The longitudinal effect of biologic use on patient outcomes (disease activity, function, and disease severity) within a rheumatoid arthritis registry

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Abstract

Introduction Biologics effectively manage symptoms and disease activity in rheumatoid arthritis (RA), but their long-term effects remain unclear.

Method Longitudinal data were examined from the Brigham and Women’s Rheumatoid Arthritis Sequential Study (BRASS) registry. Linear regression modeled the effect of biologic exposure on changes in disease activity (Disease Activity Score-28 with C-reactive protein [DAS28-CRP]), functional status (modified Health Assessment Questionnaire [mHAQ]), and RA severity (Routine Assessment of Patient Index Data [RAPID3]). Biologic exposure was the ratio of time on a biologic relative to time participating in the BRASS cohort.

Results The analysis included 1395 RA patients, 82.3% female, with 6783 unique study visits from 2003 to 2015. At the patient’s first visit, mean (SD) age was 56.3 (14.2) years and mean (SD) duration of RA was 12.7 (11.9) years. Average follow-up duration was 5.59 years (range, 1–13). Over time, DAS28-CRP, mHAQ, and RAPID3 scores decreased as the biologic exposure ratio increased. In repeated measures regression models, increased biologic exposure was significantly associated with decreased DAS28-CRP score ($\beta = -0.647; P < 0.001$), decreased mHAQ score ($\beta = -0.096; P < 0.001$), and decreased RAPID3 score ($\beta = -0.724; P < 0.001$) during follow-up. Methotrexate use at baseline predicted decreased DAS28-CRP, mHAQ, and RAPID3 scores during follow-up. Biologic use at baseline predicted increased DAS28-CRP or mHAQ during follow-up.

Conclusions Increased biologic exposure is associated with decreased disease activity, function impairment, and RA severity. Future studies should examine whether earlier initiation of biologics improves patient outcomes in RA.

Trial Registration ClinicalTrials.gov, NCT01793103

Key Points

• Biologics effectively manage symptoms and disease activity in rheumatoid arthritis (RA), but their long-term effects remain unclear.
Introduction

Biologics have become the standard of care for treating moderate to severe rheumatoid arthritis (RA) in patients with an inadequate response to small molecule disease-modifying antirheumatic drugs (DMARDs) [1–3]. Although biologics have been proven effective in managing RA symptoms and disease activity across numerous clinical studies [4], their long-term effect on disease activity and patient function remains unclear. A national registry study in Sweden and a real-world prospective study of 4651 adults with RA in the USA both reported reduced disability due to RA after biologic treatment options were introduced in 1998, compared with the period before biologics were available [5, 6].

Control of inflammatory processes is known to limit joint damage and disease progression. It stands to reason that use of biologic DMARDs, especially early in disease, could help to reduce the overall severity of disease within the RA population [7–9]. Combining biologic therapy with nonbiologic DMARD therapy early in patients with recent onset of RA reduces joint damage and disability compared with sequential or “step-up” combination therapy with nonbiologic DMARDs followed by biologic therapy [7–9]. When tumor necrosis factor inhibitors became available for the treatment of RA, patients often initiated biologic therapy later in the treatment course, when disability was already more severe [10]. Later initiation of biologic therapy in patients with established RA has a similar effect on joint damage, but it may reduce disability less effectively [7, 11].

Treatment guidelines for RA have evolved over time to support earlier and more aggressive use of biologics [1, 12, 13]. Older guidelines recommended initiating biologic therapy in patients with early RA (<6 months) only for patients with features of poor prognosis [12, 13]. Based on the available low-to-moderate evidence, current guidelines recommend either adding biologic therapy or combining nonbiologic DMARDs in a patient whose disease activity remains moderate or high despite therapy with a small molecule DMARD, regardless of whether the patient has early RA or established RA [1]. The optimal time to start biologic therapy in RA remains unclear, but there is some evidence that starting biologic therapy earlier is associated with greater chance of achieving low disease activity [14].

Many studies of disease burden in RA patients are limited to 1 year or less in duration. Registry study data can be used to examine whether longer-term exposure to a biologic maintains the benefits for reduced disease activity and function impairment. Previous analyses of registry data have shown that RA patients who respond to biologic therapy in the first 6–12 months usually experience persistent benefits if they continue biologic therapy for several years [15–24]. The objective of this analysis was to examine the effect of cumulative exposure to biologics on patient outcomes over time, as well as predictors for improvement in outcomes.

Materials and methods

Data source

The Brigham and Women’s Rheumatoid Arthritis Sequential Study (BRASS) registry includes clinical and patient-reported outcome data from a cohort of more than 1400 RA patients from a single academic medical center at varying stages of disease progression. BRASS is an observational and longitudinal registry that follows patients over a period of 15 years.

The registry was established in March 2003 at the Robert Breck Brigham Arthritis Center in Boston, MA. Approximately 2700 variables are collected as part of the registry, including patient-reported data, physician assessments, and laboratory tests. Patients complete surveys to record patient-reported outcomes twice annually, while physician assessments and laboratory tests are performed at annual study visits.

Patients have entered the registry over time, with significant enrollment in the early years of the registry and in the early 2010s. Individual patients have contributed from 1 to 13 years of data depending, in part, on timing of enrollment, with an average of 5.59 years of participation. Years in which patients contributed data were not necessarily continuous. For example, a patient contributing 5 years of data could have provided survey data in 5 noncontinuous years of the registry. The BRASS registry has produced a large prospective database; analyses of the BRASS dataset have yielded insight into diagnosis, biomarkers, treatment regimens, and patient outcomes for the RA patient population [25–31].

Keywords Biologics · Disease activity · Exposure · Function · Rheumatoid arthritis · Severity
Study procedures and written informed consent were approved by the Institutional Review Board of Brigham and Women’s Hospital, Partners Health Care, Boston, MA. A copy of the protocol and the informed consent form for enrollment in the BRASS registry were submitted to the Institutional Review Board of Brigham and Women’s Hospital, and written approval was obtained.

**Biologic utilization**

Any exposure to a given medication or class of medications for RA in a calendar year was determined from the twice-annual patient self-report surveys. The biologic exposure ratio was calculated at each time point as the ratio of the patient’s total time on biologics (from the patient self-report surveys), divided by their total time in the BRASS registry, which resulted in a continuous proportion for each patient that ranged from 0 (no biologic use) to 1 (continuous biologic use). Because the biologic exposure ratio was calculated at every time point, an individual’s biologic exposure could vary over the course of the study.

**Outcomes**

Three outcomes were assessed in this analysis to understand clinical and functional status and disease severity: the Disease Activity Score-28 with C-reactive protein (DAS28-CRP), the modified Health Assessment Questionnaire (mHAQ), and the Routine Assessment of Patient Index Data (RAPID3). All outcome data were derived from the annual study visit. The DAS28 is a clinical composite score derived from patient-reported data, physician assessments, and laboratory tests that provide an assessment of disease activity [32]. The BRASS registry includes the DAS28-CRP, which is composed of tender and swollen joint counts (28 joints assessed), a C-reactive protein laboratory value, and a patient’s global assessment score for disease activity, using a visual analog scale. Greater DAS28-CRP scores indicate greater disease activity, with cutoffs for moderate and high severity at 3.2 and 5.1, respectively [33].

The BRASS registry also includes the mHAQ, a patient-reported questionnaire that monitors functional status in patients with rheumatic diseases [34]. The questionnaire includes eight activities of daily living (dressing; getting in/out of bed; lifting a full cup/glass; walking on flat ground; washing/drying entire body; bending down to pick up clothing off the floor; turning taps on/off; and getting in/out of a bus, car, train, or airplane). Scores range from 0 to 3, with lower scores indicating a better functional status.

The RAPID3 is another patient-reported scale in the BRASS registry that provides a quick assessment of RA severity [35]. It includes a subset of variables within the multidimensional Health Assessment Questionnaire, a patient global assessment for pain, and a patient global assessment for overall health. Scores range from 0 to 10, with cutoffs for remission (0.0–1.0), low disease activity (> 1.0–2.0), moderate disease activity (> 2.0–4.0), and high disease activity (> 4.0).

**Statistical methods**

Descriptive statistics (mean and standard deviation [SD] for continuous variables; frequencies and percentages for categorical variables) were calculated for patient demographics. Baseline values were derived from the patient’s first annual study visit in the BRASS registry.

The effect of the biologic exposure ratio on RA patient outcomes was assessed using linear mixed effects repeated measures models with binary and categorical variables treated as classes. These models controlled for the correlations among repeated measures on the same individual and were not affected by random missing data. Covariates included in initial models were to control for various patient demographic and treatment factors known to be clinically relevant to RA outcomes (Table 1). Model development began with analysis of clinically relevant covariates and culminated with the best fit model; clinically relevant covariates were determined by the study team, which included practicing rheumatologists. Covariates included in the final best fit model for each outcome were determined through an iterative process that considered overfitting and collinearity, among other factors. The best fit model was identified based on model fit statistics, significance of individual covariates, and the contribution of individual covariates to model fit as indicated by the t-value. Model assumptions were checked with both distribution analyses of the raw data and the residuals of the model. Model residuals were normally distributed, supporting use of linear mixed effects repeated measures models without transformation of the data. Only variables included in the best fit models are presented.

Three sets of Pearson zero-order correlation coefficients (among outcome variables, among covariates, and between covariates and outcomes) were performed for each year of the study to examine possible changes in variable relationships over absolute time. Covariates that correlated with outcomes with a $p \leq 0.05$ and had an $r \geq 0.10$ for at least 1 year of the study were included in the preliminary full models; all covariates met these criteria. Correlation coefficients among covariates with an $r \geq 0.90$ were considered to have potentially high levels of collinearity. Further analyses of collinearity among covariates and interaction terms were conducted as part of model construction. All variables included in the best
fit models had a variance inflation index < 5 or a condition index < 30 and thus had low levels of collinearity. Patient covariates and outcomes were treated as the between-subject effects and calendar year as the within-subject effect. The restricted maximum likelihood (REML) estimation method was used for all iterations. Alternative covariance structures and degrees of freedom methods were evaluated using Akaike’s index criterion (AIC), the restricted likelihood ratio test, and the Bayes index criterion (BIC). All best fit models used the between-within degrees of freedom method. The contribution of individual covariates to model fit was evaluated using the p value and t-value for individual covariates and the overall model fit statistics to determine the best fit model. Variables with a p value > 0.25 were retained if they contributed substantially to model fit, which was defined as reducing the −2 residual log likelihood or AIC by more than 10 or the BIC by more than 2. Biologic exposure ratio was retained in the models regardless of p value.

Within-subject effects:

\[ Y_{ij} = \beta_{0i} + \beta_{1i} \text{Time}_{ij} + \beta_{2i} \text{BER}_{ij} + \beta_{3i} (\text{Time} \times \text{BER})_{ij} + \varepsilon_{ij} \]

Between-subject effects:

\[ \beta_{0i} = \gamma_{00} + \gamma_{01} \text{Covariate}_i + U_{0i} \]
\[ \beta_{1i} = \gamma_{10} + \gamma_{11} \text{Covariate}_i + U_{1i} \]
\[ \beta_{2i} = \gamma_{20} + \gamma_{21} \text{Covariate}_i + U_{2i} \]
\[ \beta_{3i} = \gamma_{30} + \gamma_{31} \text{Covariate}_i + U_{3i} \]

\( i \) 1 ... N individuals
\( j \) 1 ... ni observations f or individual i
\( Y_{ij} \) Value of outcome measure at observation j for individual i
\( \text{Time}_{ij} \) Year of study at observation j for individual i
\( \text{BER}_{ij} \) Value of biologic exposure ratio at time j for individual i
\( \beta_{0i} \) Subject-specific intercept for individual i
\( \beta_{1i} \) Subject-specific slope for variable Time for individual i
\( \beta_{2i} \) Subject-specific slope for variable BER for individual i
\( \beta_{3i} \) Subject-specific slope for Time by BER interaction for individual i
\( \gamma_{00} \) The mean intercept across all individuals
\( \gamma_{10} \) The mean slope across all individuals for the variable Time

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Table 1 Covariates evaluated in initial models

| Variable                          | Sample | Scale       | Description                                                      |
|----------------------------------|--------|-------------|------------------------------------------------------------------|
| Female                           | Static | Binary      | 0 = Male
|                                  |        |             | 1 = Female                                                       |
| Education at baseline            | Static | Categorical | 0 = Did not graduate high school
|                                  |        |             | 1 = Graduated high school                                        |
|                                  |        |             | 2 = Attended technical college                                   |
|                                  |        |             | 3 = Graduated technical college                                   |
|                                  |        |             | 4 = Graduated college                                             |
|                                  |        |             | 5 = Attended graduate school                                      |
|                                  |        |             | 6 = Completed graduate education                                  |
| Baseline mHAQ                    | Baseline | Continuous (0–3) | Lower score indicates reduced function impairment |
| Baseline DAS28-CRP               | Baseline | Continuous (0–9.4) | Lower score indicates reduced disease activity |
| Biologic use prior to enrollment | Baseline | Binary      | 0 = No biologic use                                              |
|                                  |        |             | 1 = Biologic use prior to enrollment                             |
| Biologic use at baseline         | Baseline | Binary      | 0 = No biologic use                                              |
|                                  |        |             | 1 = Biologic use at baseline                                     |
| Year of study                    | Annual | Categorical | 2003 to 2015                                                    |
| Age                              | Annual | Continuous  | Age in years                                                     |
| Disease duration                 | Annual | Continuous  | Disease duration in years                                        |
| Methotrexate use                 | Annual | Binary      | 0 = Not taking methotrexate                                       |
|                                  |        |             | 1 = Taking methotrexate                                           |
| Smoking status                   | Annual | Categorical | 0 = Never smoked                                                 |
|                                  |        |             | 1 = Smoked in the past                                           |
|                                  |        |             | 2 = Current smoker                                               |

DAS28-CRP Disease Activity Score-28 with C-reactive protein, mHAQ modified Health Assessment Questionnaire
Overall, 82.3% of patients were female. Mean (SD) age and disease duration at enrollment were 56.3 (14.2) years and 12.7 (11.9) years, respectively. Patients included in the analyses had an average of 4.9 study visits (range, 1–13 visits) during the study period. Most patients (66.4%) enrolled during the first 3 years after registry initiation, between 2003 and 2005. After the first year of the registry, returning patients comprised between 68.9% (in 2012) and 99.8% (in 2009) of the study sample.

Over time, the proportion of patients with any medication use for RA in that year increased for biologics (from 41% in 2003 to 68% in 2015) and methotrexate (from 45 to 57%), and decreased for other nonbiologic DMARDs (from 36 to 25%), steroids (from 32 to 26%), and NSAIDs (from 61 to 41%) (Fig. 1). The biologic exposure ratio was calculated as the ratio of the patient’s total time on biologics, divided by their time in the BRASS registry. Using these calculations, mean (SD) biologic exposure ratio also increased during the study period, with a 1.6-fold increase from 0.41 (0.49) in 2003 to 0.65 (0.42) in 2015.

**Disease activity: DAS28-CRP**

Over time, DAS28-CRP scores consistently decreased as biologic exposure ratios increased (Fig. 2a). In the repeated measures regression model (Table 3), increased biologic exposure ratios were significantly associated with decreased DAS28-CRP score during follow-up ($\beta = -0.647; P < 0.001$). Other statistically significant ($P < 0.05$) predictors of decreased DAS28-CRP score during follow-up were methotrexate use at baseline and study year, with increased effect size in later years of the study. Statistically significant ($P < 0.05$) predictors of increased DAS28-CRP score during follow-up were a higher baseline DAS28-CRP score, any biologic use at baseline or before enrollment, longer disease duration, a higher baseline mHAQ score, or currently smoking.

**Functional status: mHAQ**

Over time, mHAQ scores for function impairment decreased as biologic exposure ratios increased (Fig. 2b). In the repeated measures regression model (Table 3), increased biologic exposure ratios were significantly associated with decreased mHAQ score during follow-up ($\beta = -0.096; P < 0.001$); methotrexate use at baseline also was a statistically significant ($P < 0.05$) predictor of decreased mHAQ score during follow-up. Statistically significant ($P < 0.05$) predictors of increased mHAQ score during follow-up were higher baseline mHAQ score, longer disease duration, or any biologic use at baseline.
RA severity: RAPID3

Over time, RAPID3 scores decreased as biologic exposure ratios increased (Fig. 2c). In the repeated measures regression model (Table 3), increased biologic exposure ratios were significantly associated with decreased RAPID3 score during follow-up ($\beta = -0.724; P < 0.001$, where a biologic was not taken at baseline). Other statistically significant ($P < 0.05$) predictors of decreased RAPID3 score during follow-up were methotrexate use at baseline and completed graduate education. Statistically significant ($P < 0.05$) predictors of increased RAPID3 score during follow-up were a higher baseline mHAQ score, a higher baseline DAS28-CRP score, any biologic use before enrollment, currently smoking, and biologic exposure ratio × biologic use at baseline ($\beta = 0.938; P = 0.024$, for biologic exposure ratio where a biologic was taken at baseline).

Discussion

This study examined the effect of biologic exposure ratio, a measure of cumulative biologic use, on disease severity and patient functional status using longitudinal mixed-effects linear models in a sample of nearly 1400 RA patients in a prospective registry with up to 13 years of follow-up. From 2003 to 2015, as biologic use increased over time, there was a trend towards decreased disease activity (DAS28-CRP), decreased impairment of functional status (mHAQ), and decreased disease severity (RAPID3). A recent analysis of another RA disease registry reported similar results [36]. In that study, progressive decreases in Clinical Disease Activity Index (CDAI) and HAQ scores were observed with progressive increases in biologic use over time. Our study extends those results by showing that other measures of RA disease burden also improved during the period that biologic use increased. Our study also used repeated measures regression models to examine predictors for improved outcomes in RA patients. In

Table 2  BRASS longitudinal sample: demographic and clinical characteristics

| Year | Total patients | New entrants | Returning patients | Mean age (years) | Female | Mean disease duration (years) | DAS28-CRP | mHAQ | RAPID3 |
|------|----------------|--------------|--------------------|------------------|--------|-------------------------------|-----------|------|--------|
| 2003 | 507            | 507          | –                  | 58.0             | 82.1%  | 15.2                          | 466       | 4.20 | 3.05   |
| 2004 | 742            | 355          | 387                | 57.8             | 82.7%  | 15.1                          | 678       | 3.78 | 2.86   |
| 2005 | 680            | 64           | 616                | 58.6             | 83.2%  | 15.9                          | 622       | 3.52 | 2.77   |
| 2006 | 600            | 23           | 577                | 59.2             | 83.5%  | 16.4                          | 554       | 3.45 | 2.79   |
| 2007 | 612            | 56           | 556                | 59.2             | 83.2%  | 17.0                          | 531       | 3.22 | 2.72   |
| 2008 | 648            | 86           | 562                | 59.3             | 82.4%  | 16.9                          | 615       | 3.16 | 2.65   |
| 2009 | 533            | 1            | 532                | 59.7             | 82.6%  | 18.4                          | 505       | 3.09 | 2.72   |
| 2010 | 458            | 49           | 409                | 59.3             | 83.4%  | 18.0                          | 414       | 3.10 | 2.73   |
| 2011 | 367            | 1            | 366                | 60.0             | 83.4%  | 19.0                          | 314       | 3.02 | 2.62   |
| 2012 | 541            | 168          | 373                | 59.4             | 83.9%  | 17.1                          | 452       | 2.79 | 2.61   |
| 2013 | 493            | 28           | 465                | 61.0             | 82.8%  | 17.9                          | 392       | 2.67 | 2.57   |
| 2014 | 443            | 13           | 430                | 61.2             | 83.7%  | 18.0                          | 340       | 2.49 | 2.51   |
| 2015 | a              | 159          | 44                 | 60.1             | 78.6%  | 17.8                          | 21        | 2.56 | 2.67   |

DAS28-CRP Disease Activity Score-28 with C-reactive protein, mHAQ modified Health Assessment Questionnaire, RAPID3 Routine Assessment of Patient Index Data

* In 2015, only data from the first 6 months were available for this analysis

Fig. 1 Medication use by year increased for biologics and methotrexate, and decreased for other nonbiologic DMARDs, steroids, and NSAIDs.

*Nonbiologic DMARDs excludes methotrexate, which is plotted separately. DMARDs disease-modifying antirheumatic drugs, NSAIDs nonsteroidal anti-inflammatory drugs
Disease activity, function impairment, and RA severity decreased over time with increased biologic exposure ratio. 

**a** Disease activity (Disease Activity Score-28 with C-reactive protein [DAS28-CRP]).

**b** Function impairment (modified Health Assessment Questionnaire [mHAQ]).

**c** RA severity (Routine Assessment of Patient Index Data [RAPID3]).

Each data point represents the mean annual value ± standard error. See Table 2 for annual sample sizes.
these models, the biologic exposure ratio was a significant predictor of improved outcomes for RA disease activity, functional status, and disease severity. Previous research has shown that initiating biologic therapy in earlier RA (≤ 3 years) or in later RA (> 3 years) is associated with similar response rates and structural improvement, but patients with earlier RA report greater improvement in disability with biologic therapy compared with patients with later RA [7].

Several other covariates also were significant predictors of disease activity, function impairment, or disease severity. The convergence of disparate covariates across these models indicates that multiple factors in addition to biologic use may influence functional status in RA patients. For instance, methotrexate is a standard treatment for RA that is known to be effective in both biologic and nonbiologic treatment regimens [1]. In this study, any use of methotrexate in a given year was a significant predictor of improved patient outcomes across all models. Conversely, smoking is a known environmental risk factor [37–39], and “currently smoking” was a significant predictor of worsened disease activity and disease severity.

| Effect                                      | DAS28-CRP | mHAQ | RAPID3 |
|---------------------------------------------|-----------|------|--------|
| Intercept                                   | 1.713     | 0.070| 1.508  |
| Biologic exposure ratio                     | –0.647    | –0.096| –0.724 |
| Taking methotrexate (vs not)                | –0.108    | 0.033| –0.176 |
| Smoking status (vs never)                   |           |      |        |
| Current smoker                              | 0.199     | 0.304| 0.293  |
| Smoked in past                              | 0.048     | 0.020| 0.120  |
| Education (vs did not graduate high school) |           |      |        |
| Completed graduate education                | –0.184    | –0.042| –0.546 |
| Attended graduate school                    | –0.120    | 0.024| 0.584  |
| Graduated college                           | –0.118    | 0.031| 0.373  |
| Graduated technical college                 | 0.013     | 0.004| 0.925  |
| Attended technical college                  | –0.011    | 0.024| 0.505  |
| Current smoker                              | –0.083    | 0.031| –0.305 |
| Smoking status (vs never)                   |           |      |        |
| mHAQ at baseline                            | 0.157     | 0.748| 2.317  |
| DAS28-CRP at baseline                       | 0.578     | 0.748| 0.191  |
| Biologic use (vs no) before enrollment       | 0.378     |      | 0.393  |
| Biologic use (vs no) at baseline            | 0.598     |      | 0.204  |
| Disease duration                            | 0.008     |      |        |
| Study year (vs 2003)                        |           |      |        |
| 2004                                        | –0.283    | <0.001|        |
| 2005                                        | –0.551    | <0.001|        |
| 2006                                        | –0.552    | <0.001|        |
| 2007                                        | –0.765    | <0.001|        |
| 2008                                        | –0.718    | <0.001|        |
| 2009                                        | –0.868    | <0.001|        |
| 2010                                        | –0.814    | <0.001|        |
| 2011                                        | –0.869    | <0.001|        |
| 2012                                        | –0.976    | <0.001|        |
| 2013                                        | –1.146    | <0.001|        |
| 2014                                        | –1.305    | <0.001|        |
| 2015                                        | –1.086    | <0.001|        |

Variables not included in best fit models for all outcomes were not populated for that outcome

– not included in regression model
DAS28-CRP Disease Activity Score-28 with C-reactive protein, mHAQ modified Health Assessment Questionnaire, RAPID3 Routine Assessment of Patient Index Data
Previous use of biologic agents, which is likely to identify patients with severe disease, as well as measures of function impairment and disease activity, were also found in patients with worse function impairment and disease activity over the course of the study.

Calendar year was a significant predictor of disease activity in the model for DAS28-CRP. Over the study period, the proportion of patients in a given year who used any biologic increased, with the greatest increase between 2003 and 2009. From 2010 to 2015, more than half of the patients received a biologic in any given year. Mean scores for DAS28-CRP decreased from 4.2 in the first year of the registry to below 2.6 (the cutoff for RA remission) only in the last 2 years of the analysis. In addition to possible dropouts of patients with higher disease activity, factors such as length of treatment and use of more biologics may have contributed to this finding. The relationship may be partially related to the increasing availability and use of biologic agents throughout the study period. As the BRASS registry does not include data on timing and duration of biologic use prior to enrollment, this analysis could not fully control for the duration of a patient’s lifetime biologic use or the potential effect of biologic use early versus late in disease progression. Although disease duration for the full sample was between 15 and 20 years throughout the course of the study, new enrollees in each annual population tended to have a slightly lower disease duration compared with returning patients. Further study into the long-term relationship between radiographic damage and biologic use is warranted, especially where the timing of biologic use in disease progression is known.

Comorbidities were not examined as possible predictors of disease activity in this analysis. Previous registry analyses have shown that the presence and number of comorbid conditions in RA are associated with lower chance of achieving remission [23, 40]. Assessment of outcomes of biologic therapy in real-world practice remains challenging due to the varied trajectory of disease across patients and the lack of a long-term control group, because patients whose disease progresses beyond nonbiologic DMARD management typically receive biologics. The BRASS database represents a rich source of data that includes 2700 variables composed of demographics, patient reported outcomes, and clinical data. This type of longitudinal data is uncommon and provides insight into the natural course of RA and its treatment over time. The registry sample, which primarily comprised middle-aged, Caucasian women with long-standing RA, was largely representative of the RA population and included both biologic-experienced and biologic-naïve patients.

As with all analyses, there were some notable limitations in this study, as follows. Mean disease duration at study entry was 12.7 years; thus, a substantial portion of active disease was not in the registry, especially during early disease when changes may be more rapid compared with established disease. The registry does not include data on disease severity at the time of biologic initiation or duration of biologic initiation prior to enrollment. Additional data on disease severity at the time of biologic initiation or duration of biologic initiation prior to enrollment in the registry may allow for additional relationships between biologic exposure and patient outcomes to be elucidated. The BRASS database only includes data from patients treated at the Robert Breck Brigham Arthritis Clinic. Previous research suggests that patients in the BRASS database who are less educated and have more severe disease activity are more likely to drop out of the registry [41]. Improvement of patient outcomes over time in this analysis may be explained, at least partially, by greater dropout for patients with more severe disease activity than those with less severe disease activity. The registry also lacks pharmacy data to confirm patient reports of drug utilization. Each of these may limit the external validity and generalizability of study findings to other populations. Multivariable analyses of clinical outcomes were conducted but there is a large potential for confounding from other factors not included in the study. Biologic adherence cannot be assessed from the BRASS registry database.

Biologic exposure may be affected by stopping and starting biologic therapy for many reasons, such as tolerability (i.e., infection), efficacy, insurance coverage, and patient preference. Discontinuation of biologic therapy for efficacy may include not only discontinuation for lack of efficacy but also for symptomatic response; patients who discontinue therapy when they feel better may need to restart biologic therapy when disease worsens again. Patient education may address these situations to improve patient adherence to prescribed biologic therapy.

In conclusion, greater cumulative exposure to biologics was significantly associated with improvements in disease activity, function impairment, and RA severity. Several other covariates also were significant predictors, indicating that various clinical and patient characteristics may contribute to the trajectory of RA. The findings from this study suggest that cumulative exposure to biologics may help to limit the effects of inflammation on patient outcomes over time. A causative effect of early versus late use of biologics was not directly assessed in this analysis. Future studies should examine whether earlier use of biologics is associated with improved long-term outcomes. Additional study of the potential effect of the timing of biologic treatment, including both duration of exposure and disease status at initiation, is needed to support the optimal use of biologics in the treatment of RA.
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Compliance with ethical standards

Conflict of interest At the time of the analysis, B.S.S., D.C., and A.M. were employees and stockholders of Amgen Inc. N.M.G. and B.L.B. were employees of Health Analytics, LLC, which received funding from Amgen Inc. to conduct the study. N.A.S. has been a consultant to Bristol-Myers Squibb and has received research grants from Amgen, Mallinckrodt, UCB, Crescendo Biosciences, Sanofi, Bristol-Myers Squibb, and DiTerity. M.E.W. has been a consultant to Amgen, Bristol-Myers Squibb, Crescendo Bioscience, and UCB and has received research grants from Amgen, Bristol-Myers Squibb, Crescendo Bioscience, UCB, DiTerity, and Sanofi/Regeneron. M.L.F., C.I., and J.C declare that they have no competing interests.

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