of suicide attempt [n [%]]*</td>
<td>51 (6.21) 26 (4.78)</td>
<td></td>
</tr>
<tr>
<td>Hostile behavior [n [%]]*</td>
<td></td>
<td>19 (10.00) 51 (6.18)</td>
</tr>
<tr>
<td>Global baseline score CGI-SCH [mean (SD)]</td>
<td>4.34 (0.93) 4.81 (0.87)</td>
<td>2.83 (0.43) 2.79 (0.49)</td>
</tr>
<tr>
<td>Positive symptoms score at baseline CGI-SCH [mean (SD)]</td>
<td>3.90 (1.32) 4.21 (1.22)</td>
<td></td>
</tr>
<tr>
<td>Negative symptoms score at baseline CGI-SCH [mean (SD)]</td>
<td>3.90 (1.13) 4.65 (1.07)</td>
<td></td>
</tr>
<tr>
<td>Depressive symptoms score at baseline CGI-SCH [mean (SD)]</td>
<td>3.53 (1.17) 4.33 (1.09)</td>
<td></td>
</tr>
<tr>
<td>Cognitive symptoms score at baseline CGI-SCH [mean (SD)]</td>
<td>3.80 (1.01) 4.21 (1.12)</td>
<td></td>
</tr>
<tr>
<td>EVA-EQ5D baseline score [mean (SD)]</td>
<td>49.33 (18.40) 51.80 (40.00)</td>
<td>71.26 (14.72) 67.89 (14.46)</td>
</tr>
</tbody>
</table>

*The relative frequencies have been calculated on the total of observations available in each variable, which, in some cases, was less than the total of the group (n)
Remission and relapse in the ambulatory treatment of patients with schizophrenia.
Outcomes at 3 years

Figure 2
A) Odds ratios and 95% confidence intervals of the independent variables used in the multivariate logistic regression model of the likelihood of showing remission throughout the three-year follow-up period. B) hazard rates and 95% confidence intervals of the variables considered in the Cox proportional hazards model of time to relapse;
reasons as for the analysis of remission, however, those patients who did not have an a score equal to or less than 3 at any time of follow-up were also not included since relapse can only be considered when there a previous favorable clinical state has existed. In all, 190 patients (18.7%) met the criteria for relapse. The sociodemographic and clinical characteristics at baseline (the first visit in which the patients had an overall CGI-SCH score equal or lower than 3) were similar between the groups of patients that relapsed and that did not relapse, with the exception of a greater proportion of substance abusers and concomitant need for anticholinergics among patients who relapsed (table 2).

The adjusted analysis revealed that patients who started treatment with quetiapine had a significantly greater risk of relapse (or, in other words, that relapse occurred in a significantly shorter period) than those who started treatment with olanzapine (p-value: 0.0047). Similarly, patients who received concomitant anticholinergics were at significantly greater risk of relapse than those who did not (p-value: 0.0010). A marginally significant association between a longer duration of the illness with a lower risk of relapse was also found (p-value: 0.0470) (fig. 2 B).

The longest times to relapse corresponded to patients in the clozapine and risperidone cohorts, followed by patients in the olanzapine cohort (the upper lines in figure 3 B); while the shortest times were found among patients receiving two or more antipsychotics (fig. 3 B).

On the other hand, the risk of being hospitalized (fig. 2 C) was greater in patients taking quetiapine than in those on olanzapine. A worse cognitive status at baseline, having a background of suicide attempts, concomitant prescription of mood-stabilizers or the occurrence of gynaecomastia and galactorrhoea were all associated with a greater risk of relapse. Conversely, the risk decreased with age and the duration of the disease; although the respective point estimates of the latter two factors were close to the unity (fig. 2 C).

DISCUSSION

The SOHO study is a large, three-year long European study on antipsychotic treatment in out-patients. Its observational design makes it possible to analyze the evolution of
the disorder in the routine clinical practice and its size has helped to used complex statistical techniques to study the evolution predictors. The clinical remission and relapse, analyzed in the present work, is one of the most relevant results for the patient's well-being.

The factors associated to a lower likelihood of remission were greater baseline clinical severity, worse social functioning, greater disease evolution time, and the use of a concomitant treatment with anxiolytics, also probably associated a greater clinical severity. These results reinforce previous studies that indicate the good prognosis associated to good social functioning and the tendency of schizophrenia to have lower likelihood of improvement as the disorder evolves. We have also observed that the likelihood of remission increases with time of treatment after a relapse and that the likelihood of relapse is also maintained even years after the remission. This fact reinforces the notion of the importance of continued treatment to improve the outcomes. The inverse association found between body mass index at baseline and the likelihood of remission was greater among patients who initiated treatment with this drug.

Consistent with results obtained in both randomized clinical trials and naturalistic studies, the clinical evolution, measured as likelihood of remission, has been better in patients who initiated olanzapine in the baseline visit compared to those who used typical antipsychotics. However, other reviews did not find the same results. It is likely that factors such as treatment compliance grade, its maintenance and profile of side effects in addition to the intrinsic antipsychotic efficacy of the drug, influence the long-term clinical efficacy of olanzapine versus typical antipsychotics. These aspects are not generally adequately assessed in the clinical trials. In this sense, the SOHO study shares some of these characteristics with the clinical trial on antipsychotic efficacy of several interventions (Clinical Antipsychotic Trial of Intervention Efficacy [CATIE]), in which greater improvement and longer time of discontinuation with olanzapine than with other atypical antipsychotics has been demonstrated.

The 18.7% incidence of relapse found in our analysis is much lower than what had been reported in previous studies. The finding may be because the appearance of relapses was only evaluated at discrete time points in the control...
visits made every 6 months, determining the possibility of compliance of relapse criteria between visits only in the cases in which there was a hospitalization. Furthermore, it is likely that the participating psychiatrists, because this was a three-year follow-up study, tended to include patients who complied with the study treatment. Among all the factors considered, only treatment with quetiapine compared to olanzapine and the need for concomitant treatment with anticholinergics showed sufficient predictive capacity to explain future relapses. The concomitant treatment with anticholinergic may be associated to the use of antipsychotics that cause side effects, which in turn are generally associated to worse therapeutic compliance26.

It is not surprising that patients with suicide attempts have a greater risk of being hospitalized. Because the risk was also greater in patients requiring concomitant mood stabilizers, affective symptoms appear as an important factor influencing in the need for admissions. The increased risk in patients with a worse cognitive status is also important. Taken together, these data suggest that once we have achieved an effective control of psychotic symptoms with continued effective therapy, the target to reduce in-hospital stay should be the cognitive and affective impairments of these patients.

The SOHO study has some general limitations that affect the analysis of remission and relapse. In the first place, the non-randomized allocation has been already mentioned. However, the use of multivariate statistical techniques that take into account the baseline differences among patient who initiate different treatments has been shown as valid to control this bias27. However, there is no assurance that all the factors that may influence the outcome have been considered, as the cohorts may be also imbalanced by other unmeasured parameters. Furthermore, the small number of patients in some subgroups (as is the case of the cohorts of patients with polytherapy or treated with clozapine) may compromise the robustness of estimations made by regression analysis with these treatments. The presence of information biases also cannot be ruled out, as participating psychiatrists were obviously not blinded to the treatment the patients received. Nevertheless, an attempt was made to reduce the influence of this bias in the design stage. This was done by procuring as much flexibility as possible in the patients’ treatment and thus avoiding interference with the routine clinical practice. In fact, a specific assessment made on such effects could not demonstrate the existence of observer biases in the SOHO study27. It is also possible that there was a tendency of the participating psychiatrists to keep patients in the study as long as possible by delivering some extra efforts because of economic incentives calculated on the basis of study visits reported. Consequently, the presence of a selection bias cannot be ruled out, as cases having worse prognosis were eliminated from the study in benefit with the less severe patients.

In conclusion, the data presented here are relevant because they come from a considerably large sample of almost 2000 patients with schizophrenia followed-up for as long as three years within Spain in the frame of an observational prospective clinical study. Patients who require change of antipsychotic treatment for clinical reasons may benefit from substantial improvements in terms of high remission and low relapse rates. Because of the long-term favorable outcomes, olanzapine has been shown to be of great value in routine clinical practice.

ACKNOWLEDGEMENTS

This study has been financed by Eli Lilly and Company Limited, S.A.

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