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Clinical impact of Ocrelizumab extended interval dosing during the COVID-19 pandemic and associations with CD19⁺ B-cell repopulation

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Summary
In this retrospective clinical audit, 136 patients with MS received Ocrelizumab during the COVID-19 pandemic. There was no significant difference in clinical relapse rate or radiological activity between 67 patients who received extended interval dosing (EID) (≥30 weeks, mean 48.3 ± 5.9 weeks) compared to standard interval dosing (<30 weeks) with average follow-up of over 4 months. CD19⁺ B-cell repopulation occurred in 94% (p<0.001) of EID patients at re-infusion and correlated strongly with re-dosing interval (rs=0.738, p<0.0001) but was not associated with inflammatory disease activity. EID did not impact short-term disease activity, despite significant CD19⁺ B-cell repopulation and warrants long-term prospective study.

1. Introduction

The COVID-19 pandemic has significantly impacted the healthcare of many people with multiple sclerosis (PwMS) causing cancelled or delayed appointments, investigations and disease modifying therapy (DMT) (Vogel et al., 2020).

Anti-CD20 therapies were posited to increase risk of hospitalisation or intensive care admission with COVID-19 infection (Simpson-Yap et al., 2021). The Association of British Neurologists suggested delaying Ocrelizumab re-infusion during highly infectious periods using CD19⁺ B-cell repopulation (1% of total lymphocyte population) (Coles et al., 2020). From 15th July 2020, PwMS on Ocrelizumab at the Royal Stoke MS centre were routinely offered blood sampling for CD19⁺ B-cell population of total lymphocytes (CD19⁺ %). With increasing use of telemedicine and hospital avoidance behaviours (Portaccio et al., 2021), practically CD19⁺ % was obtained when patients attended hospital for infusions.

In this retrospective audit, we reviewed the clinical impact of Ocrelizumab extended interval dosing (EID) resulting from the COVID-19 pandemic at our tertiary neuroscience centre and explore associations of CD19⁺ % with EID, clinical relapses and radiological activity.

2. Methods

We retrospectively reviewed electronic records of all PwMS on Ocrelizumab and identified patients who received standard interval dosing (SID) (<30 weeks) versus EID (≥30 weeks) during the COVID-19 pandemic.

Inclusion criteria were prior completion of first Ocrelizumab cycle (two 300 mg infusions 14 days apart), subsequent 600 mg infusion during the COVID-19 pandemic and CD19⁺ % sampling. Data was extracted for age, sex, MS phenotype, disease duration, Expanded Disability Status Scale (EDSS) score, previous ocrelizumab cycles and dates of previous and subsequent ocrelizumab cycles. We reviewed clinical notes for relapses and MRI reports since previous ocrelizumab cycles and dates of previous and subsequent ocrelizumab cycles. We reviewed clinical notes for relapses and MRI reports since previous ocrelizumab cycles and dates of previous and subsequent ocrelizumab cycles. We reviewed clinical notes for relapses and MRI reports since previous ocrelizumab cycles and dates of previous and subsequent ocrelizumab cycles.
CD19+ % (>1%) or re-dosing interval (as independent variable) could predict binary (yes/no) outcomes for clinical relapses or radiological activity (dependant variables) with demographic variables as covariates.

3. Results

From 152 patients who previously received a full cycle of Ocrelizumab, 16 patients were excluded; 4 patients who did not have a subsequent infusion and 12 patients who did not have CD19+ % sampling. Of 136 patients included, 69 patients had SID and 67 patients had EID (see Table 1).

EID and SID patients did not differ significantly in terms of age, gender, disease duration, clinical phenotype, DMT use or baseline EDSS. However, EID appeared more common in patients with one prior Ocrelizumab infusion whilst patients with 3 or more prior infusions had no delays (p<0.001). The mean re-dosing interval was 48.3 weeks for EID versus 25.4 weeks for SID (mean difference 22.9 weeks). Retrospective follow-up was 19.1 weeks after EID and 20.8 weeks after SID. There were no significant differences in clinical relapses or radiological activity between EID and SID groups.

EID patients had significantly higher CD19+ % (p<0.001) and CD19+ B-cell counts (p<0.001), with 94% of EID patients CD19+ replete (>1%) by next ocrelizumab cycle compared with 17% of SID patients (p<0.001). CD19+ % correlated strongly with re-dosing interval (r=0.738, p<0.0001) but was not associated with clinical relapses or radiological activity (see Fig. 1).

Binary logistic regression analyses found neither re-dosing interval nor CD19+ % could predict clinical relapses of radiological activity.

4. Discussion

In this retrospective study, 48.3 week mean EID showed no effect on inflammatory disease activity over 19.1 weeks follow-up, despite CD19+ repopulation in almost (94%) EID patients.

Our real-world data mirrors results from the Ocrelizumab phase II extension trial, where annualised relapse rates and 6-month disability progression remained low during 12–18 months treatment-free with fewer adverse effects and infections (Baker et al., 2020). The small number of patients imaged had no radiological activity. CD19+ B-cells took median 15 months to return to baseline or lower limit of normal. Although, we did not have baseline CD19+ counts, 39% of EID patients had CD19+ counts above the lower limit of normal (100 cells/μL).

Similar observations of EID have been reported during the COVID-19 pandemic, albeit with shorter EID intervals and follow-up duration. Personalised Ocrelizumab dosing using CD19+ B-cell counts led to EID in most patients, with low disease activity (van Lierop et al., 2021). Recently, a small retrospective analysis of 33 patients, showed no clinical consequences from mean EID of 33 weeks, although lacked radiological data (Barun et al., 2021). Moreover, a retrospective multicentre cohort study showed stable disease activity 3 months following EID in 116 patients with median delay of nearly 9 weeks. CD19+ B-cell counts were similarly not associated with re-emergent disease activity, although the majority of EID patients remained CD19+ deplete (Rolfe et al., 2021).

Tailoring EID to memory B-cell (CD19+ CD27+ co-expression)
repopulation may be better, as early CD19+ repopulation produces immature and immune naïve B-cells from the bone marrow rather than pathogenic memory B-cells (Baker et al., 2020). This has been shown to reduce reinfusion requirements, whilst maintaining reduction of disease activity in PwMS on Rituximab. (Novi et al., 2020) EID to memory B-cells may also allow improved vaccine responses through naïve B-cell repopulation (Baker et al., 2020).

This study has limitations. Relatively low relapse rates in both groups suggest treatment efficacy, but limit statistical power to detect differences between dosing groups. Radiological activity may be missed as patients were not scanned regularly and not all patients were imaged, although >60% of our patients had MRI scans. Similarly, disability progression was not measured, as EDSS follow-up data was not acquired during the pandemic. We also cannot determine the longer-term consequences of EID, either from a single delayed dose or continued EID regimen.

In conclusion, our real-world observational data suggests that Ocrelizumab EID does not impact short-term disease activity, despite CD19+ B-cell repopulation and offers practical reassurance if delaying Ocrelizumab re-dosing to mitigate risk of adverse events. However, prospective studies of continued EID versus SID are necessary to address long-term efficacy, safety, and cost-effectiveness, as well as to identify optimum dosage interval and monitoring.

Declaration of Competing Interest

None.

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