Mortality in patients with interstitial lung disease treated with rituximab or TNFi as a first biologic

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ABSTRACT

Objectives Guidelines cautioned prescribing of tumour necrosis factor inhibitors (TNFi) to patients with rheumatoid arthritis and interstitial lung disease (RA-ILD) after reports of new or worsening of ILD. Less is known about outcomes among patients with RA-ILD who receive rituximab (RTX). This study compares mortality in patients with RA-ILD who received RTX or TNFi as their first biologic.

Methods Participants with RA-ILD recruited to the British Society for Rheumatology Biologics Register for RA were included. Death rates were calculated and risk comparisons were made using Cox regression. Causes of death, including the frequency in which ILD was recorded on death certificates were examined.

Results 43 patients on RTX and 309 on TNFi were included. RTX recipients had shorter disease duration and less disability. Death rates were 94.8 (95%CI: 74.4 to 118.7) and 53.0 (22.9 to 104.6) per 1000 person years, respectively. The adjusted mortality risk was halved in the RTX cohort, but the difference was not statistically significant (HR 0.53, 95%CI: 0.26 to 1.10). ILD was the underlying cause of death in 1 of 7 RTX deaths (14%) and 12 of 76 TNFi deaths (16%).

Conclusions Patients with RA-ILD who received RTX had lower mortality rates compared to TNFi. The absence of information on ILD severity or subtype prevents conclusions of which drug represents the best choice in patients with RA-ILD and active arthritis.

BACKGROUND

Pulmonary involvement, including interstitial lung disease (ILD), is common in patients with rheumatoid arthritis (RA).1 2 Such extra-articular manifestations are widely recognised to be associated with increased mortality.1–4 Despite overall mortality falling in recent decades in patients with RA, RA-ILD mortality rates appear to be increasing.5–15

The treatment of active arthritis in patients with RA-ILD is challenging due to concerns about exacerbation of lung disease with certain conventional synthetic disease-modifying antirheumatic drugs (csDMARDs)6 and tumour necrosis factor inhibitors (TNFi).7

Indeed, the British Society for Rheumatology (BSR) has specifically cautioned prescribing TNFi to patients with RA-ILD.3 Rituximab (RTX), which has been successfully used in the treatment of refractory ILD,8 may be considered as an alternative biologic in these patients.9 However, ILD has been reported as a complication of treatment in lymphoma patients treated with RTX.10 11 and less is known regarding the outcomes and mortality risk in patients with RA-ILD who receive RTX.12–14

Previous analyses within the British Society for Rheumatology Biologics Register for RA (BSRBR-RA)15 reported no significant difference in mortality rates between patients with RA-ILD treated with TNFi or csDMARDs. However, a greater proportion of deaths were attributed to RA-ILD in the TNFi cohort. This

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ORIGINAL ARTICLE
The current study aims to investigate 5-year mortality rates and causes of death among patients with RA-ILD treated with either RTX or TNFi as their first-line biologic therapy.

METHODS
The BSRBR-RA is a national, prospective, observational cohort study which has recruited patients with RA commencing a biologic therapy in order to examine the long-term safety of these medications. Patients starting a TNFi (infliximab (Remicade), etanercept (Enbrel) or adalimumab (Humira)) were recruited between 2001 and 2007 and again from 2010, at which point the register opened again to recruit patients starting these three original TNFi as well as certolizumab pegol (Cimzia). Patients starting RTX were actively recruited between beginning of 2008 and 30 September 2011, during which patients starting RTX as either a first-line or second-line drug were registered.

Participants eligible for this analysis had physician’s diagnosed RA-ILD (‘Has the patient ever had pulmonary fibrosis?’) and were recruited prior to 30 September 2011 (to allow a full 5 years of follow-up at the point of data analyses), at the point of commencing either RTX (Mabthera) or a TNFi: infliximab (Remicade), etanercept (Enbrel) or adalimumab (Humira) as their first biologic therapy and had returned at least one follow-up form.

Data
Baseline data collected included sex, age, ethnicity, smoking history, disease duration, disease activity, measured using the 28-joint count disease activity score, and past csDMARDs, corticosteroid use and comorbidities including presence of asthma and chronic obstructive pulmonary disease (COPD) and disability (Stanford Health Assessment Questionnaire (HAQ)). The primary outcome measure for this analysis was death, which was captured by: (1) rheumatologist follow-up questionnaire (returned 6 or 12 month follow-up questionnaire depending on time since treatment commenced), (2) family report to the register and (3) ‘flagging’ with the UK Office for National Statistics, which provided a copy of the death certificate, including cause of death. Reporting of death, including cause of death, is a mandatory requirement for all deaths occurring in the UK. Secondary outcomes included (1) the proportion of deaths which listed RA-ILD as the underlying cause of death and (2) the proportion of deaths which listed RA-ILD anywhere on the death certificate. RA-ILD was identified using International Classification of Diseases-10 codes J84.1 ‘interstitial pulmonary disease with fibrosis’ and J84.9 ‘interstitial pulmonary disease, unspecified’ and M05.1 ‘rheumatoid lung’.

Statistical analysis
Baseline characteristics were compared between the cohorts, using Pearson $\chi^2$ tests and Student’s $t$-tests. Years of follow-up time were calculated from the date of starting biologics until death, or 5 years following first registration, whichever came first. Mortality rates, per 1000 person years (pyrs) with 95% CI were calculated using an ever-exposed model, assuming a Poisson distribution of cases. Kaplan-Meier survival curves for mortality were generated. The risk of death between cohorts was compared using Cox regression. Confounders taken forward to a fully adjusted model included the a priori selected variables age, sex, and disease duration. Further any additional factors which shifted the unadjusted risk of mortality between RTX and TNFi by more than 10% were included as confounders. Additional candidate confounders included ethnicity, smoking history, disability (measured using the HAQ), disease activity (measured using the DAS28 (disease activity score-28)) previous DMARD and current steroid use, number of comorbidities and diagnoses of asthma and COPD. Missing data were accounted for in variables of interest using multiple imputation, with 20 imputed datasets. The variables on which multiple imputation was conducted and the proportion of missing data are shown in table 1. Variables included in the imputation model included those offered as candidate confounders, the natural log of follow-up time and death.

All analyses were conducted using Stata V.13.1 (StataCorp).

RESULTS
Of 1632 recruited to the RTX and 15644 participants recruited to the TNFi cohort by 30 September 2011, 352 patients (43 RTX (6.8%) and 309 TNFi (2.0%)) satisfied criteria for inclusion in the analysis.

Age and gender were comparable in the two cohorts (table 1). The mean disease activity using the 28-joint count disease activity score (DAS28), of those commencing RTX was lower than that of those commencing TNFi (mean: 6.3 (SD:1.1) vs 6.6 (1.0)). Those commencing RTX had significantly shorter median disease duration (median: 5.5 (IQR 3–13) vs 12 years (7–20)), had, on average, used fewer previous csDMARDs (3.2 (1.2) vs 4.0 (1.6)) and had less disability (HAQ: 1.6 (0.7) vs 2.1 (0.5)) than patients starting TNFi.

All patients commencing RTX did so after 2008 and all patients commencing TNFi did so prior to 2008 and thus there was no overlap in patient recruitment. Finally, the prevalence of pulmonary comorbidities was significantly greater in the RTX cohort (41.9%) compared with the TNFi cohort (25.6%, p<0.03), driven by the presence of COPD (RTX: 33.3% (n=14) vs TNFi: 18.8% (n=58), p<0.03).

During 801.3 pyrs, 76 deaths occurred in the TNFi cohort and 8 deaths occurred within 150.7 pyrs in the RTX cohort. The respective all-cause mortality rates per 1000 pyrs were 94.8 (74.7–118.7) and 53.0 (22.9–104.6). The unadjusted 5-year risk of mortality in the RTX-treated patients was approximately half that in the TNFi-treated patients, although this was not statistically
### Table 1  Baseline characteristics of the cohorts

|                  | RTX (n=43)       | All TNFi (n=309) | p Value |
|------------------|------------------|-----------------|---------|
| Age (years)      | 64.7 (11.4)      | 62.5 (10.4)     | 0.2     |
| Female, n (%)    | 19 (44.2)        | 179 (57.9)      | 0.09    |
| Smoking history, n (%) |               |                 |         |
| Current          | 7 (16.3)         | 64 (20.7)       | 0.6     |
| Former           | 27 (62.8)        | 169 (54.7)      |         |
| Never            | 9 (20.9)         | 75 (24.6)       |         |
| Ethnicity        |                  |                 |         |
| White            | 37 (86.0)        | 271 (87.7)      | 0.97    |
| Other            | 1 (2.4)          | 7 (2.3)         |         |
| Missing, n (%)   | 5 (11.6)         | 31 (10.3)       |         |
| DAS28            | 6.3 (1.1)        | 6.6 (1.0)       | 0.03    |
| HAQ              | 1.6 (0.7)        | 2.1 (0.54)      | <0.001  |
| Disease duration (years), median (IQR)* | 5 (3–13) | 12 (7–20)      | <0.001  |
| Baseline steroid use, n (%) | 27 (62.8) | 177 (57.3)     | 0.5     |
| Number of prior csDMARDs | 3.2 (1.2) | 4.0 (1.6)      | <0.01   |
| Comorbidity, n (%)† |               |                 |         |
| None             | 10 (23.3)        | 103 (33.3)      | 0.3     |
| 1 comorbidity    | 14 (32.6)        | 113 (36.7)      |         |
| 2 comorbidity    | 13 (30.2)        | 64 (20.7)       |         |
| ≥3 comorbidity   | 6 (13.9)         | 29 (9.4)        |         |
| Hypertension     | 18 (42.9)        | 123 (39.9)      | 0.7     |
| Ischaemic heart disease (angina + MI) | 9 (20.9) | 39 (12.6)     | 0.1     |
| Stroke           | 5 (11.9)         | 16 (4.9)        | 0.09    |
| Renal            | 2 (4.6)          | 11 (3.6)        | 0.7     |
| Diabetes         | 4 (9.3)          | 17 (5.6)        | 0.3     |
| Liver disease    | 1 (2.3)          | 12 (3.9)        | 0.7     |
| Depression       | 8 (18.6)         | 49 (16.4)       | 0.6     |
| Lung (asthma + COPD) | 18 (41.9) | 80 (25.6)     | 0.03    |
| Asthma           | 5 (11.9)         | 32 (10.4)       | 0.8     |
| COPD             | 14 (33.3)        | 58 (18.8)       | 0.03    |
| Year of registration‡ |            |                 |         |
| <2008            | 0 (0)            | 309 (100)       | NA      |
| ≥2008            | 43 (100)         | 0 (0)           |         |

Values are mean (SD), unless otherwise stated; categorical variables compared using Pearson chi-squared tests and continuous variables compared using Student’s t-tests; information about missing data only reported for variables in which missingness occurred.

*Comparison conducted using Wilcoxon rank-sum test.

†Hypertension, coronary heart disease (MI or angina), stroke, lung (asthma or bronchitis/emphysema), diabetes mellitus, depression, renal disease and liver disease.

‡Due to differences in recruitment periods for cohorts, between group comparisons are not appropriate for year of registration.

COPD, chronic obstructive pulmonary disease; csDMARDs, conventional synthetic disease modifying antirheumatic drug; DAS28, disease activity score-28; HAQ, Health Assessment Questionnaire; MI, myocardial infarction; pyrs, person years; NA, not applicable; RTX, rituximab; TNFi, tumour necrosis factor inhibitor.
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Figure 1 Kaplan-Meier survival curves for death following exposure to TNFi or RTX over the first 5 years following therapy commencement, within an intention to treat analysis. Numbers in table represent the number of patients at risk of death at the specific follow-up time points. RTX, rituximab; TNFi, tumour necrosis factor inhibitor.

In total, 12 (16%) 12 (16%) TNFi deaths and 1 (14%) RTX deaths had RA-ILD listed as the underlying cause of death (table 2). However, RA-ILD was more commonly reported anywhere on the death certificates of the RTX cohort. Death certificates were unavailable for two patients (one in each cohort) who died outside the UK.

In summary, this study provides some evidence to suggest that patients with RA with ILD who received RTX as their first biologic for RA appeared to have better long-term survival compared with patients who received TNFi. This finding persisted after adjusting for key markers of RA disease severity. The proportion of deaths attributed to RA-ILD did not differ between the two groups.

This study has a number of limitations. First, these are observational data and the impact of non-randomisation to treatment is confounded by the fact that the two cohorts of patients included in this analysis were recruited over different time frames, where the approach to diagnosis and management of RA, and therefore possibly mortality, have differed. Although differences in disease severity and duration were identified and controlled for in our analysis, the early period of recruitment for TNFi also included a time prior to any guidelines regarding biologic treatment and RA-ILD, while later periods of recruitment may span a time when, in response to guidelines, patients with the most severe ILD may not have been prescribed a biologic at all. In part, these differential periods of recruitment were driven by the overall design of the BSRBR-RA, but there were 2 years of recruitment overlap during which no patients recruited to the register starting a TNFi had a history of RA-ILD.

Second, although this represents one of the largest outcome studies of RA-ILD in biologic-treated patients, including comprehensive drug exposure data and outcome data, the sample size, especially of patients exposed to RTX, was low, which may have affected power and also our ability to adjust further for other potential confounders using propensity scores.

Though different from a typical cohort of patients with RA starting a biologic, those included in this analysis were reflective of patients with RA-ILD, increasing the external validity of these findings. Unfortunately, data regarding the subtype or severity of the underlying RA-ILD, which have been associated with differences in mortality, were not available. It is recognised that prognosis is worse among patients with RA with usual interstitial pneumonia compared with those with non-specific interstitial pneumonia\(^{14,20}\) and that one of the strongest predictors of mortality is pulmonary function.\(^{21}\) Therefore, the data presented cannot be used to make firm conclusions regarding relative safety of RTX and TNFi. Similarly, smoking data were not recorded over time and it would be beneficial to replicate this analysis in a cohort which included longitudinal data on ILD severity and smoking data.

Although it was not possible to establish whether the progression of the pulmonary disease was altered by the biologic agent, reassuringly, most of the deaths were not attributed to ILD. In previous analyses, a higher proportion of deaths were attributed to RA-ILD in patients commencing TNFi compared with those using sDMARDs.\(^{15}\) Here, RA-ILD was identified as the underlying cause of death in 1 of 7 RTX deaths (14%) and 12 of 76 TNFi deaths (16%). Nevertheless, RA-ILD was listed more often on the death certificates of patients in the RTX cohort. This may reflect that RA-ILD may have been a prominent feature that informed biologic choice among patients starting RTX.

In summary, this study provides some evidence to suggest that patients with RA with ILD who received RTX as their first biologic experienced better long-term survival compared with patients starting TNFi. However, the high proportion of death certificates on which ILD was present in the RTX cohort, the lack of information on severity or subtype of ILD and differences in time of recruitment precludes firm conclusions about comparative drug safety in this population.

| Cohort | Registration | 1 year | 2 years | 3 years | 4 years | 5 years |
|--------|-------------|--------|---------|---------|---------|---------|
| TNFi   | 309         | 269    | 252     | 242     | 233     |         |
| RTX    | 43          | 42     | 39      | 35      | 35      |         |

Discussion

Patients with physician-recorded RA-ILD who received RTX as their first biologic for RA appeared to have better long-term survival compared with patients who received TNFi. This finding persisted after adjusting for key markers of RA disease severity. The proportion of deaths attributed to RA-ILD did not differ between the two groups.

This study has a number of limitations. First, these are observational data and the impact of non-randomisation to treatment is confounded by the fact that the two cohorts of patients included in this analysis were recruited over different time frames, where the approach to diagnosis and management of RA, and therefore possibly mortality, have differed. Although differences in disease severity and duration were identified and controlled for in our analysis, the early period of recruitment for TNFi also included a time prior to any guidelines regarding biologic treatment and RA-ILD, while later periods of recruitment may span a time when, in response to guidelines, patients with the most severe ILD may not have been prescribed a biologic at all. In part, these differential periods of recruitment were driven by the overall design of the BSRBR-RA, but there were 2 years of recruitment overlap during which no patients recruited to the register starting a TNFi had a history of RA-ILD.

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Table 2  Mortality rates and cause of death in patients with ILD treated with first-line TNFi or RTX in an intention to treat analysis

|                          | RTX n=42 | All TNFi n=309 |
|--------------------------|----------|---------------|
| Total follow-up time (pyrs) | 256.0    | 2564.0        |
| Total follow-up time (pyrs) censored at 5-year follow-up | 150.7    | 801.3         |
| Median (IQR) follow-up per person (years) | 6.3 (5.3–7.3) | 9.1 (5.0–12.0) |
| Median (IQR) follow-up per person (years) censored at 5-year follow-up | 3.9 (2.6–4.4) | 2.7 (1.2–4.1) |

Deaths within first 5 years following treatment start

All-cause (n) 8 76
All-cause mortality/1000 pyrs (95% CI) 53.0 (22.9 to 104.6) 94.8 (74.7 to 118.7)

Cox regression models of relationship between treatment group and mortality*

|                          | Unadjusted model | Adjusted model† |
|--------------------------|------------------|-----------------|
| Treatment TNFi           | HR 1.0           |      1.0        |
| Treatment RTX            | 0.53             | 0.26–1.10       |

Cause of death

Deaths (n) 8 76
ILD certificate information not yet reported to register (n) 1‡ 1‡
ILD present on a death certificate (%)§ 5 (71.4) 27 (36.5)
ILD listed as the underlying cause of death (%)§ 1 (14.3) 12 (16.0)
ILD present in section I on death certificate, n(%)§ 2 (40.0) 20 (27.0)
ILD on a death certificate I(a) (%)§ 1 (20.0) 5 (6.8)
ILD on a death certificate I(b) (%)§ 0 14 (18.9)
ILD on a death certificate I(c) (%)§ 1 (20.0) 1 (1.3)
ILD on a death certificate II (%)§ 3 (42.8) 7 (9.5)

*Analysis conducted using multiple imputation for missing data.
†Age, disability (HAQ), disease activity (DAS28), disease duration and sex adjusted.
‡Death occurred outside of the UK and therefore no death certificate and the cause of death cannot be determined.
§Percentage of those with death certificate data available.
DAS28, disease activity score-28; ILD, interstitial lung disease; HAQ, Health Assessment Questionnaire; pyrs, person years; RTX, rituximab; TNFi, tumour necrosis factor inhibitors.

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Competing interests  None declared.

Patient consent  Detail has been removed from this case description/these case descriptions to ensure anonymity. The editors and reviewers have seen the detailed information available and are satisfied that the information backs up the case the authors are making.

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