Adherence and Persistence to Biological Drugs for Psoriasis: Systematic Review with Meta-Analysis

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Abstract:
Despite the large number of biologics currently available for moderate-to-severe psoriasis, poor adherence and persistence to therapy represent the main issues for both the clinical and economic management of psoriasis. However, the data about adherence and persistence to biologics in psoriasis patients are conflicting. Our aim was to produce summary estimates of adherence and persistence to biologicals in adult patients with psoriasis. We performed a systematic review and meta-analysis of observational studies, searching two databases (PubMed and Embase). Sixty-two records met the inclusion criteria, and a meta-analysis was conducted on fifty-five studies. Overall, the proportion of adherent and persistent patients to biological therapy was 0.61 (95% confidence interval: 0.48–0.73) and 0.63 (0.57–0.68), respectively. The highest proportions were found for ustekinumab, while the lowest ones were found for etanercept. The proportions of adherence and persistence to biological drugs in psoriasis patients are sub-optimal. Notably, both proportions largely differ between drugs, suggesting that a more rational use of biologics might ensure better management of psoriasis.

Keywords: psoriasis; biological drugs; anti-TNF-α; anti-IL-17; anti-IL-12/23; adherence; persistence

1. Introduction
Psoriasis affects about 30 million adults worldwide [1]. Genetic factors, as well as lifestyle (smoking, alcohol consumption, and diet), certain drugs, environmental factors, and various metabolic conditions, can promote the development and progression of psoriasis [2,3]. Although the etiopathogenesis of psoriasis is multifactorial, its clinical manifestation mainly results from both uncontrolled keratinocyte proliferation and the overproduction of inflammatory mediators, such as tumor necrosis factor-α (TNF-α), interleukin (IL)-17, IL-12, and IL-23. In particular, the activation of these pro-inflammatory molecules triggers a vicious circle that progressively exacerbates psoriasis [2].

Due to its peculiar clinical manifestation, psoriasis has a negative psychological impact on patients, deeply affecting their quality of life [4]. Moreover, patients with psoriasis usually have several comorbidities that further aggravate their clinical condition [1]. Therefore, adequate pharmacological treatment might ameliorate both disease severity and, indirectly, the psychosocial sphere of the individual.

The therapeutic armamentarium currently available for the management of psoriasis is mainly represented by anti-inflammatory drugs and immunomodulators. In particular, topical (i.e., corticosteroids, vitamin D3 derivatives, and keratolytic products) and systemic drugs, such as methotrexate and retinoids, are commonly used in the mild-to-moderate forms of psoriasis, while targeted biological drugs are recommended for patients with severe forms who fail to respond to first-line therapy. TNF-α inhibitors were the first...
biologics to obtain marketing authorization and reimbursement for psoriasis, and they include etanercept (ETN), infliximab (INF), and adalimumab (ADA) [5]. Other biologics are IL17A inhibitors (ixekizumab, IXE; secukinumab, SECU) [6] and ustekinumab (UST), which is an anti-IL12/23 human monoclonal antibody [7].

Despite the large number of therapeutic options for the clinical management of psoriasis, two key contributors to both treatment failure and scarce relapse control are poor adherence and persistence to therapy. Adherence reflects “the extent to which a patient acts in accordance with the prescribed interval, and dose of a dosing regimen”, while persistence, also known as drug survival, is “the duration of time from initiation to discontinuation of therapy” [8]. In addition, suboptimal adherence and persistence deeply impact the economic management of psoriasis in healthcare systems [9], especially for the more expensive drugs (i.e., biologics). Therefore, improving medication-taking behaviors may help patients to better control therapy, as well as limiting the economic health expenditure. Currently, the data about adherence and persistence to biological therapy in psoriasis patients are scarce and conflicting, and previous systematic reviews, although quite recent [8,10], do not provide an exhaustive and quantitative synthesis of the literature. Moreover, real-world data about adherence and persistence to individual biologics are discordant, thus hindering the rational use of these drugs in clinical practice.

Hence, the aim of this systematic review and meta-analysis is to provide overall, updated adherence and persistence proportions to biologics, as well as reporting a stratification of results based on the individual biological drugs.

2. Materials and Methods

The protocol for this systematic review and meta-analysis was registered in the PROSPERO database (CRD42021245065).

2.1. Eligibility Criteria

We included prospective, retrospective, and cross-sectional observational studies evaluating adherence and persistence (or drug survival) to biologic drugs among participants aged 18 years or older with psoriasis. We considered studies irrespective of patient gender, comorbidities, or concomitant drugs. Biological drugs belonging to the following 3 classes were considered: TNF-\(\alpha\) inhibitors (ETN, ADA, INF); IL17A inhibitors (IXE, SECU); and IL12/23 inhibitors (UST). The outcomes were adherence and persistence to biologics, as reported in the included studies.

2.2. Information Sources and Search Strategy

We searched Medline and EMBASE for studies published from inception to 18 January 2021. The search strategy (Supplementary Material S1) reports psoriasis as the first term; drug or therapy adherence, persistence, compliance, and switching as the second term; and the considered biologic drugs as the third term (etanercept, ustekinumab, adalimumab, infliximab, ixekizumab, and secukinumab). The three terms were combined using the Boolean operator “AND”.

2.3. Selection Process

Titles and abstracts of papers identified by the search strategy were screened by two authors independently, E.P. and D.P. Each paper was categorized as not relevant or potentially included according to the eligibility criteria. Any disagreement was discussed with another author, E.L.

The full text of the potentially includible articles was retrieved or, if not available, directly requested from the authors of the study. Two authors (E.P. and D.P.) checked the full texts for the eligibility criteria and excluded studies not fitting them.

The selection process was managed using bibliographic management software Mendeley Desktop (v1.19.6, Mendeley Ltd., London, UK).
2.4. Data Extraction Process

We extracted the following information: study design, outcome (adherence or persistence), and objective; number and general characteristics of participants included in the studies, such as age, gender, comorbidities, and concomitant drugs (drugs used for the treatment of psoriasis, as well as other drugs); definition of adherence/persistence as reported in the study; number of adherent/persistent patients; and reasons for discontinuation/switching. Furthermore, the data relating to any stratifications were retrieved. The data extraction was carried out by two authors independently, E.P. and D.P., and any discrepancies were resolved through consultation with a third reviewer, E.L.

For the data collection, spreadsheet software Microsoft Excel was used (version 2102 build 13801.20864).

2.5. Study Risk of Bias Assessment

The methodological quality of included studies was assessed according to risk of bias in prevalence studies developed by Hoy et al. [11]. The tool considers ten domains concerning characteristics of prevalence studies, each rated in terms of risk of bias and applicability to research question. Risk of bias was judged from 0 (high risk) to 10 (low risk). The risk of bias was evaluated by two authors independently, E.P. and D.P., and any discrepancies were resolved through consultation with a third reviewer, E.L.

2.6. Effect Measures

We evaluated the study-specific prevalence of adherence or persistence (drug survival) to biologics by calculating the proportion of adherent or persistent subjects on the total number of participants for each study. Where the study provided adherence/persistence as a percentage or where the non-adherence/non-persistence was provided, appropriate calculations were performed.

2.7. Synthesis Methods

As adherence and persistence refer to two different concepts that cannot be matched and pooled, we separately analyzed these parameters, as previously reported by others [12]. In detail, three outcomes were evaluated in our meta-analysis: (1) adherence; (2) good adherence, generally reported as the medication possession ratio (MPR) or proportion of days covered (PDC) ≥ 80%; and (3) persistence.

Study-specific means of adherence were pooled using random effect models and the generic inverse variance method. Study-specific adherence/persistence proportions were pooled using random effect models with Freeman–Tukey transformation.

The heterogeneity for both methods was quantified through the Higgins heterogeneity index ($I^2$) and was tested through the chi-square test for mean adherence and Cochran’s Q test for adherence/persistence proportion.

Subgroup analyses were conducted according to study design (retrospective observational, prospective observational, or cross-sectional), type of biologics, the type of biologic users (biological-naïve subjects, i.e., subjects who have never used a biological drug, and biological-experienced subjects, i.e., subjects who have already had experience with this type of treatment), and study quality (high quality, score ≥ 8, vs. low quality, score < 8). Differences between groups was considered statistically significant if the heterogeneity test was significant.

$p$-value < 0.10 was considered statistically significant.

The “metagen” and “metaprop” routines within the META package in R (version 4.12) was used for analyses [13].

3. Results

3.1. Systematic Review

A flowchart of the search is presented in Figure 1. We identified 1285 records from the PUBMED search and 2698 from EMBASE. In total, 62 studies, including 169,371 par-
participants, met the inclusion criteria and were included in the qualitative synthesis. Three studies \[14–16\] did not show data on persistence or the number of persistent patients, while one did not show data on adherence \[17\]; two studies were conducted on patients not only affected by psoriasis \[18,19\] and did not report adherence data for psoriasis patients; one study \[20\] did not report the number of patients treated with each biological drug but only adherence as percentage. Fifty-five studies \[21–75\] on 161,748 participants were included in the quantitative synthesis (meta-analyses).

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In 13 studies \[24,25,28,32,34,35,38,39,55,64,66,70,75\], the sample was composed of patients with other chronic inflammatory autoimmune conditions, including osteoarticular diseases (such as ankylosing spondylitis and rheumatoid arthritis), bowel diseases (such as ulcerative colitis and Crohn’s disease), and psoriatic arthritis. The extraction of data, in this case, focused on the cohorts of patients suffering from psoriasis regardless of other conditions.

Among the included studies (Table 1), 5 studies presented data on adherence \[22,27,33,44,53\], 16 studies on good adherence \[21,25–27,32,33,35,40,44,46,49,53,63,68,71,73\], 46 studies on persistence data \[21,23–25,27–31,33,34,36–39,41–45,47–62,64–70,72,74,75\], and 8 studies reported data on both adherence and persistence \[21,25,27,33,44,49,53,68\]. Regarding study design (Table S1), 51 were retrospective cohort studies \[15,18,19,21–49,51–57,59–62,64,66–68,70,72,74,75\], 5 were prospective cohort studies \[14,50,58,65,69\], and 6 were cross-sectional studies \[16,17,20,63,71,73\]. The mean age of the participants was 47 years, of which about 45% were female. Thirty-two studies reported no use of concomitant drugs \[14,16–19,21,23–26,28–31,34,37,39,42,51,52,55,58,61,62,64,67,68,70,71,73,75\]. Twenty-four studies presented data on biological-naïve patients \[26–32,37–39,41,43,46,48,50,51,53,55,60,61\].
while four studies [37,39,50,75] reported data on biological-experienced patients. Twenty-eight studies reported data on ADA adherence/persistence [26,27,29,31–33,35,36,39,48–53,55,56,63,66–68,70,74,75], fifteen on INF [27,29,36,39,47,49,50,52,53,55,57,64,67,70,75], twenty-five on ETN [22,26,27,29–32,36,39,48–53,55,56,63,66–68,70,74,75], four on IXE [28,33,44,54], ten on SECU [24,28,32,34,44,59,61,68–70], and twenty-one on UST [23,26,27,31,32,39,40,48,50,51,53,60,61,63,64,67–70,72,75]. Finally, forty-five studies were included in biological drug subgroup analysis [21,23,24,26–36,39,40,42–44,46–56,59–61,63–70,72–75] and twenty-eight in experienced/naïve subgroup analysis [26–32,35,37–39,41,43,46,48,50,51,53,55,60,61,66,68,74,75].

Table 1. Details of calculation methods in considered outcomes.

| No. of Studies (No. of Patients) | Adherence | Good adherence | Persistence |
|----------------------------------|-----------|----------------|-------------|
| **Adherence**                    |           |                |             |
| MPR/PDC mean                     |           |                |             |
| during a period of 12 months     | 2 (4832)  | [27,53]        |             |
| during a period of >12 months    | 3 (6297)  | [22,33,44]     |             |
| **Good adherence**               |           |                |             |
| MPR/PDC ≥ 80%                    |           |                |             |
| during a period of 12 months     | 6 (29,256)| [25–27,49,53,68]|
| during a period of >12 months    | 5 (11,516)| [32,33,35,40,44]|
| Other definitions during different or not-specified periods a | 5 (4480) | [21,46,63,71,73] |
| **Persistence**                  |           |                |             |
| No discontinuation or gap a or switch |         |                |             |
| during a period of <12 months   | 2 (1179)  | [23,72]        |             |
| during a period of 12 months    | 24 (114,864)| [24,27,28,31,37–39,43,45,48,49,51,53–55,60–62,64–66,68,74,75]|
| during a period of >12 months   | 11 (24,246)| [29,33,34,41,42,44,50,56,58,59,67,69] |
| during a not-specified period   | 1 (84)    | [52]           |             |
| Still on treatment              |           |                |             |
| after a period of <12 months    | 1 (378)   | [25]           |             |
| after a period of 12 months or more | 4 (2336) | [30,36,47,57] |
| Other definitions during different or not-specified periods | 2 (13,714) | [21,70] |

* Different permissible gaps (from 7 to 150 days).

Table 1 shows details on adherence, good adherence, and persistence. The measures were highly heterogeneous: 5 studies gave the mean of adherence using MPR or PDC measures defined during different periods; 16 studies gave the proportion of adherent patients by mainly using (11 out of 16) the cut-off of 80% of MPR or PDC measures defined during different periods; 46 gave the proportion of persistent patients by mainly using (38 out of 64) discontinuation or switch or no gap (from 7 to 150 days) concepts defined during different periods.

3.2. Risk of Bias in Studies

Seventeen studies [18,25–27,32,39,45,46,48,49,53,59,61,62,67–69] obtained a total score of 10 in quality assessment based on the scale of Hoy et al. [11], while three studies [34,42,73] scored less or equal than 6 points. Details on single domains can be found in Table S2.
3.3. Results of Synthesis

3.3.1. Adherence

The meta-analysis conducted on five studies including 11,129 patients showed a mean adherence of 65% (95% confidence interval, CI: 61–70%, Figure S1) with considerable heterogeneity ($I^2 = 99$%). Among 16 studies including 45,252 patients, the proportion of good adherence was 61% (48–73, Figure 2), with considerable heterogeneity ($I^2 = 100.0$%). Only 2 out of 16 studies reported the reasons for non-adherence, which were loss of efficacy and adverse events. Qualitative descriptions of the reasons are shown in Table S3.

**Figure 2.** Forest plot of proportions, and their 95% confidence intervals, of adherent patients.

Regarding the stratification according to the type of biologic drug, the highest adherence proportion (Table 2, Figures 3 and 4) was observed for UST (72%, 48–91), followed by INF (63%, 44–80), ADA (62%, 47–76), SECU (52%, 35–68), ETN (50%, 36–65), and, finally, IXE (46%, 43–48). The difference between groups was statistically significant ($p$-value = 0.04). ADA, ETN, and UST represent the three biological drugs most considered in the included studies, as the use of each of them was evaluated in a considerable number of studies compared to the others: 10 studies for the first drug [26,27,32,33,35,40,49,53,63,68], 8 for the second [26,27,32,40,49,53,63,68], and 7 for the third [26,27,32,40,53,63,68].

**Table 2.** Pooled proportions of adherent patients stratified according to study design, type of biological drug, and type of patient.

| Source | Events | Total | Proportion | 95% CI |
|--------|--------|-------|------------|--------|
| Armstrong, 2018 | 254 | 478 | 0.53 | [0.49; 0.58] |
| Blauvelt, 2020a | 1658 | 4314 | 0.36 | [0.37; 0.40] |
| Blauvelt, 2020b | 762 | 1797 | 0.42 | [0.40; 0.45] |
| Chan, 2013 | 29 | 29 | 1.00 | [0.86; 1.00] |
| Degli Esposti, 2014 | 139 | 378 | 0.37 | [0.32; 0.42] |
| Dommasch, 2018 | 9849 | 13,862 | 0.71 | [0.70; 0.72] |
| Doshi, 2018 | 1020 | 2707 | 0.38 | [0.36; 0.40] |
| Feldman, 2019 | 1900 | 5022 | 0.38 | [0.36; 0.39] |
| Gendelman, 2018 | 113 | 136 | 0.83 | [0.76; 0.89] |
| Ito, 2010 | 232 | 247 | 0.94 | [0.90; 0.97] |
| Li, 2014 | 1373 | 3249 | 0.42 | [0.41; 0.44] |
| Malatestinic, 2017 | 1377 | 3848 | 0.36 | [0.34; 0.37] |
| Murage, 2019a | 1112 | 2130 | 0.52 | [0.50; 0.54] |
| Sorensen, 2017 | 415 | 478 | 0.87 | [0.83; 0.90] |
| Wu, 2020 | 4491 | 6331 | 0.71 | [0.70; 0.72] |
| Zschocke, 2017 | 143 | 246 | 0.58 | [0.52; 0.64] |
| Total | 45,252 | 61 [48; 73] | 99.7% | 5205.80 | 0 |

**Table 2.** Pooled proportions of adherent patients stratified according to study design, type of biological drug, and type of patient.
Biological drug

| Drug  | Type of patient | Events | Total | Proportion | 95% CI | Heterogeneity: $\chi^2 = 2263.54 \ (p < 0.001), I^2 = 100\%$
|-------|----------------|--------|-------|------------|--------|---------------------------------
| ADA   | Biological naïve | 1368   | 3668  | 0.37       | [0.36; 0.39] | $\chi^2 = 0.04 \ (p > 0.05), I^2 = 0\%$
| ADA   | Not specified   | 441    | 1084  | 0.41       | [0.38; 0.44] | $\chi^2 = 0.04 \ (p > 0.05), I^2 = 0\%$
| ETN   | Biological naïve | 114    | 2858  | 0.42       | [0.40; 0.43] | $\chi^2 = 0.04 \ (p > 0.05), I^2 = 0\%$
| ETN   | Not specified   | 107    | 2664  | 0.40       | [0.37; 0.44] | $\chi^2 = 0.04 \ (p > 0.05), I^2 = 0\%$
| INF   | Biological naïve | 113    | 2686  | 0.42       | [0.39; 0.45] | $\chi^2 = 0.04 \ (p > 0.05), I^2 = 0\%$
| INF   | Not specified   | 107    | 2503  | 0.43       | [0.40; 0.47] | $\chi^2 = 0.04 \ (p > 0.05), I^2 = 0\%$
| UST   | Biological naïve | 457    | 982   | 0.47       | [0.43; 0.50] | $\chi^2 = 0.04 \ (p > 0.05), I^2 = 0\%$
| UST   | Not specified   | 96     | 198   | 0.49       | [0.45; 0.54] | $\chi^2 = 0.04 \ (p > 0.05), I^2 = 0\%$
| IXE   | Biological naïve | 52     | 70    | 0.74       | [0.68; 0.81] | $\chi^2 = 0.04 \ (p > 0.05), I^2 = 0\%$
| IXE   | Not specified   | 79     | 108   | 0.73       | [0.67; 0.80] | $\chi^2 = 0.04 \ (p > 0.05), I^2 = 0\%$
| SECU  | Biological naïve | 157    | 318   | 0.49       | [0.44; 0.55] | $\chi^2 = 0.04 \ (p > 0.05), I^2 = 0\%$
| SECU  | Not specified   | 135    | 234   | 0.58       | [0.51; 0.64] | $\chi^2 = 0.04 \ (p > 0.05), I^2 = 0\%$
| Not specified | 5     | 4380  | 65.03 | 0.04       | [0.00; 0.08] | $\chi^2 = 0.04 \ (p > 0.05), I^2 = 0\%$

Total: 10,340.62
Heterogeneity: $\chi^2 = 2263.54 \ (p < 0.001), I^2 = 100\%$

Biological drug = INF

| Drug  | Type of patient | Events | Total | Proportion | 95% CI | Heterogeneity: $\chi^2 = 33.15 \ (p < 0.001), I^2 = 94\%$
|-------|----------------|--------|-------|------------|--------|---------------------------------
| INF   | Biological naïve | 3     | 650   | 0.94       | [0.85; 1.00] | $\chi^2 = 0.04 \ (p > 0.05), I^2 = 0\%$
| INF   | Not specified   | 79     | 108   | 0.73       | [0.67; 0.80] | $\chi^2 = 0.04 \ (p > 0.05), I^2 = 0\%$
| Not specified | 5     | 4380  | 65.03 | 0.04       | [0.00; 0.08] | $\chi^2 = 0.04 \ (p > 0.05), I^2 = 0\%$

Total: 10,340.62
Heterogeneity: $\chi^2 = 33.15 \ (p < 0.001), I^2 = 94\%$

Biological drug = ETN

| Drug  | Type of patient | Events | Total | Proportion | 95% CI | Heterogeneity: $\chi^2 = 1444.93 \ (p < 0.001), I^2 = 100\%$
|-------|----------------|--------|-------|------------|--------|---------------------------------
| ETN   | Biological naïve | 30     | 39    | 0.77       | [0.61; 0.89] | $\chi^2 = 0.04 \ (p > 0.05), I^2 = 0\%$
| ETN   | Not specified   | 135    | 234   | 0.58       | [0.51; 0.64] | $\chi^2 = 0.04 \ (p > 0.05), I^2 = 0\%$
| Not specified | 5     | 4380  | 65.03 | 0.04       | [0.00; 0.08] | $\chi^2 = 0.04 \ (p > 0.05), I^2 = 0\%$

Total: 10,340.62
Heterogeneity: $\chi^2 = 1444.93 \ (p < 0.001), I^2 = 100\%$

Biological drug = SEC

| Drug  | Type of patient | Events | Total | Proportion | 95% CI | Heterogeneity: $\chi^2 = 0.28 \ (p > 0.05), I^2 = 0\%$
|-------|----------------|--------|-------|------------|--------|---------------------------------
| SEC   | Biological naïve | 128    | 275   | 0.47       | [0.41; 0.53] | $\chi^2 = 0.04 \ (p > 0.05), I^2 = 0\%$
| SEC   | Not specified   | 463    | 1152  | 0.40       | [0.37; 0.43] | $\chi^2 = 0.04 \ (p > 0.05), I^2 = 0\%$
| Not specified | 5     | 4380  | 65.03 | 0.04       | [0.00; 0.08] | $\chi^2 = 0.04 \ (p > 0.05), I^2 = 0\%$

Total: 1251.80
Heterogeneity: $\chi^2 = 0.28 \ (p > 0.05), I^2 = 0\%$

Biological drug = UST

| Drug  | Type of patient | Events | Total | Proportion | 95% CI | Heterogeneity: $\chi^2 = 128.11 \ (p < 0.001), I^2 = 98\%$
|-------|----------------|--------|-------|------------|--------|---------------------------------
| UST   | Biological naïve | 149    | 160   | 0.93       | [0.88; 0.97] | $\chi^2 = 0.04 \ (p > 0.05), I^2 = 0\%$
| UST   | Not specified   | 1655   | 2023  | 0.82       | [0.80; 0.83] | $\chi^2 = 0.04 \ (p > 0.05), I^2 = 0\%$
| Not specified | 5     | 4380  | 65.03 | 0.04       | [0.00; 0.08] | $\chi^2 = 0.04 \ (p > 0.05), I^2 = 0\%$

Total: 1679.72
Heterogeneity: $\chi^2 = 128.11 \ (p < 0.001), I^2 = 98\%$

Biological drug = NotSpecified

| Drug  | Type of patient | Events | Total | Proportion | 95% CI | Heterogeneity: $\chi^2 = 114.71 \ (p < 0.001), I^2 = 97\%$
|-------|----------------|--------|-------|------------|--------|---------------------------------
| NotSpecified | 19    | 15681  | 63.81 | 0.01       | [0.00; 0.02] | $\chi^2 = 0.04 \ (p > 0.05), I^2 = 0\%$

Total: 114.71
Heterogeneity: $\chi^2 = 114.71 \ (p < 0.001), I^2 = 97\%$

Test for subgroup differences: $\chi^2 = 13.03 \ (p = .04)$. $\chi^2 = 33.04 \ (p < 0.001)$. $\chi^2 = 144.49 \ (p < 0.001)$.
There were differences stratifying by study design, with the cross-sectional design (85%; 55–100) showing a higher adherence compared to the retrospective cohort design (54%; 43–66) ($p$-value from subgroup test = 0.06) (Table 2 and Figure S2); however, only 3 studies had a cross-sectional design in contrast with 13 retrospective cohort studies. There were no differences stratifying by biological-naive patients and not-specified patients ($p$-value = 0.24) (Table 2 and Figure S3) or stratifying by risk of bias ($p$-value = 0.40) (Figure S4).

3.3.2. Persistence

The meta-analysis conducted on 46 studies including 156,801 patients showed a persistence proportion of 63% (57–68, Figure 5), with considerable heterogeneity ($I^2 = 100\%$).

Less than half of the studies (19 out of 46) reported the reasons for drug discontinuation or switching. The most common reasons were loss of efficacy and adverse events (nine studies) followed by ineffectiveness (three studies). Qualitative descriptions of the reasons are shown in Table S3.

![Figure 5. Forest plot of proportions, and their 95% confidence intervals, of persistent patients.](image)

Regarding the stratification according to the type of biological drug, the highest persistence (Table 3, Figures 6 and 7) was found for UST (77%, 70–84), followed by SECU (72%, 58–84), IXE (70%, 52–85), INF (64%, 60–68), ADA (57%, 50–63), and ETN (53%, 42–65).
The heterogeneity between groups was statistically significant ($p$-value < 0.01). ADA, ETN, and UST represent the three biological drugs most considered in the included studies, as the use of each of them was evaluated in a considerable number of studies compared to the others: 22 studies for the first drug [29,31,36,39,48,50–52,55,61,64–67,69,70,74,75], 19 for the second [29–31,36,39,48,50–52,55,56,66,67,70,74,75], and 17 for the third [23,31,39,48,50,51,60,61,64,67,69,70,72,75].

Table 3. Pooled proportions of persistent patients stratified according to study design, type of biological drug, and type of patient.

| Study design               | No. of Studies | No. of Patients | Persistence, % [CI 95%] | $I^2$ | Q      | $p$-Value for Heterogeneity within Strata | $p$-Value for Heterogeneity between Strata |
|----------------------------|----------------|----------------|------------------------|------|-------|------------------------------------------|------------------------------------------|
| Overall                    | 46             | 156,801        | 63 [57; 68]            |      |       | 19,429.06                               | 0                                        |
| Study design               |                |                |                        |      |       |                                          |                                          |
| Retrospective cohort       | 42             | 146,657        | 62 [56; 68]            | 100.0% | 16,496.95 | 0                                      |                                          |
| Prospective cohort         | 4              | 10,144         | 71 [63; 77]            | 96.1% | 76.69 | <0.001                                   |                                          |
| Biological drug            |                |                |                        |      |       |                                          |                                          |
| ADA                        | 22             | 21,176         | 57 [50; 63]            | 99.0% | 2428.85 | 0                                      |                                          |
| ETN                        | 19             | 12,914         | 53 [42; 65]            | 99.0% | 2770.05 | 0                                      |                                          |
| INF                        | 14             | 1465           | 64 [60; 68]            | 56.0% | 29.85 | 0.0049                                   |                                          |
| UST                        | 17             | 11,869         | 77 [70; 84]            | 98.0% | 1045.48 | <0.001                                 |                                          |
| IXE                        | 4              | 2155           | 70 [52; 85]            | 98.0% | 176.12 | <0.001                                 |                                          |
| SECU                       | 9              | 3053           | 72 [58; 84]            | 99.0% | 585.42 | <0.001                                 |                                          |
| Not specified              | 12             | 90,014         | 55 [44; 66]            | 100.0% | 9286.07 | 0                                      |                                          |
| Type of patient            |                |                |                        |      |       |                                          |                                          |
| Biological naïve           | 21             | 66,821         | 56 [49; 64]            | 100.0% | 5408.54 | 0                                      |                                          |
| Biological experienced     | 4              | 43,097         | 50 [35; 65]            | 100.0% | 1638.78 | <0.001                                 |                                          |
| Not specified              | 25             | 46,583         | 67 [60; 74]            | 100.0% | 5961.35 | 0                                      |                                          |

Different persistence proportions ($p$-value from heterogeneity test = 0.05) were observed among 21 studies on biological-naïve patients (56%, 49–64), 4 studies on biological-experienced patients (50%, 35–65), and 25 studies where it was not specified (67%, 60–74) (Table 3 and Figure S5). There were also statistical differences in study design stratification ($p$-value from subgroup test = 0.07) (Table 3 and Figure S6). However, only 4 studies had a cross-sectional design in contrast with 42 retrospective cohort studies. There were no statistical differences in the risk of bias stratification ($p$-value = 0.78) (Figure S7).
Figure 6. Forest plot of proportions, and their 95% confidence intervals, of persistent patients stratified according to biological drugs.
Another systematic review on rheumatoid arthritis [77] reported a median adherence value of 63% for both ETN and ADA. Finally, two recent meta-analyses on psoriasis showed that 66% of patients were persistent at 1 year [78] and 53.2% at two years [79].

In the studies included in our systematic review, the main reported reasons for drug discontinuation, switching, or non-adherence were loss of efficacy and adverse events. However, many other aspects could affect the patient’s behavior toward biological therapy. The female gender, recent disease onset, smoking, the presence of comorbidities, and a lack of efficacy of the previous treatments have been reported as predictors of non-persistence/non-adherence [42,71,80–82]. On the contrary, the presence of psoriatic arthritis has been generally associated with sustained drug survival of biological agents [69,81].

The variability in the included studies is reflected in the heterogeneity of our analysis. We found that biological-naïve patients were more persistent than biological-experienced patients. However, only four studies evaluated persistence in biological-experienced patients. Moreover, we observed a high percentage of adherent and persistent patients among cross-sectional and prospective cohort studies, respectively, compared to retrospective cohort ones. This is expected, even if only three studies evaluating adherence had a cross-sectional design and four studies evaluating persistence had a prospective cohort design. We did not investigate whether the inclusion of different definitions of the concepts of adherence and persistence influenced our results because few studies used the same definition. This represents a limitation of our study, as well as other meta-analyses aimed at pooling adherence and persistence. The proposal of a unified set of definitions might be useful to make the results of future studies more consistent and comparable [83].

At present, the data about adherence and persistence to individual biological drugs are quite scarce. This evaluation is essential to guide clinicians toward a more rational therapeutic choice, which is fundamental for both medical and economic purposes. In our study, the highest adherence was found for the human antibody UST (72%), followed by INF (63%), ADA (62%), SECU (52%), ETN (50%), and IXE (46%). Similar proportions were found for persistence as, in descending order, they were UST with 77%, SECU with 72%, IXE with 70%, INF with 64%, ADA with 57%, and ETN with 53%.

The variability in both adherence and persistence to specific biologics could derive, to a minimal extent, from the differences in the efficacy of treatments, which can reasonably affect patients’ satisfaction and, consequently, adherence/persistence to therapy. Future studies are required to elucidate on comparative efficacy because few data derived from direct “head-to-head” comparisons, and short-term efficacy outcomes were mainly evaluated [84–87]. A role for body mass index (BMI) in the patient’s attitude toward biological treatment has been recently proposed [88,89]. Indeed, the efficacy of TNF-α inhibitors and
UST is reduced in obese/overweight patients with psoriasis [90–92], with consequences for both adherence and persistence [93,94].

The difference between adherence and persistence to biological therapies can be certainly explained by discussing the origin, therapeutic class, administration route and timing, and toxicity profile of the biologics. Firstly, the immunogenic potential of chimeric antibodies (i.e., INF) might cause acute anaphylactic reactions following infusion, as well as hypersensitivity reactions (such as influenza-like syndrome, local skin reactions, and pyrexia) [95]. These phenomena can be counteracted with concomitant immunosuppressive therapy, with serious consequences on patient compliance and medication adherence/persistence [96]. On the contrary, pharmacological treatment with fully human antibodies (ADA, SECU, and UST) is less associated with anti-drug antibody production, although residual immunogenicity has been reported even for the most innovative biological drugs [97].

All biological drugs share the common risk of mild-to-moderate adverse events, including headache, cutaneous and upper respiratory tract infections, and injection site reactions, which can dramatically reduce quality of life [98]. Generally, these reactions do not require additional therapy, but they can be prevented by combining biological drugs with immunomodulators [99], with predictable detrimental effects on the patient’s compliance. Notably, TNF-α inhibitors are generally associated with a higher risk of severe infections, and they can induce hypersensitivity reactions [100] and dermatological disorders [101]. Hence, the peculiar toxicity profile of TNF-α inhibitors might explain, at least in part, the sub-optimal medication adherence and persistence to ETN, ADA, and INF in psoriatic patients.

Among the TNF-α inhibitors, ETN is self-administered using pre-filled syringes or pens up to twice a week. Both self-administration and short intervals between administrations might reduce compliance [102]. ADA is administered subcutaneously every 2 weeks. Therefore, the interval between administrations is longer than that reported for ETN, partially justifying the better adherence and persistence proportions to ADA rather than ETN. Finally, INF is intravenously administered at weeks 0, 2, and 6 after initiation and then at an 8- to 12-week interval [103]. The outpatient administration of INF ensures periodic support is provided to psoriasis patients, as well as contributing to a more assiduous monitoring of therapy by clinicians. Importantly, patients with scheduled appointments do not forget to take drugs, and they do not make mistakes, which instead can occur in self-administered therapy.

Concerning IL17A inhibitors, SECU is self-administered once a week for 4 weeks and then every 4 weeks [104], while IXE is self-administered every 2 weeks for the first 12 weeks and then once a month [105].

The highest adherence and persistence proportions were found for the IL12/23 inhibitor UST (72% for adherence and 77% for persistence). UST is characterized by high efficacy in the treatment of moderate-to-severe forms of psoriasis [106] and a favorable safety profile [107]. Moreover, it is administered every 12 weeks, exclusively under the guidance of an experienced physician. Both the longest administration interval and the supervision of a healthcare provider might favorably impact adherence and persistence [108]. The subcutaneous administration of UST using pre-filled syringes or pens might also explain the wide difference in the adherence and persistence proportions from another biological drug administered in hospitals or clinics, namely, INF (63% vs. 64%, respectively). Indeed, the latter therapy requires a slow 2 h infusion followed by an additional monitoring period of 2 h; it is a demotivating protocol that might partially contribute to scarce medication adherence and persistence.

In accordance with our findings, a comparative meta-analysis showed that UST has the longest persistence at 5 years after initiation compared with TNF-α inhibitors (ETN, ADA, and INF) [79]. In a meta-analysis of real-world evidence, UST appeared as the biological drug less frequently discontinued due to loss of efficacy [78], thus confirming its clinical relevance in the pharmacological treatment of psoriasis. On the contrary, ETN showed
the worst persistence and the highest number of therapy interruptions for low efficacy, supporting the results of our meta-analysis.

Even if UST is one of the most expensive biological drugs, it is endowed with one of the most favorable cost-efficacy profiles among the biological drugs for psoriasis [109]. Indeed, both sustained adherence and persistence and a sporadic dose regimen reduce the direct costs of treatment in the long term [110], but great attention should be paid to obese patients requiring high dosage [111]. In addition, UST is associated with minor indirect costs for the healthcare system, as it reduces hospital visits for non-responders; treatment failure; and resultant drug switching, which is associated with a 7–17% increase in annual costs [112]. There are, however, some crucial aspects that must be considered before initiating biological therapy with UST. Of course, UST must be avoided in patients with hypersensitivity to this biological drug or any of the excipients [110]. Moreover, health insurance coverage does not apply in all cases in real clinical practice. This latter aspect is reported to be responsible for short-term intermittent treatment with UST [113], as uninsured patients cannot afford the economic burden of continuous treatment with this biologic drug. Hence, the expansion of insurance coverage might ameliorate both patients’ satisfaction and adherence/persistence toward biological therapy. Finally, patient preferences should also be considered before starting therapy with UST, as involving patients in treatment decisions can influence both adherence to treatment and the outcomes of therapy [114].

5. Conclusions

The adherence and persistence to biological therapy in psoriasis patients are sub-optimal; however, the initial therapeutic choice might be crucial to ensure better medication adherence/persistence. Psoriasis patients are more adherent and persistent to therapies with a favorable safety profile and that are characterized by less frequent administrations (i.e., UST). However, several aspects regarding comorbidities, insurance coverage, patient preferences, and costs must be considered before initiating therapy with UST. We suggest that constant real-life therapeutic discussions between health providers (dermatologists, general practitioners, pharmacists, and nurses) and their patients, as well as specific support programs, might promote the optimal levels of adherence and persistence to biological drugs for both clinical and economic purposes.

Our study has several strengths, including the high number of studies identified and the large sample size, which gives consistency to the results. Our study also has some limitations, such as having considered work from all over the world; therefore, it cannot be excluded that adherence and persistence to treatment may have a link with reimbursement policies that vary from country to country. In addition, the follow-up period was variable from study to study, although most papers were aligned in considering 12 months as the follow-up period. We are also aware that we had to exclude some studies because of the lack of usable data, even though they met all inclusion and exclusion criteria. The age and sex of the participants could influence adherence/persistence; however, in the studies included in our meta-analysis, patients were very similar in terms of age and sex. Finally, although we included all drugs approved before May 2021 (data of our literature search) to manage moderate-to-severe psoriasis, we did not include studies investigating new groups of drugs, for example, selective inhibitors of IL-23 in a recently published study [115], and this represents a limitation of our systematic review.

Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/jcm11061506/s1, Supplementary Material S1: Search strategy; Table S1: Summary of findings of included studies; Table S2: Risk of bias for prevalence studies; Figure S1: Forest plot of mean adherence; Table S3: Qualitative description of the main reasons for biological drug discontinuation or switching; Figure S2: Forest plot of proportions of adherent patients with psoriasis to biological drugs stratified by study design; Figure S3: Forest plot of proportions of adherent patients with psoriasis to biological drugs stratified by type of patient; Figure S4: Forest plot of proportions of adherent patients with psoriasis to biological drugs stratified by risk of bias; Figure S5: Forest plot of proportions of prevalent patients with psoriasis to biological drugs stratified by...
type of patient; Figure S6: Forest plot of proportions of prevalent patients with psoriasis to biological drugs stratified by study design; Figure S7: Forest plot of proportions of prevalent patients with psoriasis to biological drugs stratified by risk of bias.

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**References**

1. Parisi, R.; Iskandar, I.Y.K.; Kontopantelis, E.; Augustin, M.; Griffiths, C.E.M.; Ashcroft, D.M. National, regional, and worldwide epidemiology of psoriasis: Systematic analysis and modelling study. *BMJ* 2020, 369, m1590. [CrossRef] [PubMed]
2. Kamiya, K.; Kishimoto, M.; Sugai, J.; Komine, M.; Ohtsuki, M. Risk factors for the development of psoriasis. *Int. J. Mol. Sci.* 2019, 20, 4347. [CrossRef] [PubMed]
3. Ito, T.; Takahashi, H.; Kawada, A.; Iizuka, H.; Nakagawa, H. Epidemiological survey from 2009 to 2012 of psoriatic patients in Japanese Society for Psoriasis Research. *J. Dermatol.* 2018, 45, 251–333. [CrossRef] [PubMed]
4. Bhosle, M.J.; Kulkarni, A.; Feldman, S.R.; Balkrishnan, R. Quality of life in patients with psoriasis. *Health Qual. Life Outcomes* 2006, 4, 35. [CrossRef]
5. Silva, L.C.R.; Ortigosa, L.C.M.; Benard, G. Anti-TNF-α agents in the treatment of immune-mediated inflammatory diseases: Mechanisms of action and pitfalls. *Immunotherapy* 2010, 2, 817–833. [CrossRef]
6. Giunta, A.; Ventura, A.; Chimenti, M.S.; Bianchi, L.; Esposito, M. Spotlight on ixekizumab for the treatment of moderate-to-severe plaque psoriasis: Design, development, and use in therapy. *Drug Des. Devel. Ther.* 2018, 12, 737–747. [CrossRef]
7. Ergen, E.N.; Yusuf, N. Inhibition of interleukin-12 and/or interleukin-23 for the treatment of psoriasis: What is the evidence for an effect on malignancy? *Exp. Dermatol.* 2018, 27, 737–747. [CrossRef]
8. Belinchón, I.; Rivera, R.; Blanch, C.; Comellas, M.; Lizán, L. Adherence, satisfaction and preferences for treatment in patients with psoriasis in the European union: A systematic review of the literature. *Patient Prefer. Adherence* 2016, 10, 2357–2367. [CrossRef]
9. De Vera, M.A.; Mailman, J.; Galo, J.S. Economics of Non-Adherence to Biologic Therapies in Rheumatoid Arthritis. *Curr. Rheumatol. Rep.* 2014, 16, 460. [CrossRef]
10. Aleshaki, J.S.; Cardwell, L.A.; Muse, M.E.; Feldman, S.R. Adherence and resource use among psoriasis patients treated with biologics. *Expert Rev. Pharm. Outcomes Res.* 2018, 18, 609–617. [CrossRef]
11. Hoy, D.; Brooks, P.; Woolf, A.; Blyth, F.; March, L.; Bain, C.; Baker, P.; Smith, E.; Buchbinder, R. Assessing risk of bias in prevalence studies: Modification of an existing tool and evidence of interrater agreement. *J. Clin. Epidemiol.* 2012, 65, 934–939. [CrossRef] [PubMed]
12. Ozaki, A.F.; Choi, A.S.; Le, Q.T.; Ko, D.T.; Han, J.K.; Park, S.S.; Jackevicius, C.A. Real-World Adherence and Persistence to Direct Oral Anticoagulants in Patients with Atrial Fibrillation: A Systematic Review and Meta-Analysis. *Circ. Cardiovasc. Qual. Outcomes* 2020, 13, e005969. [CrossRef] [PubMed]
13. Balduzzi, S.; Rücker, G.; Schwarzer, G. How to perform a meta-analysis with R: A practical tutorial. *Evid. Based Ment. Health* 2019, 22, 153–160. [CrossRef] [PubMed]
14. Dávila-Seijo, P.; Daudein, E.; Carretero, G.; Ferrandiz, C.; Vanaclocha, F.; Gómez-García, F.J.; Herrera-Ceballos, E.; De la Cueva-Dobao, P.; Belinchón, I.; Sánchez-Carazo, J.-L.; et al. Survival of classic and biological systemic drugs in psoriasis: Results of the BIOBADADERM registry and critical analysis. *J. Eur. Acad. Dermatol. Venereol.* 2016, 30, 1942–1950. [CrossRef]
15. Gniadecki, R.; Bang, B.; Bryld, L.E.; Iversen, L.; Lasthein, S.; Skov, L. Comparison of long-term drug survival and safety of biologic agents in patients with psoriasis vulgaris. *Br. J. Dermatol.* 2015, 172, 244–252. [CrossRef]
16. Yeung, H.; Wan, J.; Van Voorhees, A.S.; Callis Duffin, K.; Krueger, G.G.; Kalb, R.E.; Weisman, J.D.; Sperber, B.R.; Brod, B.A.; Schleicher, S.M.; et al. Patient-reported reasons for the discontinuation of commonly used treatments for moderate to severe psoriasis. J. Am. Acad. Dermatol. 2013, 68, 64–72. [CrossRef]

17. Wang, Q.; Luo, Y.; Lv, C.; Zheng, X.; Zhu, W.; Chen, X.; Shen, M.; Kuang, Y. Nonadherence to Treatment and Patient-Reported Outcomes of Psoriasis During the COVID-19 Epidemic: A Web-Based Survey. Patient Prefer. Adherence 2020, 14, 1403–1409. [CrossRef]

18. Bergman, M.; Patel, P.; Chen, N.; Jing, Y.; Saffore, C.D. Evaluation of Adherence and Persistence Differences Between Adalimumab Citrate-Free and Citrate Formulations for Patients with Immune-Mediated Diseases in the United States. Rheumatol. Ther. 2020, 8, 109–118. [CrossRef]

19. Marshall, J.K.; Bessette, L.; Shear, N.H.; Lebovic, G.; Glass, J.; Millson, B.; Gaetano, T.; Gazel, S.; Latour, M.G.; Laliberté, M.-C.; et al. Canada’s Study of Adherence Outcomes in Patients Receiving Adalimumab: 3-year Results from the COMPANION Study. Clin. Ther. 2018, 40, 1024–1032. [CrossRef]

20. Ichiyama, S.; Ito, M.; Funasaka, Y.; Abe, M.; Nishida, E.; Muramatsu, S.; Nishihara, H.; Kato, H.; Morita, A.; Imafuku, S.; et al. Assessment of medication adherence and treatment satisfaction in Japanese patients with psoriasis of various severities. J. Dermatol. 2018, 45, 727–731. [CrossRef]

21. Armstrong, A.W.; Foster, S.A.; Comer, B.S.; Lin, C.-Y.; Malatestinic, W.N.; Burge, R.; Goldblum, O. Real-world health outcomes in adults with moderate-to-severe psoriasis in the United States: A population study using electronic health records to examine patient-perceived treatment effectiveness, medication use, and healthcare resource utilization. BMC Dermatol. 2018, 18, 4. [CrossRef] [PubMed]

22. Bhosle, M.J.; Feldman, S.R.; Camacho, F.T.; Timothy Whitmire, J.; Nahata, M.C.; Balkrishnan, R. Medication adherence and health care costs associated with biologics in Medicaid-enrolled patients with psoriasis. J. Dermatol. Treat. 2006, 17, 294–301. [CrossRef] [PubMed]

23. Clemmensen, A.; Spon, M.; Skov, L.; Zachariae, C.; Gniadecki, R. Responses to ustekinumab in the anti-TNF agent-naïve vs. anti-TNF agent-exposed patients with psoriasis vulgaris. J. Eur. Acad. Dermatol. Venereol. 2011, 25, 1037–1040. [CrossRef] [PubMed]

24. Conesa-Nicolás, E.; García-Lagunar, M.H.; Núñez-Bracamonte, S.; García-Simón, M.S.; Mira-Sirvent, M.C. Persistence of secukinumab in patients with psoriasis, psoriatic arthritis, and ankylosing spondylitis. Famn. Hosp. 2020, 45, 16–21. [CrossRef] [PubMed]

25. Degli Esposti, L.; Sangiorgi, D.; Perrone, V.; Radice, S.; Clementi, E.; Perone, F.; Buda, S. Adherence and resource use among patients treated with biologic drugs: Findings from BEETLE study. Clinicoecon. Outcomes Res. 2014, 6, 401–407. [CrossRef] [PubMed]

26. Dommasch, E.D.; Lee, M.P.; Joyce, C.J.; Garry, E.M.; Gagne, J.J. Drug utilization patterns and adherence in patients on systemic medications for the treatment of psoriasis: A retrospective, comparative cohort study. J. Am. Acad. Dermatol. 2018, 79, 1061–1068.e1. [CrossRef]

27. Doshi, J.A.; Takeshita, J.; Pinto, L.; Li, P.; Yu, X.; Rao, P.; Viswanathan, H.N.; Gelfand, J.M. Biologic therapy adherence, discontinuation, switching, and restarting among patients with psoriasis in the US Medicare population. J. Am. Acad. Dermatol. 2016, 74, 1057–1065.e4. [CrossRef]

28. Egberg, A.; Bryld, L.E.; Skov, L. Drug survival of secukinumab and ixekizumab for moderate-to-severe plaque psoriasis. J. Am. Acad. Dermatol. 2019, 81, 173–178. [CrossRef]

29. Esposito, M.; Gisondi, P.; Cassano, N.; Ferrucci, G.; Del Giglio, M.; Loconsole, F.; Giunta, A.; Vena, G.A.; Chimenti, S.; Girolomoni, G. Survival rate of antitumour necrosis factor-α treatments for psoriasis in routine dermatological practice: A multicentre observational study. Br. J. Dermatol. 2013, 169, 666–672. [CrossRef]

30. Esposito, M.; Gisondi, P.; Cassano, N.; Babino, G.; Cannizzaro, M.V.; Ferrucci, G.; Chimenti, S.; Giunta, A. Treatment adherence to different etanercept regimens, continuous vs. intermittent, in patients affected by plaque-type psoriasis. Drug Dev. Res. 2014, 75 (Suppl. 1), S31–S34. [CrossRef]

31. Feldman, S.R.; Zhao, Y.; Navaratnam, P.; Friedman, H.S.; Lu, J.; Tran, M.H. Patterns of medication utilization and costs associated with the use of etanercept, adalimumab, and ustekinumab in the management of moderate-to-severe psoriasis. J. Manag. Care Spec. Pharm. 2015, 21, 201–209. [CrossRef] [PubMed]

32. Feldman, S.R.; Zhang, J.; Martinez, D.J.; Lopez-Gonzalez, L.; Marchlewicz, E.H.; Shrady, G.; Mendelsohn, A.M.; Zhao, Y. Real-world treatment patterns and healthcare costs of biologics and apremilast among patients with moderate-to-severe plaque psoriasis by metabolic condition status. J. Dermatolog. Treat. 2021, 32, 203–211. [CrossRef] [PubMed]

33. Bluemel, A.; Shi, N.; Burge, R.; Malatestinic, W.N.; Lin, C.-Y.; Lew, C.R.; Zimmerman, N.M.; Goldblum, O.M.; Zhu, B.; Murage, M.J. Comparison of real-world treatment patterns among patients with psoriasis prescribed ixekizumab or secukinumab. J. Am. Acad. Dermatol. 2020, 82, 927–935. [CrossRef] [PubMed]

34. Ferriéres, L.; Konstantinou, M.P.; Bulai Livideanu, C.; Hegazy, S.; Tauber, M.; Amelot, F.; Paul, C. Long-term continuation with secukinumab in psoriasis: Association with patient profile and initial psoriasis clearance. Clin. Exp. Dermatol. 2019, 44, e230–e234. [CrossRef]

35. Gendelman, O.; Weitzman, D.; Rosenberg, V.; Shalev, V.; Chodick, G.; Amital, H. Characterization of adherence and persistence profile in a real-life population of patients treated with adalimumab. Br. J. Clin. Pharmacol. 2018, 84, 786–795. [CrossRef]
36. Gniadecki, R.; Kragballe, K.; Dam, T.N.; Skov, L. Comparison of drug survival rates for adalimumab, etanercept and infliximab in patients with psoriatic vulgaris. *Br. J. Dermatol.* **2011**, *164*, 1091–1096. [CrossRef]

37. Hendrix, N.; Marcum, Z.A.; Veenstra, D.L. Medication persistence of targeted immunomodulators for plaque psoriasis: A retrospective analysis using a U.S. claims database. *Pharmacoepidemiol. Drug Saf.* **2020**, *29*, 675–683. [CrossRef]

38. Higa, S.; Devine, B.; Patel, V.; Baradaran, S.; Wang, D.; Bansal, A. Psoriasis treatment patterns: A retrospective claims study. *Curr. Med. Res. Opin.* **2019**, *35*, 1727–1733. [CrossRef]

39. Howe, A.; Ten Eyck, L.; Dufour, R.; Shah, N.; Harrison, D.J. Treatment patterns and annual drug costs of biologic therapies across indications from the Humana commercial database. *J. Manag. Care Spec. Pharm.* **2014**, *20*, 1236–1244. [CrossRef]

40. Husu, D.Y.; Gniadecki, R. Patient Adherence to Biologic Agents in Psoriasis. *Dermatology* **2016**, *232*, 326–333. [CrossRef]

41. Iskandar, I.Y.K.; Warren, R.B.; Lunt, M.; Mason, K.J.; Evans, I.; McElhone, K.; Smith, C.H.; Reynolds, N.J.; Ashcroft, D.M.; Griffiths, C.E.M. Differential Drug Survival of Second-Line Biologic Therapies in Patients with Psoriasis: Observational Cohort Study from the British Association of Dermatologists Biologic Interventions Register (BADBIR). *J. Investig. Dermatol.* **2018**, *138*, 775–784. [CrossRef] [PubMed]

42. Jacobi, A.; Rustenbach, S.J.; Augustin, M. Comorbidity as a predictor for drug survival of biologic therapy in patients with psoriasis. *Int. J. Dermatol.* **2016**, *55*, 296–302. [CrossRef] [PubMed]

43. Khalid, J.M.; Fox, K.M.; Globe, M.; Maguire, A.; Chau, D. Treatment patterns and therapy effectiveness in psoriasis patients initiating biologic therapy in England. *J. Dermatolog. Treat.* **2014**, *25*, 67–72. [CrossRef] [PubMed]

44. Blauvelt, A.; Shi, N.; Burge, R.; Malastenic, W.N.; Lin, C.-Y.; Lew, C.R.; Zimmerman, N.M.; Goldblum, O.M.; Zhu, B.; Murage, M.J. Comparison of Real-World Treatment Patterns Among Psoriasis Patients Treated with Ixekizumab or Adalimumab. *Patient Prefer. Adherence* **2020**, *14*, 517–527. [CrossRef]

45. Lee, S.; Xie, L.; Wang, Y.; Vaidya, N.; Basar, O. Evaluating the Effect of Treatment Persistence on the Economic Burden of Moderate to Severe Psoriasis and/or Psoriatic Arthritis Patients in the U.S. Department of Defense Population. *J. Manag. Care Spec. Pharm.* **2018**, *24*, 654–663. [CrossRef] [PubMed]

46. Li, Y.; Zhou, H.; Cai, B.; Kahler, K.H.; Tian, H.; Gabriel, S.; Arccona, S. Group-based trajectory modeling to assess adherence to biologics among patients with psoriasis. *Clinicoecon. Outcomes Res.* **2014**, *6*, 197–208. [CrossRef]

47. Magis, Q.; Jullien, D.; Gaudy-Marquèze, C.; Baumstark, K.; Viguier, M.; Bachelez, H.; Guibal, F.; Delaporte, E.; Karimova, E.; Montaudüh, H.; et al. Predictors of long-term drug survival for infliximab in psoriasis. *J. Eur. Acad. Dermatol. Venereol.* **2017**, *31*, 96–101. [CrossRef]

48. Mahlich, J.; Alba, E.; El Hadad, L.; Leisten, M.-K.; Peitsch, W.K. Drug Survival of Biological Therapies for Psoriasis Treatment in Germany and Associated Costs: A Retrospective Claims Database Analysis. *Adv. Ther.* **2019**, *36*, 1684–1699. [CrossRef]

49. Malastenic, W.; Nordstrom, B.; Wu, J.J.; Goldblum, O.; Solotkin, K.; Lin, C.-Y.; Kistler, K.; Fraeman, K.; Johnston, J.; Hawley, L.L.; et al. Characteristics and Medication Use of Psoriasis Patients Who May or May Not Qualify for Randomized Controlled Trials. *J. Manag. Care Spec. Pharm.* **2017**, *23*, 370–381. [CrossRef]

50. Menter, A.; Papp, K.A.; Gooderham, M.; Pariser, D.M.; Augustin, M.; Kerdel, F.A.; Fakharzadeh, S.; Goyal, K.; Calabro, S.; Langhoff, W.; et al. Drug survival of biologic therapy in a large, disease-based registry of patients with psoriasis: Results from the Psoriasis Longitudinal Assessment and Registry (PSOLAR). *J. Eur. Acad. Dermatol. Venereol.* **2016**, *30*, 1148–1158. [CrossRef]

51. Mercadal Orfila, G.; Ventayol Bosch, P.; Maestre Fullana, M.A.; Serrano L. Drug survival and discontinuation rates, and safety profile of secukinumab in real-world patients: A 152-week, multicenter, retrospective study. *J. Dermatolog. Treat.* **2021**, *32*, 664–674. [CrossRef] [PubMed]

52. Mocanu, M.; Toader, M.-P.; Rezus, E.; Tararu, T. Aspects concerning patient adherence to anti-TNFx therapy in psoriasis: A decade of clinical experience. *Exp. Ther. Med.* **2019**, *18*, 4987–4992. [CrossRef] [PubMed]

53. Murage, M.J.; Anderson, A.; Casso, D.; Oliveria, S.A.; Ojeh, C.K.; Muram, T.M.; Merola, J.F.; Zbrozek, A.; Araujo, A.B. Treatment patterns, adherence, and persistence among psoriasis patients treated with biologics in a real-world setting, overall and by disease severity. *J. Dermatolog. Treat.* **2019**, *30*, 141–149. [CrossRef] [PubMed]

54. Murage, M.J.; Gilligan, A.M.; Tran, O.; Goldblum, O.; Burge, R.; Lin, C.-Y.; Qureshi, A. Ixekizumab treatment patterns and healthcare utilization and costs for patients with psoriasis. *J. Dermatolog. Treat.* **2021**, *32*, 56–63. [CrossRef]

55. Bonañeda, M.; Fox, K.M.; Watson, C.; Princic, N.; Gandra, S.R. Treatment patterns in the first year after initiating tumour necrosis factor blockers in real-world settings. *Adv. Ther.* **2012**, *29*, 664–674. [CrossRef] [PubMed]

56. Onsun, N.; Güne¸ s, B.; Yabacı, A. Retention and survival rate of etanercept in psoriasis over 15 years and patient outcomes during the COVID-19 pandemic: The real-world experience of a single center. *Dermatol. Ther.* **2021**, *34*, e14623. [CrossRef]

57. Pogácsás, L.; Borsi, A.; Takács, P.; Remenyik, É.; Kemény, L.; Kárpáti, S.; Holló, P.; Wikonkál, N.; Gyulai, R.; Károlyi, Z.; et al. Long-term drug survival and predictor analysis of the whole psoriatic patient population on biological therapy in Hungary. *J. Dermatolog. Treat.* **2017**, *28*, 635–641. [CrossRef]

58. Ross, C.; Marshallman, G.; Grillo, M.; Stanford, T. Biological therapies for psoriasis: Adherence and outcome analysis from a clinical perspective. *Australas. J. Dermatol.* **2016**, *57*, 137–140. [CrossRef]

59. Ruiz-Villaverde, R.; Rodríguez-Fernandez-Freire, L.; Galán-Gutierrez, M.; Armario-Hlta, J.C.; Martínez-Pilar, L. Drug survival, discontinuation rates, and safety profile of secukinumab in real-world patients: A 152-week, multicenter, retrospective study. *Int. J. Dermatol.* **2020**, *59*, 633–639. [CrossRef]
107. Ghosh, S.; Gensler, L.S.; Yang, Z.; Gasink, C.; Chakravarty, S.D.; Farahi, K.; Ramachandran, P.; Ott, E.; Strober, B.E. Ustekinumab Safety in Psoriasis, Psoriatic Arthritis, and Crohn’s Disease: An Integrated Analysis of Phase II/III Clinical Development Programs. Drug Saf. 2019, 42, 751–768. [CrossRef]

108. Sandoval, L.F.; Huang, K.E.; Feldman, S.R. Adherence to ustekinumab in psoriasis patients. J. Drugs Dermatol. 2013, 12, 1090–1092.

109. Chi, C.-C.; Wang, S.-H. Efficacy and cost-efficacy of biologic therapies for moderate to severe psoriasis: A meta-analysis and cost-efficacy analysis using the intention-to-treat principle. Biomed Res. Int. 2014, 2014, 862851. [CrossRef]

110. Rouse, N.C.; Farhangian, M.E.; Wehausen, B.; Feldman, S.R. The cost-effectiveness of ustekinumab for moderate-to-severe psoriasis. Expert Rev. Pharm. Outcomes Res. 2015, 15, 877–884. [CrossRef]

111. D’Souza, L.S.; Payette, M.J. Estimated cost efficacy of systemic treatments that are approved by the US Food and Drug Administration for the treatment of moderate to severe psoriasis. J. Am. Acad. Dermatol. 2015, 72, 589–598. [CrossRef] [PubMed]

112. Feldman, S.R.; Evans, C.; Russell, M.W. Systemic treatment for moderate to severe psoriasis: Estimates of failure rates and direct medical costs in a north-eastern US managed care plan. J. Dermatolog. Treat. 2005, 16, 37–42. [CrossRef] [PubMed]

113. Choi, C.W.; Choi, J.Y.; Kim, B.R.; Youn, S.W. Economic Burden Can Be the Major Determining Factor Resulting in Short-Term Intermittent and Repetitive Ustekinumab Treatment for Moderate-to-Severe Psoriasis. Ann. Dermatol. 2018, 30, 179–185. [CrossRef] [PubMed]

114. Xu, Y.; Sudharshan, L.; Hsu, M.-A.; Koenig, A.S.; Cappelleri, J.C.; Liu, W.F.; Smith, T.W.; Pasquale, M.K. Patient preferences associated with therapies for psoriatic arthritis: A conjoint analysis. Am. Health Drug Benefits 2018, 11, 408–416. [PubMed]

115. Torres, T.; Puig, L.; Vender, R.; Lynde, C.; Piaserico, S.; Carrascosa, J.M.; Gisondi, P.; Daudén, E.; Conrad, C.; Mendes-Bastos, P.; et al. Drug Survival of IL-12/23, IL-17 and IL-23 Inhibitors for Psoriasis Treatment: A Retrospective Multi-Country, Multicentric Cohort Study. Am. J. Clin. Dermatol. 2021, 22, 567–579. [CrossRef]