Evolution of Drug Survival with Biological Agents and Apremilast Between 2012 and 2018 in Patients with Psoriasis from the PsoBioTeq Cohort

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Drug survival reflects treatment effectiveness and safety in real life. There is limited data on the variation of drug survival with the availability of systemic treatments with additional biological disease-modifying antirheumatic drugs (bDMARDs) or synthetic disease-modifying antirheumatic drugs (sDMARDs). The aim of this study was to determine whether the increasing number of available systemic treatments for psoriasis affects drug survival over time. Patients were selected from the PsoBioTeq cohort, a French prospective observational cohort enrolling patients with moderate to severe psoriasis. All patients initiating a first bDMARD or sDMARD were included. The primary outcome was comparison of drug survival over time. A multivariate Cox proportional hazard ratio model was computed. A total of 1,866 patients were included; 739 females (39%), median age 47 years. In the multivariate Cox model, no association was found between the calendar year of initiation and drug survival (hazard ratio) overlapping from 0.80 (0.42–1.52) to 1.17 (0.64–2.17), p = 0.633. In conclusion, drug survival in psoriasis is not affected by the year of initiation.

Key words: psoriasis; drug survival; treatment; biologic; apremilast.

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Psoriasis is a chronic inflammatory skin disease with a prevalence ranging from 0.9% to 8.5% of the population in Europe (1, 2), and an incidence ranging from 78.9/100,000 to 230/100,000 (1, 2). The last 2 decades have witnessed the approval of numerous biological agents, i.e. tumour necrosis factor (TNF) inhibitors, interleukin (IL)-12–23 inhibitors, IL-17 inhibitors, IL-23 inhibitor and/or apremilast, creating a wide therapeutic armamentarium (3). Most randomized clinical trials (4) include short-term evaluation of efficacy and safety (4). In a real-life setting, due to a loss of effectiveness, or to adverse events, some patients may discontinue or switch treatment, potentially leading to diseases flares and/or a reduced quality of life. To further evaluate and compare treatments, drug survival through time, also called “persistence” is frequently used (5–7). Drug survival is a surrogate criterion representing a good balance between effectiveness and safety in a real-life setting (5).

Several studies have evaluated drug survival in patients with moderate to severe psoriasis. Variations in drug survival were observed between cohorts with up to 79% retention rates at 1 year from 2008 to 2013 in the Spanish BIOBADERM cohort (8), 77% from 2007 to 2014 in the British BADBIR cohort (6), and only 62% according to the French healthcare system between 2009 and 2016 (9). An Israeli cohort reported that the calendar year of initiation could be associated with a significant variation

SIGNIFICANCE

Drug survival is thought to represent a balance between effectiveness and safety in real life. Nevertheless, there is limited data on the variation of drug survival with the availability of additional biological or synthetic systemic treatments. This study found that drug survival in psoriasis is not affected by the year of initiation, suggesting that drug survival of patients with moderate to severe psoriasis is not sensitive to the increasing therapeutic armamentarium, and therefore strengthens its use to estimate balance between treatment effectiveness and adverse reactions over years.
in drug survival over time, patients initiating a systemic treatment more recently having shorter drug survival (10). In France, drug approval was granted in 2005 for etanercept and infliximab, 2008 for adalimumab, 2010 for ustekinumab, 2016 for ixekizumab, secukinumab, and apremilast. This widening of the armamentarium of psoriasis treatments may have promoted a reduction in drug survival, by giving patients and physicians additional options in case of suboptimal effectiveness and safety. Hence, it was hypothesized that the increasing number of available treatments would shorten the drug survival over the years. The objective of this study was to assess the evolution of drug survival according to the calendar year of initiation in patients with moderate to severe psoriasis.

PATIENTS AND METHODS

Overview and data source

The PsoBioTeq cohort is a French prospective observational cohort that enrols and follows prospectively adult patients (aged >18 years) with moderate to severe psoriasis. The objectives of PsoBioTeq are to describe the use, benefit and risks of systemic treatments with conventional, synthetic disease-modifying antirheumatic drugs (sDMARDs) and biological disease-modifying antirheumatic drugs (bDMARDs) in a real-life setting. The choice of treatment is left to the clinicians and is not influenced by participation in the cohort. PsoBioTeq is currently recruiting patients from 41 departments of dermatology across France. Several ancillary studies have already been published with the PsoBioTeq study (11, 12). The study protocol was approved by the “Comité d’Évaluation de l’Ethique des projets de Recherche Biomédicale (CEERB) du GHU Nord” (authorization number JMD/MDM/177-11). The ClinicalTrials.gov identifier is NCT01617018. Written informed consent was obtained from all patients before study inclusion.

Population and data collection

All patients initiating a first therapeutic line of biological disease-modifying antirheumatic drugs (bDMARDs) or a synthetic disease-modifying antirheumatic drugs (sDMARD) between 1 July 2012 and 31 December 2018 were included in the analysis. Patients treated only with methotrexate (MTX), acitretin and cyclosporine were not included. Also excluded were patients treated with brodalumab, certolizumab pegol and guselkumab, given their late French approval, occurring after 2018. Patients with only 1 inclusion visit were also excluded. The follow-up was conducted until 31 December 2019.

Demographic variables, including age and sex, were assessed. The study also assessed previous treatments including previous cyclosporine use, previous MTX use and previous phototherapy, concomitant MTX treatment at baseline, body mass index (BMI), main comorbidities (including chronic human immunodeficiency virus (HIV), hepatitis B virus (HBV), hepatitis C virus (HCV), infections, psoriatic arthritis (PsA), intestinal bowel disease (IBD), uveitis, diabetes, history of major adverse cardiovascular event (MACE), history of diabetes, dyslipidaemia, chronic obstructive pulmonary disease (COPD), chronic renal failure and positive tuberculous screening), and baseline disease severity according to Psoriasis Area and Severity Index (PASI).

The primary outcome was the rate of drug survival at 1 year, defined by evidence of a switch of treatment, or a discontinuation of more than 180 days. The covariate of interest was the calendar year of initiation. All patients were censored at 1 year if they had not stopped the treatment. The study also evaluated factors associated with drug survival as exploratory outcomes.

Statistical analysis

Descriptive statistics. Quantitative variables are presented as median (interquartile range; IQR) or mean ± standard deviation (SD), and categorical variables are presented as absolute values (percentage). The Kaplan–Meier method was used to display the rate of drug survival at 1 year with its 95% confidence interval (95% CI). To avoid informative censorship over time and biased estimators due to discrepancies in the follow-up between the different years of initiation, the study analysed only 1-year drug survival.

Inference. To determine whether the calendar year of initiation would impact the rate of drug survival at 1 year, 2 different models were used (see below). All tests were 2-tailed and p-value < 0.05 were considered significant. All statistics were performed with R software (R CRAN 3.6.2 (R foundation for statistical computing, Vienna, Austria).

Survival model. Using Kaplan–Meier’s method, the crude drug survival rates were calculated for all the treatments during the all-study period (from 2012 to 2018), then per year of initiation. Hazard ratios (HRs) and their 95% confidence interval (95% CI) were then computed using a Cox proportional HR model. The multivariate analysis included the calendar year of initiation and the variables with a known effect on drug survival in the medical literature (6–8), including treatments, previous treatment history, age, sex, body mass index, concomitant MTX, the presence of comorbidities, and PASI at baseline. It was decided not to impute missing data. Interactions were tested. The assumption of proportionality was assessed by visualizing the Schoenfeld residuals and the assumption of log-linearity by visualizing the Martingale residuals.

Fig. 1. Flow chart of the included population.
Subgroup analyses were conducted considering each drug separately (9). For IL-17 inhibitors and apremilast, the study period was from 2016 (date of the first commercialization in France) to 2018. Multivariate analyses could not be performed with IL-17 inhibitors due to small sample sizes.

**Time series analysis.** Secondly, a linear model was used to determine whether the calendar semesters would impact the rate of drug survival at 1 year. Compared with other former treatments, 2 highly effective bDMARDs treatments were marketed in 2016 (ixekizumab and secukinumab) (4). Thus, 2016 was considered as an intervention date, with a pre-period (from 2012 to 2015) and post-period (from 2016 to 2018). The dependent variable was the drug survival rate at 1 year computed from the Kaplan–Meier representation. The independent variable was the calendar semester of initiation. In case of no change in baseline level nor trend between the 2 periods, a more parsimonious model was computed, i.e. a simple linear regression with estimation of the time effect (through the calendar semester). Such analyses were not performed for IL-17 inhibitors and apremilast related to the date of the first commercialization (2016).

**RESULTS**

**Population**

A total of 1,866 patients were included (Fig. 1); 739 females (39%) with a median age of 47 years (IQR 36–57 years). Their characteristics are presented in Table I. PsA was found in 343 (19%) patients, and the overall median PASI at baseline was 9.6 (4.5–15.6). Regarding first-line therapy, 1,106 patients (59%) were initially treated with TNF inhibitors, including 665 receiving adalimumab, 389 etanercept and 52 infliximab. A total of 518 patients (28%) received ustekinumab, 80 patients (4%) received a IL-17 inhibitor, including 44 treated with secukinumab and 36 with ixekizumab. A total of 162 patients (9%) received apremilast. The numbers of patients initiating a first biologic per year and per drug are available in the Table S1. A total of 579 patients discontinued their treatment within the first year, indicating a global drug survival rate of 0.67 (95% CI 0.65–0.69) according to the Kaplan–Meier representation. The drug survival of the different treatments is shown in Fig. S1. During the study period (2012 to 2018), the crude drug survival rates at 1 year were 0.65 (95% CI 0.62–0.68) for TNF inhibitors, including 0.60 (0.55–0.65) for etanercept, 0.56 (0.44–0.72) for infliximab and 0.69 (0.65–0.72) for adalimumab. The crude drug survival rate at 1 year were 0.84 (95% CI 0.80–0.87), 0.77 (95% CI 0.70–0.86) for ustekinumab and IL17 inhibitors respectively, but only 0.23 (0.17–0.31) for apremilast.

**Inference**

Regarding the primary outcome, the Kaplan–Meier representation of drug survival according to the calendar year of initiation is shown in Fig. 2, indicating no difference according to the year of initiation. The calendar year of initiation was not associated with a reduction in 1-year drug survival over time in multivariate analysis, with HR overlapping from 0.80 (0.42–1.52) to 1.17 (0.64–2.17), and a global p-value of 0.63 (Table II).

The subgroup analyses considering biological treatments 1 by 1 showed results consistent with the primary analysis (Fig. 2 and Table SII), with HR overlapping from 0.52 (0.22–1.25) to 0.67 (0.28–1.59) for adalimumab, 0.37 (0.12–1.16) to 0.59 (0.10–1.74) for ustekinumab and 0.68 (0.19–2.48) to 1.71 (0.82–3.57) for etanercept. The yearly causes of discontinuation are shown in Table SIII. Discontinuation because of ineffectiveness appeared

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**Table I. Characteristics of the included population**

|                           | Overall (n = 1,866) | TNF-inhibitors (n = 41,106) | Ustekinumab (n = 518) | IL17-inhibitors (n = 80) | Apremilast (n = 162) |
|---------------------------|---------------------|-----------------------------|-----------------------|-------------------------|---------------------|
| Females, n (%)            | 739 (39)            | 433 (39)                    | 200 (39)              | 38 (47)                 | 70 (43)             |
| Age, years, median (IQR)  | 47 (36–57)          | 48 (36–58)                  | 44.0 (33.3–54.7)      | 43.8 (34.2–55.0)        | 50.5 (41.3–60.7)    |
| BMI, kg/m², median (IQR)  | 26.7 (23.4–30.8)    | 27.0 (23.4–31.1)            | 26.0 (23.2–30.1)      | 26.8 (24.2–30.8)        | 26.7 (23.9–30.5)    |
| PASI at baseline (median, IQR) | 9.5 (4.5–15.6)      | 10.5 (5.2–16.1)             | 8.9 (3.7–15.0)        | 9.0 (4.0–19.3)          | 7.4 (3.2–13.2)      |
| Psoriasis arthritis, n (%)| 343 (19)            | 259 (23)                    | 56 (11)               | 8 (10)                  | 24 (13)             |
| Intestinal bowel disease, n (%) | 7 (0.5)           | 5 (0.5)                     | 2 (0.4)               | 0 (0)                   | 0 (0)               |
| Uveitis, n (%)            | 3 (0.1)             | 2 (0.0)                     | 1 (0.2)               | 0 (0)                   | 0 (0)               |
| HIV, n (%)                | 7 (0.4)             | 3 (0)                       | 4 (1)                 | 0 (0)                   | 0 (0)               |
| Hepatitis B virus, n (%)  | 20 (1)              | 13 (1)                      | 5 (1)                 | 0 (0)                   | 2 (1)               |
| Hepatitis C virus, n (%)  | 19 (1)              | 12 (1)                      | 4 (1)                 | 0 (0)                   | 3 (2)               |
| Diabetes, n (%)           | 207 (11)            | 136 (12)                    | 46 (9)                | 6 (7)                   | 19 (11)             |
| History of MACE, n (%)    | 61 (3)              | 47 (4)                      | 6 (1)                 | 2 (1)                   | 6 (4)               |
| Dyslipidaemia, n (%)      | 282 (15)            | 186 (17)                    | 62 (12)               | 10 (12)                 | 24 (15)             |
| COPD, n (%)               | 66 (4)              | 41 (4)                      | 14 (3)                | 1 (1)                   | 10 (6)              |
| Cirrhosis, n (%)          | 35 (2)              | 28 (3)                      | 2 (0.4)               | 0 (0)                   | 5 (3)               |
| Chronic renal failure, n (%) | 30 (2)             | 23 (2)                      | 5 (1)                 | 1 (1)                   | 1 (1)               |
| History of tuberculosis, n (%) | 4 (0.2)         | 1 (0)                       | 2 (0.4)               | 1 (1)                   | 0 (0)               |
| Positive tuberculosis screening (IFN gamma release assay, n (%) | 112 (6)          | 68 (6)                      | 33 (6)                | 4 (4)                   | 7 (4)               |
| Concomitant MTX at baseline, n (%) | 221 (12)      | 148 (13)                    | 43 (8)                | 5 (6)                   | 28 (17)             |
| Previous MTX use, n (%)   | 1,113 (59)          | 718 (65)                    | 288 (56)              | 35 (44)                 | 72 (44)             |
| Previous cyclosporine use, n (%) | 251 (13)      | 154 (14)                    | 83 (16)               | 5 (4)                   | 9 (6)               |
| Previous PUVA therapy, n (%) | 1245 (67)       | 752 (68)                    | 354 (68)              | 52 (65)                 | 87 (54)             |

IQR: interquartile range; TNF: tumour necrosis factor; IL: interleukin; BMI: body mass index; PASI: Psoriasis Area and Severity Index; MACE: major adverse cardiovascular event; COPD: chronic obstructive pulmonary disease; IFN: interferon; MTX: methotrexate; PUVA: psoralen plus ultraviolet A.
to decrease slightly over time (from 77% in 2012 to 51% in 2018).

The segmented linear regression is represented in Fig. 3 and Table III. Regarding all treatment, there was no significant effect of 2016 on the change in mean drug survival rate at 1 year (−0.047 ± 0.063 ‰, p = 0.47), or on the time trend (−0.0004 ± 0.017 ‰ per semester, p = 0.98). A simple linear regression was thus computed, which showed similar results, with a global stability and a trend of −0.006 ± 0.004 ‰ per semester, p = 0.13. Subgroup analyses considering ustekinumab, etanercept and adalimumab showed consistent results, with no significant effect of 2016 and a stability over years, except for etanercept with a higher survival drug after 2016 (Fig. 3 and Tables SII–SIV).

Factors associated with drug survival during the study period

Regarding treatments, ustekinumab and IL-17 inhibitors were associated with a decreased risk of discontinuation compared with TNF inhibitors with HR of 0.42 (0.32–0.56), p < 10^{-4} and 0.36 (0.18–0.71), p = 0.004, respectively. In contrast, apremilast was associated with an increased risk of discontinuation, with an HR of 3.60 (2.67–4.83), p < 10^{-4}. Regarding clinical factors associated with drug survival, female sex and BMI > 30 kg/m² were associated with an increased risk of discontinuation with respective HR of 1.31 (1.08–1.58), p = 0.007 and 1.36 (1.07–1.73), p = 0.011 in multivariate analysis.

**DISCUSSION**

This large real-life setting study, did not find any association between calendar year of initiation and drug survival among patients with moderate to severe psoriasis conducting a multivariate Cox model and a segmented linear regression.
Regarding precisely the risk of drug discontinuation over time, the results of this study seem to be at variance with those of Shalom et al. (10) and Graier et al. (13), who reported that the calendar year of initiation was associated with an increased risk of discontinuation, particularly after the marketing of IL-17 inhibitors, using a Cox model and a Kaplan–Meier representation (10, 13). In this study, the authors assessed the drug survival for patients with psoriasis treated with biologics, irrespective of the number of treatment lines already received. To take into account time-period effects from the release of new drugs, they defined 2 periods; before and after 2016. Biologics’ drug survival decreased over time at an overall relative HR of 1.5 for the time-period (before and after 2016). However, interdependence analysis on drug survival using Cox regression revealed no treatment-independent effects of time-period. Such results were consistent with the current analyses. Moreover, contrary to Graier et al. (13), we adjusted for many confounders in order to assess the effect of the calendar year of initiation. The current results were also confirmed when we replicated the analyses for each biological agent. Therefore, the current findings suggest that, against the odds, drug survival of patients with moderate to severe psoriasis is not sensitive to the increasing therapeutic armamentarium, and that patients do not discontinue their treatments due to a high number of alternatives, but mostly in case of a suboptimal effectiveness/adverse reactions balance. Nevertheless, regarding etanercept specifically and contrary to expectations, an increasing survival rate was found after 2016. This may be explained by a sampling fluctuation in line with the decreasing number of patients treated with etanercept over years, making its drug survival less robust/precise than other treatments.

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alternatives, but mostly in case of a suboptimal effectiveness/adverse reactions balance. Nevertheless, regarding etanercept specifically, and contrary to expectations, the results showed an increasing survival rate after 2016. This may be explained by a sampling fluctuation in line with the decreasing number of patients treated with etanercept over years, making its drug survival less precise than other treatments.

This study confirms the good and similar drug survival of IL-17 inhibitors and ustekinumab (14). It also confirms the poor drug survival of apremilast, already observed by Sbidian et al. (15) compared with methotrexate. Here, the results are unambiguous compared with biologics, indicating its difficult placing in the increasing therapeutic armamentarium. These observations may be in line with the lower efficacy compared with biologics and with the higher rate of adverse events, with 79% of patients presenting at least one adverse event in the ESTEEM-1 trial (16).

Regarding age, sex and prevalence of PsA, the current results were similar to those of other cohorts (6–8). Nonetheless, in the current study, the overall mean retention rate is 0.67 at 1 year, which is lower than the 2 drug survival studies of the BADBIR cohort, with 0.77 and 0.79 retention rates at 1 year (6, 7), and the BIOBADADERM cohort, with 0.79 drug survival at 1 year (8). This difference is partially explained by the inclusion of apremilast and etanercept in the current model, which are likely to be associated with a decreased survival compared with other systemic treatments (15, 18). Nonetheless, taking separately, the mean retention rate at 1 year of adalimumab, ustekinumab or IL17 inhibitors are still lower than the BADBIR study, with 0.78 (0.77–0.79), 0.88 (0.87–0.89) and 0.88 (0.86–0.91), respectively. Hence, this difference might also be explained by a lower overall drug survival in France compared with other European countries, potentially associated with the particularities of the French health system (9).

Strengths

This study has several strengths. The first is its prospective large scale, associated with the real-life setting, as well as the capture of many clinical parameters, which are relevant in order to investigate drug survival. Moreover, the use of 2 different statistical models and sensitivity analyses, resulting in no effect of the calendar year of initiation on drug survival, strengthens these findings.

Limitations

This study also has several limitations. The first is a potential lack of power, mostly for subgroup analyses, due to scarce number of patients. Moreover, we cannot exclude that the profile of patients receiving bDMARDs and sDMARDs has changed over the years. Nevertheless, the analyses were adjusted on potential confounders, but could not include the Dermatology Life Quality Index (DLQI) because of too many missing values. Moreover, this study does not capture pharmacokinetics/pharmacodynamics parameters that could be relevant for drug discontinuation, especially the drug concentrations, such as the presence of antibodies. In addition, the study considered only first-line therapies in the analyses, but no second- or third-line therapies, whereas refractory patients with many previous therapeutic failures may be maintained on their treatment due to a lack of alternatives, and be switched to better ones after their approval. Last, but not least, it cannot be excluded that the length of study is insufficient over years, and that drug survival would decrease for patients initiating a treatment after 2018, e.g. in 2019 or 2020 and subsequent years, or after 1 year of follow-up, mainly because of the marketing of IL23p19 inhibitors.

Conclusion

Despite the increasing therapeutic armamentarium over the years, the calendar year of initiation does not seem to have any impact on drug survival in patients with moderate to severe psoriasis, which seems more linked to treatment, sex, or obesity. This finding suggests that drug survival of patients with moderate to severe psoriasis is not sensitive to this increasing armamentarium, and therefore strengthens its use to estimate balance between treatment effectiveness and adverse reactions over time. Nonetheless, they require confirmation on larger cohorts or databases with an extended follow-up.

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Conflicts of interest: CP has been an investigator and consultant for AbbVie, Amgen, Boehringer Ingelheim, Celgene, GSK, Janssen,
LEO Pharma, Lilly, Novartis and Pfizer. HB has undertaken activities as a paid consultant, advisor or speaker for Abbvie, Almirall, Amgen, Baxalta, Biogen, Boehringer Ingelheim, Celgene, Janssen, Leo Pharma, Lilly, Novartis, Pfizer, UCB and Sun Pharmaceuti-
cals. He has also received grant funding from Pfizer; MB-B offers a
consultancy service and is an investigator for Abbvie, Amgen, Cel-
gene, Janssen, Leo Pharma, Lilly, MSD, Novartis and Pfizer; MV has undertaken activities as a paid consultant, advisor or speaker
for Janssen, Abbvie, MSD, Pfizer, Leo Pharma, Medac, Boehringer
Ingelheim, Novartis, Lilly and Arrow. EM is a consultant for No-
varris, Abbvie, Pfizer and Janssen, has also been an investigator for
Leo Pharma, Amgen, AstraZeneca, Abbvie, Novartis and Pfizer,
and has received speaker remuneration from Abbvie, Janssen,
Novartis and Pfizer. NB has been an investigator for Pfizer and
Novartis. She is a consultant for Janssen and has received speaker
remuneration from Janssen. DDJ is a consultant for MEDAC UCB
Fresenius Kabi, Biogen, Abbvie, Celgen, Novartis, Lilly, Janssen,
Pfizer and MSD. MAR is a consultant for Pfizer, Leo Pharma,
Janssen, Galderma, AbbbVi, Novartis, Pierre Fabre, Merck and
BMS. PJ is a consultant for Roche, GSK, Lilly, Principiabio and
Sanofi Aventis. FT is Head of the Clinical Research Unit at Pitie-
Salpetriére Hospital and the Centre de Pharmaco epidemiologie
(Cephepi) (Pharmaco-epidemiology Centre) of the Assistance
Pública – Hopitaux de Paris (Paris Hospitals). She has received
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is a member of the PsoBioTeq cohort scientific committee, OC is
not receive any personal remuneration from these companies. OC
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