CLINICAL STUDY

Persistence at 1 year of oral antiosteoporotic drugs: a prospective study in a comprehensive health insurance database

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Abstract

Objective: Treatments against osteoporosis have demonstrated fracture risk reduction but persistence to therapy remains a major issue. Intermittent regimens have been developed to improve persistence. The aim of this 1-year prospective study was to compare, in the general population, the persistence of various oral regimens of antiosteoporotic treatment.

Methods: We conducted this prospective study in the French comprehensive public health insurance database of the Rhône-Alpes region. Women aged 45 years or older who had a first reimbursement of an oral antiosteoporotic treatment during February 2007 composed the study cohort. Persistence was defined by the proportion of patients refilling a prescription in the pharmacist delivery register (ERASME). Using statistical analyses like Kaplan–Meier survival curves and log-rank tests, we compared the treatment persistence of strontium ranelate, raloxifene, and daily-, weekly-, and monthly bisphosphonates.

Results: Two thousand four hundred and nineteen patients were included over a period of 1 month and followed up for 12 months. Two hundred and eighty-nine (11.9%) patients were treated with monthly bisphosphonates, 1298 (53.7%) with weekly bisphosphonates, and 832 (34.4%) with daily treatments (526 strontium ranelate (21.7%), 296 raloxifene (12.2%), and 10 bisphosphonates (0.4%)). At 1 year, overall persistence was 34%. Fifty percent of patients on monthly bisphosphonates were still persistent while only 37% of patients on weekly bisphosphonates, 34% on raloxifene, and 16% on strontium ranelate were persistent. Therapy monitoring with biochemical markers or bone mineral density was associated with improved persistence.

Conclusion: Overall persistence at 1 year was low, but intermittent regimens were associated with higher persistence rates, along with women who had therapy monitoring.

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Introduction

Several classes of oral drugs can be used in the treatment of postmenopausal osteoporosis. They have all shown efficacy to reduce fracture risk. Adherence to the treatment, however, remains a critical issue in the management of chronic diseases such as osteoporosis. One reason is that patients do not perceive the need for treatment because osteoporosis is an asymptomatic disease before the first fracture (1, 2). Therapeutic adherence is the extent to which a person takes medication under the physician’s recommendation. Adherence includes persistence, length of time the patient continues taking medication from initiation, and compliance, which describes how regimen and doses are taken. Compliance is commonly evaluated by the medication possession rate (MPR) corresponding to the number of pills correctly taken divided by the number of pills prescribed. A good compliance is generally above 80%.

In osteoporosis, therapeutic adherence is logically associated with fracture risk reduction (3). Independent correlation of persistence with fracture risk reduction was also observed in two US claim databases including 35 537 women receiving bisphosphonates. Only 43% of patients were refill compliant and 20% persisted after 2 years of follow-up (4). In the PHARMO-RLS study, adherence to bisphosphonate therapy was evaluated through compliance (MPR) in new postmenopausal female bisphosphonate users. During the follow-up, a steady increase in noncompliant patients (MPR <80%) was observed: 42% after 1 year, 51% after 2 years, and 60% after 3 years. Noncompliant bisphosphonate use was associated with a 40% increased risk of osteoporotic
fracture requiring hospitalization (5). In another observational study of 11,252 postmenopausal women extracted from a drug dispensation database, an antosteoporotic medication covering at least 80% of the time was independently associated with a fracture risk reduction of 16% (6). In the UK, the General Practice Research Database (GPRD) 1-year persistence rate for risedronate or alendronate was 58% but decreased to 24% after 5 years. Persistence of at least 24 months showed a hip fracture risk reduction of 34%, and hip fracture hazard rates diverged after 1 year of treatment (7). The study conducted among the new postmenopausal female alendronate users of the Belgian national social security database showed a low global persistence (40%) at 12 months, but persistent patients had a 60% hip fracture relative risk reduction. MPR was also a significant predictor of hip fracture with an increased adjusted risk of 0.4% for each 1% decrease in the MPR. MPR was significantly higher for patients who received weekly alendronate (70.5%) compared with daily regimen (58.6%), raising the issue of optimal regimen (8). Similarly, in a US claim database, a 42% persistence rate was observed with alendronate after 24 months of follow-up. Patients with weekly regimen were more likely to be persistent (49%) than nonpersistent (38%) (9).

All these studies, however, have not been conducted in the general population. Some of them used private insurance claim databases and some used regrouping of general practitioners (GPs). In addition, all these studies were retrospective, so potential predictors of adherence could not be ascertained before starting therapy. Thus, we have conducted a prospective study to assess the persistence of several antosteoporotic drugs and whether monthly regimen was associated with a better persistence in a universal coverage health care system database providing data of the general population.

**Materials and methods**

**Database**

The Rhône-Alpes area has six million inhabitants representing ~10% of the French population. We have prospectively used the Rhône-Alpes ERASME database (Extraction, Recherche et Analyse pour un Suivi Médico-Économique), which means Extraction, Research and Analysis of a Medico-Economical follow-up. This database is run by the French public social insurance (Sécurité Sociale). The ERASME database contains fee-for-service claims for in- and outpatient medical services supplied to 80% of the residents of France, as well as all drugs and tests that are ordered by physicians for community-dwelling patients. The remaining 20% of the French residents are covered by special subdivisions of the French social health care system, depending on their job. Before being nationally centralized, claims are first centralized at the regional level by each regional center of the CNAMTS (Caisse Nationale d’Assurance Maladie des Travailleurs Salariés). The organization of data collection and the database has been previously described in detail (10).

We created an inception cohort of all women aged 45 years or older living in the Rhône-Alpes area with a first reimbursement of an oral antosteoporotic treatment (i.e. raloxifene, strontium ranelate, and daily-, weekly-, and monthly brand and generic bisphosphonates) between 1st and 28th February 2007. Patients treated with teriparatide or intravenous bisphosphonate were excluded. Patients previously treated with any antosteoporotic drugs including teriparatide or intravenous bisphosphonates were also excluded.

**Inclusion data**

We used the date of first reimbursement as the date of entry, and we collected for each woman the type of antosteoporotic treatment with raloxifene, strontium ranelate, daily-, weekly-, and monthly brand and generic bisphosphonates; age; presence of severe chronic comorbidity; and prior bone mineral density (BMD) reimbursement. For the definition of chronic severe comorbidity, we used the ALD (Affection de Longue Durée or Chronic Severe Comorbidity (SCC)) status as a proxy. The ALD status is given to patients who have chronic diseases, e.g. diabetes, cancer, cardiac failure, renal deficiency, rheumatoid arthritis, and multiple sclerosis. It is considered that these diseases imply costly and long-term treatments and monitoring so that the patients have no co-payment.

**Follow-up**

Each woman in the cohort was then followed for 1 year for reimbursement of oral antosteoporotic treatment, including the number of delivered pills, through record linkage in the Rhone-Alpes health insurance database. Using the number of delivered pills and the treatment regimen (monthly, weekly, and daily), assuming that all delivered pills were effectively taken by the patients, we computed the time to end of refilling in days. For example, a patient delivered 12 pills of a weekly bisphosphonate has a time to end of refilling of 84 days. In the survival analysis, we used a persistence of 365 days, corresponding to a 100% refilling treatment after 1 year of follow-up. The outcome was time to end of refilling. No refill gap was allowed. Reimbursement of serum type I collagen breakdown product (CTX) assessment 3–6 months after first reimbursement of a drug was also collected during the follow-up to know whether it could be associated with improved persistence.

**Statistical analysis**

Categorical variables were presented in n (%) and continuous variables in mean (s.d.) or median
(interquartile range (IQR)) as appropriate. Survival time was defined as the time between study entry (i.e. date of first reimbursement of oral antosteoporotic treatment between 1st and 28th February 2007) and either date of end of refilling before 28th February 2008 or 28th February 2008 as date of the end of the follow-up, which serves as the censoring time point. Kaplan–Meier curves were used to visualize persistence probabilities, with step down when a treatment has been stopped. Kaplan–Meier curves stratified on oral antosteoporotic treatment were plotted. Comparison of persistence between oral antosteoporotic treatments was done using a log-rank test. For the estimation of hazard ratios of antosteoporotic treatment, a multivariate Cox proportional hazard model was fitted including potential confounders, i.e. quartiles of age, severe chronic comorbidity, prior BMD, and serum CTX assessment. We also compared duration of treatment among the various regimens using uni- and multivariate linear regression models. All comparisons were two sided and a \( P \) value of \( <0.05 \) was used to indicate a statistically significant difference. Analyses were performed using STATA 9.1 Software version (StataCorp, College Station, TX, USA) and SPSS 15.0 Software version (SPSS, Inc., Chicago, IL, USA).

**Results**

**Baseline characteristics**

Two thousand four hundred and nineteen women aged 45 years or older were included over a period of 1 month in February 2007 when they were given for the first time an oral antosteoporotic treatment. The mean (± s.d.) age was 67.7 years (± 10.5) and 36.9% (\( n = 892 \)) of the sample had an associated severe chronic comorbidity. Interestingly, 897 (37.1%) of the population underwent a BMD testing in the 6 months preceding the initial drug delivery and 172 (7.1%) had a serum CTX assessment after prescription. Among patients with BMD testing, 35.2% did not have any CTX assessment, and among patients with CTX assessment, 38.4% did not have any BMD testing. Thus, patients with BMD testing and patients with serum CTX assessment cannot be considered to be the same population.

Among the 2419 women, 1298 (53.7%) received a weekly bisphosphonate, 526 (21.7%) strontium ranelate, 296 (12.2%) raloxifene, and 289 (11.9%) monthly bisphosphonates. Only some of the patients received a daily bisphosphonate (\( n = 10 \); 0.4%). As expected, patients receiving raloxifene were significantly younger (60.2 ± 7.8 years) with less severe chronic comorbidity (17.2%) in comparison with the other groups (Table 1).

The median duration of therapy was 168 (IQR: 56–336) days with strontium ranelate and 224 (IQR: 184–365) days with raloxifene. For bisphosphonates, the median significantly increased with the dosing frequency: 84 (IQR: 28–140) days for daily, 308 (IQR: 112–365) for weekly, and 365 (IQR: 180–365) for the monthly regimen (Table 2).

### Table 1

<table>
<thead>
<tr>
<th>Total</th>
<th>Raloxifene</th>
<th>SR</th>
<th>Daily BP</th>
<th>Weekly BP</th>
<th>Monthly BP</th>
<th>( P^a )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( n) (%)</td>
<td>2419 (100)</td>
<td>296 (12.2)</td>
<td>526 (21.7)</td>
<td>10 (0.4)</td>
<td>1298 (53.7)</td>
<td>289 (11.9)</td>
</tr>
<tr>
<td>Age (years)</td>
<td>67.7 (10.5)</td>
<td>60.2 (7.8)</td>
<td>68.6 (11.1)</td>
<td>65.3 (12.5)</td>
<td>69.0 (10.4)</td>
<td>68.0 (9.6)</td>
</tr>
<tr>
<td>SCC</td>
<td>892 (36.9)</td>
<td>51 (17.2)</td>
<td>186 (35.4)</td>
<td>3 (30.0)</td>
<td>541 (41.7)</td>
<td>111 (38.4)</td>
</tr>
<tr>
<td>Prior BMD testing</td>
<td>897 (37.1)</td>
<td>126 (42.6)</td>
<td>193 (36.7)</td>
<td>2 (20.0)</td>
<td>445 (34.3)</td>
<td>131 (45.3)</td>
</tr>
<tr>
<td>CTX testing</td>
<td>172 (7.1)</td>
<td>29 (9.8)</td>
<td>44 (8.4)</td>
<td>0 (0)</td>
<td>77 (5.9)</td>
<td>22 (7.6)</td>
</tr>
</tbody>
</table>

\( ^a \)ANOVA for continuous variables; Fisher exact test for categorical variables.

SCC, severe chronic comorbidity; SR, strontium ranelate

**One-year persistence and factors associated with persistence**

The overall persistence was 34% (\( n = 818/2419 \)) and varied significantly according to the drug and the regimen (log-rank test \( P < 0.001 \); Fig. 1). Daily treatments had the smallest persistence rates ranging from 10% on daily bisphosphonates to 16% on strontium ranelate and 34% on raloxifene. Monthly bisphosphonates (50%) had a better persistence rate than the weekly bisphosphonates (37%).

In a multivariate analysis (Cox proportional hazard model) including age, severe chronic comorbidity, prior BMD testing, and serum CTX assessment after prescription as covariates, treatment regimen remained independently associated with persistence. Compared with weekly bisphosphonates taken as the reference category, monthly bisphosphonates (hazard ratio (HR) = 1.32 (1.10; 1.57), \( P = 0.002 \)) were independently associated with improved persistence. The three daily treatments were independently associated with lower persistence (Fig. 2A). The same results were observed for a persistence defined as 80% (292 days) of treatment refilling at 1 year (data not shown).

In univariate analysis, patients who underwent CTX measurement (log-rank \( P = 0.009 \); Fig. 3A) or BMD...
Table 2 Comparison of the duration of various regimens of oral antiosteoporotic treatment. Coefficients express the mean duration that patients persisted on therapy in comparison with the reference group (weekly bisphosphonates). They represent the number of additional days on therapy (monthly bisphosphonates) or the number of fewer days on therapy (raloxifene, strontium ranelate, and daily bisphosphonates), compared with weekly regimens. Data are expressed in days.

<table>
<thead>
<tr>
<th>Regimen</th>
<th>Median (Q1–Q3)</th>
<th>Coefficient (95% CI)a</th>
<th>P</th>
<th>Coefficient (95% CI)b</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Raloxifene</td>
<td>224 (184–365)</td>
<td>−25.7 (−41.9– −9.6)</td>
<td>0.002</td>
<td>−31.5 (−48.2– 14.8)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>SR</td>
<td>168 (56–336)</td>
<td>−57.8 (−70.7– −44.8)</td>
<td>&lt;0.01</td>
<td>−59.6 (−72.5– −46.7)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Daily BP</td>
<td>84 (28–140)</td>
<td>−132.5 (−212.0– −52.9)</td>
<td>0.001</td>
<td>−128.9 (−208.0– −49.8)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Weekly BP</td>
<td>308 (112–365)</td>
<td>Reference</td>
<td>−</td>
<td>Reference</td>
<td>−</td>
</tr>
<tr>
<td>Monthly BP</td>
<td>365 (180–365)</td>
<td>24.0 (7.7 – 40.3)</td>
<td>0.004</td>
<td>20.4 (4.2 – 36.7)</td>
<td>0.014</td>
</tr>
</tbody>
</table>

Q1, first quartile; Q3, third quartile; CI, confidence interval; BP, bisphosphonate; SR, strontium ranelate.

aUnivariate linear regression.

bMultivariate linear regression adjusted for age, severe chronic comorbidity, prior BMD testing, and CTX assessment.

evaluation before treatment initiation (log-rank P<0.001; Fig. 3B) had a significantly improved persistence. In our multivariate model including the previous covariates (i.e. taking BMD into account), CTX measurement remained significantly associated with persistence with an HR of 1.19 (1.04; 1.37, P < 0.05), indicating that CTX, in this population, was an independent predictor of adherence. Prior BMD testing was also an independent predictor of persistence (HR = 1.31 (1.18; 1.46), P < 0.001) in multivariate analysis after adjusting for CTX assessment (Fig. 2B).

The second quartile of age (60–68 years; HR = 1.19 (1.04; 1.37), P = 0.013) was another factor independently associated with improved persistence. No association was found with severe and/or chronic comorbidity (ALD).

In a linear regression analysis we compared duration of refilling for the different regimens. In multivariate analysis adjusted for age, severe chronic comorbidity, prior BMD testing, and serum CTX assessment, duration of treatment with the monthly regimen was on average 20.4 days longer than weekly regimens (95% confidence interval (4.2–36.7), P = 0.014). The three daily treatment durations were on average (mean) between 31.5 and 128.9 days shorter than weekly treatments (all P < 0.01; Table 2).

Discussion

In this prospective study within a comprehensive health insurance database we found that overall persistence at 1 year of all the available oral antiosteoporotic treatments was low. Persistence tended to improve with the longest interval intermittent regimen. Data regarding strontium ranelate have to be interpreted with caution because, during the study period, a European Medicines Agency warning for Drug Reaction with Eosinophilia and Systemic Symptoms (DRESS) was released in November 2007. DRESS refers to a rare adverse reaction occurring usually in the first month of exposure and associating a rash with fever and inflammation of internal organs. If patients continue with the therapy it can lead to death (11, 12, 13, 14). The onset of DRESS, shortly after strontium ranelate marketing, received high media coverage that may have temporarily increased the propensity of patients (or physicians) to stop the drug. This may have also affected treatment initiation but that does not influence our study. Nevertheless, in a recent retrospective study, strontium ranelate persistence at 1 year was also found to be very low at 21.9% (15).

We observed that persistence improved when BMD testing was carried out done before the initial prescription and when bone markers were measured after 3–6 months of therapy. Observational data have already suggested that BMD testing could increase drug prescription (16) and improve persistence (17). Moreover, patients who thought that their bone density test results did not show osteoporosis are more likely to discontinue the therapy early (2). Results with markers

Figure 1 Kaplan–Meier survival curves with time to end of refilling (days) according to the use of initial antiosteoeporotic treatment for 1 year: raloxifene, strontium ranelate, and daily-, weekly-, and monthly bisphosphonates (BPs).
are supported by previous data showing that serum CTX assessment after 3–6 months might improve patients’ persistence on therapy (18). Monitored patients had a greater adherence than the controlled arm. However, the nature of monitoring did not seem to matter because no difference was observed between the nurse-monitoring and the marker-monitoring groups (18). This study was limited by the small sample size of each group. The IMPACT study analyzed the effect on persistence of monitoring bone turnover markers (uNTX) in postmenopausal women treated with risidoثنate. A total of 2302 women were randomized between standard care and reinforcement with the marker result. IMPACT reported a surprisingly high 1-year persistence rate in both the groups with 77% in the standard care group and 80% in the reinforcement group (19). Nevertheless, the positive feedback message to the patient was associated with improved persistence. We found that intermittent regimens provided better persistence results at 1 year, confirming previous studies comparing daily with weekly regimens (8, 20). Recently, in a different French prescription database within a network of 1200 GPs around the country (Thales), Côté et al. (21) have reported a clear superiority of 1-year persistence rate with monthly bisphosphonate (47%) in comparison with weekly bisphosphonates (30%). One limitation of the Thales database is that it does not represent all prescribers and all patients. Indeed, it covers a set of GPs who are paid to participate in the network and may have greater implications for patients. Another potential bias is the absence of possibility for identifying patients who switch to out-of-network physicians. In contrast, our study cohort has the advantage of being representative of the general population. Even when patients switched to another physician, their reimbursements are still managed by the public health insurance. Interestingly, Thales network also showed that densitometry performed before treatment initiation improved adherence. The benefit of monthly regimen is sustained by the results from the PERSIST study, which randomized patients into weekly vs monthly bisphosphonate groups. Persistence in the monthly bisphosphonate group associated with a patient support program (PSP) was superior (57 vs 39%). Nevertheless, the follow-up was short (6 months) and there was no monthly arm without PSP (22). Thus, it remained difficult to interpret

![Graph](image-url)
the relative contributions of the PSP and the dosing regimen. In another study on US retail pharmacy prescription refills, no superiority with monthly regimen was found. However, the study was performed soon after the initial release of ibandronate and results are biased by the refilling gap of 30 days that disadvantaged the monthly regimen and by the co-payment variation in the US healthcare system, with many patients initiating monthly therapy for free with a voucher and not refilling the first box. There was also no information about free samples directly distributed by GPs (23).

Several other determinants of therapeutic adherence have been reported, such as hospitalization for osteoporosis (24), an initial prescription by a rheumatologist (24), patient motivation (25, 26), and the onset of bothersome side effects (2). Among the side effects, gastrointestinal problems before or during bisphosphonate therapy have been associated with a decreased persistence of about 20% at 12 months (17) and reduced compliance (24).

Our study had strengths and weaknesses. It was conducted in a comprehensive health insurance database, so it described a representative sample of the general French population, with almost no loss to follow-up. Its prospective nature allowed for collection of potential predictors associated with patient adherence. However, it was based on refilling claims so we cannot assert that those drugs that were bought were really used appropriately. The main limitation is the observational nature of the database, which does not provide any reason for interruption or any information on patients, i.e. whether they definitively stopped their treatment or switched to another antosteoporotic treatment. We also have no indication on prescriber specialty or patients’ profiles, such as the severity of osteoporosis, nature of severe comorbidity, association with steroid therapy, vitamin D status, and calcium and vitamin D supplements. Indeed, it is conceivable for instance that patients with more severe osteoporosis tend to have a higher persistence and that patients’ profiles may have influenced the medical decision to prescribe the anti-osteoporotic treatment. Our study was performed before atypical fractures with long-term bisphosphonate use were reported. It remains unknown whether this recent description has affected the 1-year persistence rate (27, 28).

In conclusion, persistence of antosteoporotic drugs at 1 year in the French general population remained low, even with intermittent regimes which, however, tended to improve the rate of persistence. Supporting the initial prescription with BMD evaluation and early bone marker evaluation also tended to increase persistence.

Declaration of interest

F Canouët-Poirine, A-M Schott, V Ambrosi, and V Tainturier have nothing to declare. C B Confavreux received lecture fees from Amgen, Ar笛, Lilly, and MSD. R D Chapurlat received lecture fees from Amgen, Lilly, Merck, Novartis, Servier, and Warner-Chilcott.

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