RESEARCH ARTICLE

Predictors of drug survival for biologic and targeted synthetic DMARDs in rheumatoid arthritis: Analysis from the TRA Clinical Electronic Registry

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Abstract

In this study we aimed to identify the predictors of drug survival for biologic and targeted synthetic DMARDs (bDMARDs and tsDMARDs) among patients with rheumatoid arthritis (RA) in a real-world setting. Data from RA patients receiving bDMARDs and tsDMARDs between 2007 and 2019 were extracted from the Taiwan Rheumatology Association Clinical Electronic Registry (TRACER). Patients were categorized into tumor necrosis factor-alpha (TNF-α) inhibitors, non-TNF-α inhibitors, and tofacitinib groups. The primary outcome was 3-year drug retention and the causes of bDMARDs and tsDMARDs discontinuation were recorded. Baseline demographic data before the initiation of bDMARDs and tsDMARDs treatment were analyzed to identify the predictors of 3-year drug survival. A total of 1,270 RA patients were recruited (TNF-α inhibitors: 584; non-TNF-α inhibitors: 535; tofacitinib: 151). The independent protective factors for 3-year drug survival were positive rheumatoid factor (RF) (HR: 0.48, 95% CI: 0.27–0.85, p = 0.013) and biologics-naïve RA (HR: 0.61, 95% CI: 0.39–0.94, p = 0.024). In contrast, positive anti-citrullinated protein antibody (ACPA) (HR: 2.24, 95% CI: 1.32–3.79, p = 0.003) and pre-existing latent tuberculosis (HR: 2.90, 95% CI: 2.06–4.09, p<0.001) were associated with drug discontinuation. RA patients treated with TNF-α inhibitors exhibited better drug retention, especially in the biologics-naïve subgroup (p = 0.037). TNF-α inhibitors were associated with lower cumulative incidence of discontinuation due to inefficacy and adverse events (both p<0.001). Baseline RF and ACPA positivity in abatacept-treated patients were associated with a better 3-year drug survival. However, negative ACPA levels predicted superior drug survival of TNF-α.
inhibitors and tofacitinib. In conclusion, bio-naïve status predicted better drug survival in TNF-α inhibitors-treated RA patients. RF and ACPA positivity predicted better abatacept drug survival. In contrast, ACPA negativity was associated with superior TNF-α inhibitors and tofacitinib survival.

Introduction

Rheumatoid arthritis (RA) is a chronic and debilitating form of arthritis, and is one of the most prevalent autoimmune inflammatory rheumatic diseases [1]. According to the American College of Rheumatology (ACR)/European League Against Rheumatism (EULAR) recommendations for management of RA, the aim of treatment should be to reach a target of sustained remission or low disease activity in every patient with either biologic disease-modifying anti-rheumatic drugs (bDMARDs) or targeted synthetic DMARDs (tsDMARDs) [2, 3]. Advances in targeted therapy of RA have shown efficacy in preventing bone erosion and joint deformities [4]. However, the optimal response rates among bDMARDs and tsDMARDs for achieving low disease activity and remission were shown to be no greater than 50% and 20%, respectively [4, 5]. Therefore, it is imperative to identify predictors of drug retention for bDMARDs and tsDMARDs in RA using registries and health care databases [6–10].

The Taiwan Rheumatology Association Clinical Electronic Registry (TRACER) is a prospective, non-randomized cohort that promotes the "treat-to-target (T2T) "strategy for RA nationwide. In Taiwan, the first bDMARD, etanercept, was made available to patients via Taiwan’s National Health Insurance (NHI) program in 2002. This marked the first time in Taiwan that a tumor necrosis factor-alpha (TNF-α) inhibitor had been used to treat active RA patients with inadequate response to methotrexate (MTX)-based conventional synthetic disease-modifying anti-rheumatic drugs (csDMARDs). Since then, a number of TNF-α and non-TNF-α inhibitors have also been approved. In December, 2014, the first Janus-kinase (Jak) inhibitor, tofacitinib, a tsDMARDs, was made available on Taiwan’s NHI. TRACER enrolls RA patients from across Taiwan. Therefore, TRACER provides a great opportunity to investigate drug survival of bDMARDs and tsDMARDs for the treatment of RA.

In a prior systemic review and meta-analysis, better drug survival was found in etanercept-treated RA patients compared with two other TNF-α inhibitors [6]. However, concomitant use of csDMARDs, longer disease duration before initiation of a bDMARDs and female sex were associated with inferior drug survival [6]. Moreover, insufficient efficacy, adverse drug reactions, and safety signals of serious infections and malignancies could all contribute to biologic discontinuation [10–12]. In addition, shorter disease duration, baseline low disease activity, and young age may predict the 6-month therapeutic response in RA [9]. However, predictors of long-term drug retention for bDMARDs and tsDMARDs in RA are still lacking.

In the 2015 ACR guideline for the treatment of RA with high disease activity, bDMARDs were classified into TNF-α inhibitors and non-TNF-α biologics [2]. Although, rheumatoid factor (RF) and/or anti-citrullinated protein antibody (ACPA), especially at high levels, have been shown to be associated with erosive disease and poor outcome in RA, they have not consistently been shown to predict response to a variety of bDMARDs [3]. For example, RF and ACPA did not appear to be predictive of the response to anti-TNF treatment [13]. However, both ACPA and RF were previously found to predict a good EULAR response to rituximab therapy [14]. Moreover, in the AMPLE Trial, RA patients with the highest quartile of ACPA levels responded more favorably to abatacept, but not adalimumab, another anti-TNF-α
bDMARD [15]. Several studies did not find an association of seropositivity and responses to tocilizumab treatment [7]. It seems that RF and ACPA positivity may predict differential therapeutic responses in TNF-α inhibitors and non-TNF-α inhibitors. In addition, whether RF and ACPA are capable of predicting therapeutic responses in tofacitinib remains largely unknown.

To answer this unsolved question, this study aimed to identify predictors of 3-year drug retention for bDMARDs and tsDMARDs in a real-world dataset, TRACER.

Materials and methods

Data source

To identify predictors for drug survival of bDMARDs and tsDMARD, data were extracted from the Taiwan Rheumatology Association Clinical Electronic Registry (TRACER, www.tracer.org.tw), an investigator-led, Taiwan Rheumatology Association (TRA)-supported nationwide project. TRACER enrolled patients with autoimmune inflammatory rheumatic diseases, including RA, systemic lupus erythematosus, ankylosing spondylitis, and psoriatic diseases. This registry is a web-based system, which allows Taiwanese rheumatologists to register baseline demographic data, disease activity, autoantibodies status, medication, therapeutic responses, and adverse events of patients with systemic autoimmune diseases before and at 3-monthly intervals during treatment with csDMARDs, bDMARDs and tsDMARDs. TRA rheumatologists from tertiary referral hospitals, community hospitals, and local clinics voluntarily contributed de-linked patient information to this program.

Study protocol

We conducted a 3-year drug survival study of bDMARDs and tsDMARDs in RA from the TRACER database. The diagnosis of established RA was done according to the ACR 1987 revised criteria and/or 2010 ACR/EULAR criteria for the classification of RA [16, 17]. Patients with active RA and who had started bDMARDs and tsDMARDs during Jan., 2007 to Aug., 2019 were eligible. They were inadequate responders to at least two csDMARDs including MTX and had a 28 joints-disease activity score (DAS28) > 5.1 [18]. The baseline demographic data, the date of RA diagnosis, comorbidities, as well as serum levels of RF or ACPA before bDMARDs or tsDMARDs therapy were collected. All electronic procedures in TRACER and anonymized data are provided in S1 File. Participants with a follow-up period of less than 3 years were excluded. Taichung Veterans General Hospital’s Ethics Committee approved the study (CE18190A), and waived the requirement for informed consent because the patients’ data were anonymized prior to analysis.

Treatment

Targeted therapies were classified as TNF-α bDMARDs (etanercept, adalimumab, golimumab), non-TNF-α bDMARDs (tocilizumab, abatacept, rituximab) and tsDMARDs (tofacitinib) treatment [2, 3]. In Taiwan, the reimbursement for bDMARDs and tsDMARDs in RA is only approved by NHI when a combination of MTX-based csDMARDs is prescribed. Therefore, these targeted therapies were administered in combination with csDMARDs unless participants were intolerant to MTX or csDMARDs.

Study outcome

The primary outcome was 3-year drug retention. It was defined as treatment duration from the start date to discontinuation date of bDMARDs or tsDMARDs or the end of the
observation period plus one dispensation interval, whichever came first. The adverse events during bDMARDs and tsDMARDs treatment were recorded. The causes of bDMARDs and tsDMARDs discontinuation reflected treatment efficacy and adverse events ascertained by treating physicians.

**Covariates of interests**
The baseline demographic data, disease duration, comorbidities, serum levels of RF or ACPA before bDMARDs or tsDMARDs therapy were extracted from TRACER. The RF IgM levels were measured by nephelometry (Dade Behring Inc., Newark, DE, USA, positive if $\geq 14$ IU/mL). The ACPA levels were determined by EliA CCP (Phadia, Nieuwegein, The Netherlands, positive if $\geq 10$ U/mL). Disease activity of RA was assessed by DAS28-ESR. Concomitant medications of glucocorticoids, csDMARDs, and previous exposure to bDMARDs or tsDMARDs were also recorded. Pre-existing comorbidities of hypertension, diabetes mellitus, cardiovascular disease, depression, and osteoporosis were obtained from medical records. Latent TB, hepatitis B carrier, and hepatitis C carrier statuses were identified following the risk management plan set forth by Taiwan’s Centers for Disease Control (CDC) and TRA [19, 20].

**Patient and public involvement**
We did not involve patients or the public in our work.

**Statistical analysis**
The demographic data of the continuous parameters are shown as mean ± standard deviation, and for the categorical variables as the number of patients. Chi-Square test and Kruskal-Wallis test were used to compare variables among patients in the TNF-$\alpha$, non- TNF-$\alpha$, and tofacitinib groups. Risk factors associated with 3-year drug survival were determined by Cox proportional hazard regression. Statistically significant variables in univariate analyses were included in a multivariable model using the enter method. The drug retention probability curves were calculated by the Kaplan-Meier method, and statistical significance among groups was analyzed by the Log-rank test. All data were analyzed using the Statistical Package for the Social Sciences (SPSS) version 23.0. Significance was set at $p < 0.05$.

**Results**

**Baseline demographic data**
A total of 1,270 RA patients (TNF-$\alpha$ inhibitors: 584; non- TNF-$\alpha$ inhibitors: 535; tofacitinib: 151) were extracted from TRACER (Table 1). Of note, RA patients in the tofacitinib group exhibited older age (years, 58.1, 46.1–65.6 vs. 53.4, 41.6–60.8, and 57.3, 47.4–64.2, $p<0.001$), shorter disease duration (years, 9.5, 5.6–13.4 vs. 12.1, 8.3–13.4 and 13.1, 8.9–14.8, $p<0.001$), lower RF/ACPA positivity rates (RF positivity rates 73.3% vs. 99.5% and 80.4%, $p<0.001$; ACPA positivity rates 69.1% vs. 82.0% and 76.6%, $p = 0.003$) and disease activity by DAS28-ESR (5.9, 5.2–6.5 vs. 6.4, 5.8–7.0, and 6.1, 5.5–6.6, $p<0.001$), lower glucocorticoid (mg per day, 5.0, 5.0–10.0 vs. 7.5, 5.0–10.0 and 7.5, 5.0–10.0, $p = 0.002$), but higher MTX doses (mg per week, 15.0, 0.0–15.0 vs. 12.5, 5.0–15.0 and 10.0, 0.0–15.0, $p<0.001$) compared with the TNF-$\alpha$ inhibitors and non- TNF-$\alpha$ inhibitors groups (by Chi-square test or Kruskal-Wallis test, TNF-$\alpha$ inhibitors vs. tofacitinib, all $p<0.01$; non-TNF-$\alpha$ inhibitors vs. tofacitinib, all $p<0.01$).
Predictors associated with 3-year drug survival

To identify independent factors associated with 3-year drug retention, Cox regression analysis was performed (Table 2). We found that RF positivity (hazard ratio, HR: 0.48, 95% CI: 0.27–0.85, \( p = 0.013 \)) and biologic-naïve status (HR: 0.61, 95% CI: 0.39–0.94, \( p = 0.024 \)) were protective factors for drug retention. However, ACPA positivity (HR: 2.24, 95% CI: 1.32–3.79, \( p = 0.003 \)) and latent TB infection (HR: 2.9, 95% CI: 2.06–4.09, \( p < 0.001 \)) were independent risk factors for drug discontinuation.

Drug survival curves by biologics-exposure status

Among all participants (Fig 1A) and bio-naïve patients (Fig 1B), the TNF-\( \alpha \) inhibitors group exhibited superior 3-year drug survival compared with the non-TNF-\( \alpha \) inhibitors group (pairwise comparison, all \( p < 0.001 \)). However, drug retention rates seemed comparable in biologics-experienced patients (Fig 1C, pairwise comparison, all \( p > 0.05 \)).
Drug survival curves by causes of discontinuation

Fig 2 displays the causes of drug discontinuation in treatment groups. In Fig 2A, it can be seen that patients taking TNF-α blockers were at lower risk for discontinuation due to inefficacy compared with those in the non-TNF-α blockers and tofacitinib groups (pairwise comparison, TNF-α inhibitors vs. non-TNF-α inhibitors, p<0.001; TNF-α inhibitors vs. tofacitinib, p = 0.001; non-TNF-α inhibitors vs. tofacitinib, p>0.05). Moreover, the tofacitinib and non-TNF-α inhibitors groups were at higher risk of discontinuation due to adverse events (Fig 2B, Table 2. Cox regression analysis of factors associated with 3-year drug survival in RA patients receiving bDMARDs and tsDMARDs treatment.

|                                | Univariate               | Multivariate              |
|--------------------------------|--------------------------|---------------------------|
|                                | HR | 95%CI | p value | HR | 95%CI | p value |
| Age                            |    |       |         |    |       |         |
| <65y                           | Reference | Reference |         | Reference | Reference |         |
| ≥65y                           | 1.54 | (1.18-2.01) | 0.002* | 0.96 | (0.64-1.42) | 0.830 |
| Gender                         |    |       |         |    |       |         |
| F                              | Reference | Reference |         | Reference | Reference |         |
| M                              | 0.83 | (0.59-1.16) | 0.272 |      |         |         |
| Disease duration, years        | 0.99 | (0.97-1.01) | 0.302 |      |         |         |
| RF positive                    | 0.66 | (0.48-0.92) | 0.015* | 0.48 | (0.27-0.85) | 0.013* |
| ACPA positive                  | 1.58 | (1.13-2.23) | 0.008** | 2.24 | (1.32-3.79) | 0.003** |
| ESR                            | 1.00 | (1.00-1.05) | 0.926 |      |         |         |
| CRP                            | 1.00 | (0.99-1.00) | 0.718 |      |         |         |
| DAS 28                         | 1.00 | (0.96-1.05) | 0.924 |      |         |         |
| Biologics-naive                | 0.62 | (0.48-0.79) | <0.001** | 0.61 | (0.39-0.94) | 0.024* |
| Hypertension                   | 1.27 | (0.99-1.62) | 0.056 |      |         |         |
| Diabetes Mellitus              | 1.55 | (1.09-2.21) | 0.016* | 1.52 | (0.94-2.45) | 0.089 |
| Cardiovascular disease         | 1.34 | (0.94-1.92) | 0.107 |      |         |         |
| Depression                     | 1.82 | (1.02-3.25) | 0.042* | 1.47 | (0.52-4.14) | 0.462 |
| Osteoporosis                   | 1.47 | (1.16-1.86) | 0.002** | 0.86 | (0.62-1.20) | 0.382 |
| Latent TB                      | 2.05 | (1.59-2.64) | <0.001** | 2.90 | (2.06-4.09) | <0.001** |
| HBV carrier                    | 0.91 | (0.57-1.45) | 0.693 |      |         |         |
| HCV carrier                    | 2.39 | (1.62-3.51) | <0.001** | 1.17 | (0.55-2.47) | 0.679 |
| bDMARDs & tsDMARDs             |    |       |         |    |       |         |
| TNF-α inhibitors               |    |       |         |    |       |         |
| Non-TNF-α inhibitors           | 1.73 | (1.35-2.23) | <0.001** | 0.78 | (0.49-1.25) | 0.301 |
| Tofacitinib                    | 1.39 | (0.92-2.12) | 0.121 | 0.65 | (0.15-2.85) | 0.566 |
| Glucocorticoids dose           | 0.99 | (0.97-1.02) | 0.665 |      |         |         |
| MTX dose                       | 0.98 | (0.96-0.99) | 0.008** | 1.00 | (0.97-1.03) | 0.942 |
| SAL                            | 0.83 | (0.65-1.05) | 0.116 |      |         |         |
| HCQ                            | 0.91 | (0.71-1.17) | 0.478 |      |         |         |
| LEF                            | 1.44 | (1.09-1.90) | 0.009** | 1.38 | (0.90-2.13) | 0.141 |
| CsA                            | 1.08 | (0.78-1.48) | 0.644 |      |         |         |

By Cox proportional hazard regression.
* p<0.05
** p<0.01
ACCP, anti-citrullinated protein antibody; CRP, C reactive protein; CSA, cyclosporine; DAS28-ESR, the 28 joints-erythrocyte sedimentation rate measurement; ESR, erythrocyte sedimentation rate; GI disease, gastrointestinal disease; HBV, hepatitis B virus; HCV hepatitis C virus; LEF, leflunomide; MTX, methotrexate; n, number of patients included in analysis; RF, rheumatoid factor; SAL, salazopyrin; TB, tuberculosis; TNF-α, tumor necrosis factor-alpha.

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pairwise comparison, TNF-α inhibitors vs. non-TNF-α inhibitors, \( p = 0.014 \); TNF-α inhibitors vs. tofacitinib & non-TNF-α inhibitors vs. tofacitinib, \( p > 0.05 \) (C) all \( p > 0.05 \).

### Individual drug survival curves by RF and ACPA positivity

Since RF and ACPA were independent factors associated with drug survival, we examined the seropositivity status and drug retention in various treatment groups (Fig 3). In the TNF-α inhibitors and tofacitinib groups, we found that ACPA negativity was associated with a better 3-year drug retention probability (Fig 3B and 3J, ACPA (+) vs. ACPA (-) in TNF-α inhibitors, \( p < 0.001 \) and tofacitinib, \( p = 0.025 \)). In contrast, RF and ACPA positivity predicted better drug
Fig 3. The 3-year Kaplan-Meier drug retention probability of TNF-α inhibitors (A, B), abatacept (C, D), tocilizumab (E, F), rituximab (G, H), and tofacitinib (I, J) by RF and ACPA positivity. TNF-α, tumor necrosis factor-alpha; RF, rheumatoid factor; ACPA, anti-citrullinated protein antibody.

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survival in abatacept-treated patients (Fig 3C and 3D, RF(+) vs. RF (-), \( p < 0.001 \) and ACPA(+) vs. ACPA(-), \( p = 0.001 \)). Interestingly, seropositivity appeared to have no impact on drug retention in the tocilizumab and rituximab groups (Fig 3E-3H, all \( p > 0.05 \)).

Discussion

In this study, we aimed to investigate the predictors of 3-year drug survival for bDMARDs and tsDMARDs in RA using a real-world dataset in Taiwan. We demonstrated that TNF-\( \alpha \) inhibitors, non-TNF-\( \alpha \) inhibitors, and tofacitinib appeared to have differential long-term drug retention rates. Our study showed that bio-naïve status predicted better drug survival in TNF-\( \alpha \) inhibitors-treated RA patients. However, concomitant latent TB infection predicted drug discontinuation. RF and ACPA positivity predicted better abatacept drug survival. In contrast, ACPA negativity was associated with superior TNF-\( \alpha \) inhibitors and tofacitinib survival. The findings presented herein are the first to demonstrate that seropositivity seems to be a potential predictor for drug survival of bDMARDs and tsDMARDs in RA. This novel finding might shed light on the use of RF and ACPA as biomarkers for precision treatment and drug retention in RA.

Our results indicate that RA patients who received TNF-\( \alpha \) inhibitors exhibited better 3-year drug retention compared with their counterparts. This could be related to the fact that etanercept and adalimumab were the first two bDMARDs approved in Taiwan. Therefore, a high proportion (95.2%) of the TNF-\( \alpha \) antagonist group comprised bio-naïve patients from TRACER. We also showed that bio-naïve status was an independent and favorable factor for drug survival. This is consistent with previously reported drug persistence rates of TNF-\( \alpha \) inhibitors, which indicate they were better as a first-line biologic agent than as a second-line therapy for RA [8]. A greater proportion of anti-TNF-treated patients in our cohort were bDMARD-naïve patients, suggesting they might have a more rapid reduction in disease activity and greater improvements in physical function related to active RA in comparison with bDMARD-experienced patients [21].

In the present study, the concomitant latent TB infection was an independent risk predictor for drug discontinuation. Taiwan is an endemic area of latent TB infection and in over 80% of latent TB cases, a prophylaxis strategy was applied [22]. We previously demonstrated that the 1-year TB risk in RA patients receiving TNF-\( \alpha \) inhibitors was higher than that found among patients receiving non-TNF-\( \alpha \) inhibitors in a nationwide population-based study between 2008 and 2012 [23]. An Italian study also demonstrated that RA patients with latent TB might discontinue anti-TNF therapy, because active TB occurred during and after anti-TNF prophylactic therapy [24]. The disruption of granuloma integrity by anti-TNF therapy contributes to increased risk of latent TB reactivation [25]. Anti-TB prophylaxis could reduce TB reactivation by 65% [26]. Screening and prophylaxis of latent TB was advocated in 2012, and since then the incidence of TB infection has decreased. Moreover, the 5-year cumulative TB incidence between TNF-\( \alpha \) inhibitors and non-TNF-\( \alpha \) were indistinguishable [27]. Since our study enrolled participants before the era of universal screening and prophylaxis for latent TB, we found that it remained a risk factor for bDMARDs and tsDMARDs discontinuation.

Our study showed that patients with RA under anti-TNF treatment were less likely to discontinue targeted therapies due to inefficacy and adverse events. Immunogenicity to monoclonal antibodies of TNF-inhibitors could lead to the formation of anti-drug antibodies, and was correlated with loss of treatment efficacy [11, 28, 29]. However, non-TNF inhibitors were less frequently associated with immunogenicity [30]. Moreover, almost all RA under bDMARDs and tsDMARDs treatment in this study were combined with MTX-based csDMARDs, and thus our results might have been biased toward overestimation of the drug retention rates of
TNF-α inhibitors [2, 3]. A small molecule tsDMARD, tofacitinib, has not been reported to elicit immunogenicity. Nevertheless, our data showed that discontinuation of non-TNF inhibitors and tofacitinib may occur after 1 year of treatment. Further studies are needed to elucidate the mechanisms whereby patients may lose clinical responses to non-TNF inhibitors and tofacitinib.

The 3-year observational study showed that the tofacitinib and non-TNF-α inhibitors groups had a higher risk of discontinuation because of adverse events compared with the TNF-α inhibitors group. Accordingly, the adverse events for discontinuation of bDMARDs and tsDMARDs were mainly infection and drug intolerance especially in elderly patients with RA [10, 12]. A prior observational study reported that abatacept was associated with lower incidence rates of serious infections and severe infusion/injection reactions [31], as well as higher drug retention rates compared with other biologic agents [12]. Meanwhile, tocilizumab exhibited similar risk of opportunistic and bacterial infection but lower TB reactivation in comparison to anti-TNF therapy [32]. However, the prior anti-TNF experience was also associated with increased incidence of infection and serious infection during tocilizumab therapy [32, 33]. The same phenomenon was observed for B cell depletion agent, with a relatively high prevalence of severe pneumonia, and reactivation of HBV hepatitis and herpes zoster [34].

Previous reports also showed that the overall risk of infection, serious infection, and mortality rates in RA with tofacitinib were similar to those of other bDMARDs [35, 36], with a particular safety signal in herpes zoster infection especially in those receiving glucocorticoids [37]. Since the RA patients under anti-TNF treatment were younger and had a higher proportion of bio-naïve status, we cannot exclude the potential effects of confounders on the risk of discontinuation by adverse events.

Our study showed that RF positivity predicted a better 3-year drug survival in RA. Accordingly, seropositive RA patients share specific genetic and environment risk factors which differ from seronegative RA in clinical course and prognosis [38]. The autoantibodies of RF and ACPA in RA have direct pathogenic contributions to disease progression and seem to be a useful biomarker as a clinical predictor of drug survival [15, 39]. Systematic reviews indicate that neither RF nor ACPA status in RA patients is a predictor associated with response to TNF-α inhibitors [15, 40, 41]. However, our result demonstrated that the absence of baseline ACPA was associated with better drug survival of TNF-α inhibitors. Previous studies showed that high serum levels of RF were associated with poor treatment response of TNF-α inhibitors [42, 43]. Since RF- and ACPA-negative RA typically showed less bony erosion and structural damage, and exhibited a modest disease course [38], we speculate these patients might respond better to TNF-α inhibitors with less discontinuation due to inefficacy, leading to better drug retention rates.

Our result demonstrated that RF- and ACPA-positive RA patients receiving abatacept treatment exhibited better drug survival compared with the seronegative group. Inhibition of T cell co-stimulation factors may selectively affect autoantibody production [44, 45]. This was in keeping with the post-hoc analysis of the AMPLE study demonstrating that in RA patients with highest baseline anti-CCP2 antibody concentrations there was a stronger correlation with better clinical response to abatacept compared with those with lower concentrations [15]. Meanwhile, an international, prospective real-world study also showed that both RF and ACPA positivity were associated with higher abatacept retention [46]. Taken together, seropositivity of RF and ACPA could predict the likelihood of drug retention of T cell co-stimulation inhibitors.

A previous systematic review and meta-analysis showed that the baseline RF positivity in RA patients predicts better response to rituximab and tocilizumab [47]. However, a large cohort study showed contradictory results, i.e., neither baseline RF nor ACPA was a predictor of better response for tocilizumab therapy in RA [7]. Our results suggest that neither RF nor
ACPA status was a good predictor of drug survival of tocilizumab or rituximab. In Taiwan, rituximab is only approved for biologic-experienced RA patients; tocilizumab was also available on the NHIC program for second-line therapy for the first 3 years that it was on the market in Taiwan. Since biologic-experienced RA patients tend to respond less favorably to therapy, our result may not be extrapolated to bio-naïve patients.

The post-hoc analysis of tofacitinib treatment indicates that treatment outcome is not markedly influenced by autoantibody seropositivity [48]. Moreover, a higher proportion of tofacitinib-treated seropositive RA patients exhibited ACR20/50/70 responses, low disease activity, and remission in comparison with seronegative RA, especially with 10mg two times a day. Surprisingly, our study suggested that seronegative RA was associated with better tofacitinib drug retention. Since tofacitinib, a Jak1 and Jak3 kinase inhibitor, targets multiple cytokine receptors and exhibits diverse in vitro effects [49, 50], we postulate that seronegative RA patients might share a common pathology with the JAK pathway that could be targeted by tofacitinib. Further study is needed to confirm our finding and elucidate the underlying mechanisms.

Our study did have several limitations. First, the baseline characteristics among different treatment groups were not equally distributed. Although Cox regression analysis was performed to analyze independent factors associated with drug survival, there may have been confounding factors that were not completely controlled for. However, this also reflected the prescription behavior of rheumatologists in real-world observational studies. Second, various bDMARDs and tsDMARDs appeared on the market across a period spanning more than 10 years. In the early years, when only two TNF-α inhibitors, etanercept and adalimumab, were available, the limited choice of bDMARDs may have predisposed the treated patients to stay on these drugs. This might have led to an over-estimation of drug retention rates of TNF-α inhibitors. However, our study analyzed data from a period of more than a decade, which may provide long-term evidence of drug survival of bDMARDs and tsDMARDs. Third, the causes of drug discontinuation were diverse. We did not re-classify RA patients whose targeted therapies showed poor efficacy into primary and secondary treatment failure. Moreover, adverse events also included multiple diverse causes. A large inception cohort is needed to investigate the precise causes of drug discontinuation.

In conclusion, this long-term real-world study using the TRACER database demonstrated that bio-naïve status, latent TB infection, and RF/ACPA seropositivity were potential predictors for 3-year drug retention of bDMARDs and tsDMARDs in RA. Bio-naïve status was associated with better drug survival in TNF-α inhibitor-treated RA patients. RF and ACPA positivity seems to predict better abatacept drug retention. Conversely, ACPA-negative RA patients appeared to tolerate TNF-α inhibitors and tofacitinib therapies better than their counterparts. Thus, clinical parameters and autoantibody status may be used as a potential guide for targeted therapy in RA patients.

Supporting information
S1 File.
(XLSX)
S2 File.
(PDF)

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Author Contributions

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References

1. See LC, Kuo CF, Chou LJ, Chiu MJ, Yu KH. Sex- and age-specific incidence of autoimmune rheumatic diseases in the Chinese population: a Taiwan population-based study. Semin Arthritis Rheum. 2013; 43:381–6. https://doi.org/10.1016/j.semarthrit.2013.06.001 PMID: 23916348.

2. Singh JA, Saag KG, Bridges SL Jr., Akl EA, Bannuru RR, Sullivan MC et al. 2015 American College of Rheumatology Guideline for the Treatment of Rheumatoid Arthritis. Arthritis Rheumatol. 2016; 68:1–26. https://doi.org/10.1002/art.39480 PMID: 26545940.

3. Smolen JS, Landewe RBM, Bijlsma JWJ, Burmester GR, Dougados M, Kerschbaumer A et al. EULAR recommendations for the management of rheumatoid arthritis with synthetic and biological disease-modifying antirheumatic drugs: 2019 update. Ann Rheum Dis. 2020; 79:685–99. https://doi.org/10.1136/annrheumdis-2019-216655 PMID: 31969328.

4. Aletaha D, Smolen JS. Diagnosis and Management of Rheumatoid Arthritis: A Review. Jama. 2018; 320:1360–72. https://doi.org/10.1001/jama.2018.13103 PMID: 30285183.

5. Smolen JS, Aletaha D. Rheumatoid arthritis therapy reappraisal: strategies, opportunities and challenges. Nat Rev Rheumatol. 2015; 11:276–89. https://doi.org/10.1038/nrrheum.2015.8 PMID: 25687177.

6. Souto A, Maneiro JR, Gómez-Reino JJ. Rate of discontinuation and drug survival of biologic therapies in rheumatoid arthritis: a systematic review and meta-analysis of drug registries and health care databases. Rheumatology (Oxford). 2016; 55:523–34. https://doi.org/10.1093/rheumatology/kev374 PMID: 26490106.

7. Pers YM, Fortunet C, Constant E, Lambert J, Godfrin-Valnet M, De Jong A et al. Predictors of response and remission in a large cohort of rheumatoid arthritis patients treated with tocilizumab in clinical practice. Rheumatology (Oxford). 2014; 53:76–84. https://doi.org/10.1093/rheumatology/ket301 PMID: 24056521.

8. Choquette D, Bessette L, Alemao E, Haraoi B, Postema R, Raynauld JP et al. Persistence rates of abatacept and TNF inhibitors used as first or second biologic DMARDs in the treatment of rheumatoid arthritis: 9 years of experience from the Rhumadata(R) clinical database and registry. Arthritis Res Ther. 2019; 21:138. https://doi.org/10.1186/s13075-019-1917-8 PMID: 31170124.

9. Xiang Y, Wang Q, Li H, Duan X, Fang Y, Yang Pet al. Chinese registry of rheumatoid arthritis (CREDIT): III. The transition of disease activity during follow-ups and predictors of achieving treatment target. Int J Rheum Dis. 2020; 23:1719–27. https://doi.org/10.1111/1756-185X.13996 PMID: 33034424.

10. Favalli EG, Pontikaki I, Becciolini A, Biggiogero M, Ughi N, Romano M et al. Real-life 10-year retention rate of first-line anti-TNF drugs for inflammatory arthritides in adult- and juvenile-onset populations: similarities and differences. Clin Rheumatol. 2017; 36:1747–55. https://doi.org/10.1007/s10067-017-3712-8 PMID: 28597133.

11. Maneiro JR, Salgado E, Gomez-Reino JJ. Immunogenicity of monoclonal antibodies against tumor necrosis factor used in chronic immune-mediated inflammatory conditions: systematic review and meta-analysis. JAMA Intern Med. 2013; 173:1416–28. https://doi.org/10.1001/jamainternmed.2013.7490 PMID: 23797949.
12. Ebina K, Hashimoto M, Yamamoto W, Hirano T, Harra R, Katayama Met al. Drug tolerability and reasons for discontinuation of seven biologics in 4466 treatment courses of rheumatoid arthritis-the ANSWER cohort study. Arthritis Res Ther. 2019; 21:91. https://doi.org/10.1186/s13075-019-1880-4 PMID: 30971306.

13. Bobbio-Pallavicini F, Alpini C, Caporali R, Avalle S, Bugatti S, Montecucco C. Autoantibody profile in rheumatoid arthritis during long-term infliximab treatment. Arthritis Res Ther. 2004; 6:R264–72. https://doi.org/10.1186/ar1173 PMID: 15142273.

14. Sellam J, Hendel-Chavez H, Rouanet S, Abbé K, Combe B, Le Loet X et al. B cell activation biomarkers as predictive factors for the response to rituximab in rheumatoid arthritis: a six-month, national, multicenter, open-label study. Arthritis Rheum. 2011; 63:933–8. https://doi.org/10.1002/art.30233 PMID: 21225699.

15. Sokolove J, Schiff M, Fleischmann R, Weinblatt ME, Connolly SE, Johnsen A et al. Impact of baseline anti-cyclic citrullinated peptide-2 antibody concentration on efficacy outcomes following treatment with subcutaneous abatacept or adalimumab: 2-year results from the AMPLE trial. Ann Rheum Dis. 2016; 75:709–14. https://doi.org/10.1136/annrheumdis-2015-207942 PMID: 26359449.

16. Arnett FC, Edworthy SM, Bloch DA, McShane DJ, Fries JF, Cooper NS et al. The American Rheumatism Association 1987 revised criteria for the classification of rheumatoid arthritis. Arthritis Rheum. 1988; 31:315–24. https://doi.org/10.1002/art.1780310302 PMID: 3388796.

17. Aletaha D, Neogi T, Silman AJ, Funovits J, Felson DT, Bingham CO 3rd et al. 2010 Rheumatoid arthritis classification criteria: an American College of Rheumatology/European League Against Rheumatism collaborative initiative. Arthritis Rheum. 2010; 62:2569–81. https://doi.org/10.1002/art.27954 PMID: 20872595.

18. Prevoo ML, van ’t Hof MA, Kuper HH, van Leeuwen MA, van den Putte LB, van Riel PL. Modified disease activity scores that include twenty-eight-joint counts. Development and validation in a prospective longitudinal study of patients with rheumatoid arthritis. Arthritis Rheum. 1995; 38:44–8. https://doi.org/10.1002/art.1002 PMID: 854236.

19. Chen YH, Su WJ, Hsieh SJ, Yu KH, Jan YJ, Lee SS et al. 2015 TRA revised recommendations for screening and management of tuberculosis infection in patients before and after anti-tumor necrosis factor-alpha biological treatment. Formosan Journal of Rheumatology. 2015; 29:1–8.

20. Chen YH, Chien RN, Huang YH, Chen DY, Lan JL, L.Y. L et al. Screening and management of hepatitis B infection in rheumatic patients scheduled for biologic therapy: consensus recommendations from the Taiwan Rheumatology Association. Formosan Journal of Rheumatology. 2012; 26:1–7.

21. Karlsson JA, Kristensen LE, Kapetanovic MC, Gulfe A, Saxne T, Gborek P. Treatment response to a second or third TNF-inhibitor in RA: results from the South Swedish Arthritis Treatment Group Register. Rheumatology (Oxford). 2008; 47:507–13. https://doi.org/10.1093/rheumatology/ken034 PMID: 18304941.

22. Centers for Disease Control T. The TB and Latent TB Epidemiology from Notifiable Infectious Disease Statistics System. 2017.

23. Lim CH, Lin CH, Chen DY, Chen YM, Chao WC, Liao TL et al. One-Year Tuberculosis Risk in Rheumatoid Arthritis Patients Starting Their First Tumor Necrosis Factor Inhibitor Therapy from 2008 to 2012 in Taiwan: A Nationwide Population-Based Cohort Study. PLoS One. 2016; 11:e0166339. https://doi.org/10.1371/journal.pone.0166339 PMID: 27832150.

24. Cantilloni F, Labranche E, Marchesoni A, Mathieu A, Olivier I, Salvarani C et al. Latent tuberculosis infection detection and active tuberculosis prevention in patients receiving anti-TNF therapy: an Italian nationwide survey. Int J Rheum Dis. 2016; 19:799–805. https://doi.org/10.1111/1756-185X.12708 PMID: 26172207.

25. Baddley JW, Cantilloni F, Goletti D, Gomez-Reino JJ, Mylonakis E, San Juan R et al. ESCMID Study Group for Infections in Compromised Hosts (ESGICH) Consensus Document on the safety of targeted and biological therapies: an infectious diseases perspective (Soluble immune effector molecules [I]: anti-tumor necrosis factor-alpha agents). Clin Microbiol Infect. 2018; 24 Suppl 2:S10–s20. https://doi.org/10.1016/j.cmi.2017.12.025 PMID: 29459143.

26. Ai JW, Zhang S, Ruan QL, Yu YQ, Zhang BY, Liu QH et al. The Risk of Tuberculosis in Patients with Rheumatoid Arthritis Treated with Tumor Necrosis Factor-alpha Antagonist: A Metaanalysis of Both Randomized Controlled Trials and Registry/Cohort Studies. J Rheumatol. 2015; 42:2229–37. https://doi.org/10.3899/jrheum.150057 PMID: 26472414.

27. Lim CH, Chen HH, Chen YH, Chen DY, Huang WN, Tsai JJ et al. The risk of tuberculosis disease in rheumatoid arthritis patients on biologics and targeted therapy: A 15-year real world experience in Taiwan. PLoS One. 2017; 12:e0178035. https://doi.org/10.1371/journal.pone.0178035 PMID: 28570568.

28. Chen YH, Tsai WC, Tseng JC, Chen YH, Hsieh CW et al. Significant associations of antidrug antibody levels with serum drug trough levels and therapeutic response of adalimumab and etanercept
treatment in rheumatoid arthritis. Ann Rheum Dis. 2015; 74:e16. https://doi.org/10.1136/annrheumdis-2013-203893 PMID: 24442879.

29. Balsa A, Sanmarti R, Rosas J, Martin V, Cabez A, Gomez S et al. Drug immunogenicity in patients with inflammatory arthritis and secondary failure to tumour necrosis factor inhibitor therapies: the REASON study. Rheumatology (Oxford). 2018; 57:688–93. https://doi.org/10.1093/rheumatology/kex474 PMID: 29365183.

30. Strand V, Balsa A, Al-Saleh J, Barile-Fabris L, Horuchi T, Takeuchi T et al. Immunogenicity of Biologics in Chronic Inflammatory Diseases: A Systematic Review. BioDrugs. 2017; 31:299–316. https://doi.org/10.1007/s40259-017-0231-8 PMID: 28612180.

31. Ozen G, Pedro S, Schumacher R, Simon TA, Michaud K. Safety of abatacept compared with other biologic and conventional synthetic disease-modifying antirheumatic drugs in patients with rheumatoid arthritis: data from an observational study. Arthritis Res Ther. 2019; 21:141. https://doi.org/10.1186/s13075-019-1921-z PMID: 31174592.

32. Bykerk VP, Ostor AJ, Alvaro-Gracia J, Pavelka K, Ivorra JA, Graninger W et al. Tocilizumab in patients with active rheumatoid arthritis and inadequate responses to DMARDs and/or TNF inhibitors: a large, open-label study close to clinical practice. Ann Rheum Dis. 2012; 71:1950–4. https://doi.org/10.1136/annrheumdis-2011-201087 PMID: 22615456.

33. Mikulska M, Lanini S, Gudiol C, Drgona L, Ippolito G, Fernandez-Ruiz M et al. ESCMID Study Group for Infections in Compromised Hosts (ESGICH) Consensus Document on the safety of targeted and biological therapies: an infectious diseases perspective (Agents targeting lymphoid cells surface antigens [I]: CD19, CD20 and CD52). Clin Microbiol Infect. 2018; 24 Suppl 2:S71–s82. https://doi.org/10.1016/j.cmi.2018.02.003 PMID: 29447988.

34. Cohen S, Radominski SC, Gomez-Reino JJ, Wang L, Krishnaswami S, Wood SP et al. Analysis of infections and all-cause mortality in phase II, phase III, and long-term extension studies of tofacitinib in patients with rheumatoid arthritis. Arthritis Rheumatol. 2014; 66:2924–37. https://doi.org/10.1002/art.38779 PMID: 25047021.

35. Cohen S, Radominski SC, Gomez-Reino JJ, Wang L, Krishnaswami S, Wood SP et al. Analysis of infections and all-cause mortality in phase II, phase III, and long-term extension studies of tofacitinib in patients with rheumatoid arthritis. Arthritis Rheumatol. 2014; 66:2924–37.https://doi.org/10.1002/art.38779 PMID: 25047021.

36. Bykerk VP, Ostor AJ, Alvaro-Gracia J, Pavelka K, Ivorra JA, Graninger W et al. Tocilizumab in patients with active rheumatoid arthritis and inadequate responses to DMARDs and/or TNF inhibitors: a large, open-label study close to clinical practice. Ann Rheum Dis. 2012; 71:1950–4. https://doi.org/10.1136/annrheumdis-2011-201087 PMID: 22615456.

37. Salgado E, Maneiro JR, Carmona L, Gomez-Reino J. Rheumatoid factor and response to TNF antagonists in rheumatoid arthritis: systematic review and meta-analysis of observational studies. Joint Bone Spine. 2014; 81:41–50. https://doi.org/10.1016/j.jbspin.2013.04.004 PMID: 23731644.

38. Bobbio-Pallavicini F, Caporali R, Alpini C, Avallone S, Epis OM, Klersy C et al. High IgA rheumatoid factor levels are associated with poor clinical response to tumour necrosis factor alpha inhibitors in rheumatoid arthritis. Ann Rheum Dis. 2007; 66:302–7. https://doi.org/10.1136/ard.2006.060608 PMID: 17079248.

39. Bugatti S, Manzo A, Montecucco C, Caporali R. The Clinical Value of Autoantibodies in Rheumatoid Arthritis. Front Med (Lausanne). 2018; 5:339. https://doi.org/10.3389/fmed.2018.00339 PMID: 30560132.

40. Lv Q, Yin Y, Li X, Shan G, Wu X, Liang D et al. The status of rheumatoid factor and anti-cyclic citrullinated peptide antibody are not associated with the effect of anti-TNFalpha agent treatment in patients with rheumatoid arthritis: a meta-analysis. PLoS One. 2014; 9:e89442. https://doi.org/10.1371/journal.pone.0089442 PMID: 24586782.

41. De Rycke L, Verhelst X, Kruithof E, Van den Bosch F, Hoffman IE, Veyes EM et al. Rheumatoid factor, but not anti-cyclic citrullinated peptide antibodies, is modulated by infliximab treatment in rheumatoid arthritis. Ann Rheum Dis. 2005; 64:299–302. https://doi.org/10.1136/ard.2004.023523 PMID: 15166003.

42. Lorenzetti R, Janowska I, Smulski CR, Frede N, Henneberger N, Walter L et al. Abatacept modulates CD80 and CD86 expression and memory formation in human B-cells. J Autoimmun. 2019; 101:145–52. https://doi.org/10.1016/j.jaut.2019.04.016 PMID: 31054942.
45. Jansen D, Emery P, Smolen JS, Westhovens R, Le Bars M, Connolly SE et al. Conversion to seronegative status after abatacept treatment in patients with early and poor prognostic rheumatoid arthritis is associated with better radiographic outcomes and sustained remission: post hoc analysis of the AGREE study. RMD Open. 2018; 4:e000564. https://doi.org/10.1136/rmdopen-2017-000564 PMID: 29657830.

46. Nusslein HG, Alten R, Galeazzi M, Lorenz HM, Nurmohamed MT, Bensen WGet al. Efficacy and prognostic factors of treatment retention with intravenous abatacept for rheumatoid arthritis: 24-month results from an international, prospective, real-world study. Clin Exp Rheumatol. 2016; 34:489–99. PMID: 26986919.

47. Maneiro RJ, Salgado E, Carmona L, Gomez-Reino JJ. Rheumatoid factor as predictor of response to abatacept, rituximab and tocilizumab in rheumatoid arthritis: Systematic review and meta-analysis. Semin Arthritis Rheum. 2013; 43:9–17. https://doi.org/10.1016/j.semarthrit.2012.11.007 PMID: 23290690.

48. Bird P, Hall S, Nash P, Connell CA, Kwok K, Witcombe D et al. Treatment outcomes in patients with seropositive versus seronegative rheumatoid arthritis in Phase III randomised clinical trials of tofacitinib. RMD Open. 2019; 5:e000742. https://doi.org/10.1136/rmdopen-2018-000742 PMID: 30886732.

49. O’Shea JJ, Schwartz DM, Villarino AV, Gadina M, McInnes IB, Laurence A. The JAK-STAT pathway: impact on human disease and therapeutic intervention. Annu Rev Med. 2015; 66:311–28. https://doi.org/10.1146/annurev-med-051113-024537 PMID: 25587654.

50. Villarino AV, Kanno Y, O’Shea JJ. Mechanisms and consequences of Jak-STAT signaling in the immune system. Nat Immunol. 2017; 18:374–84. https://doi.org/10.1038/ni.3691 PMID: 28323260.