Evaluation of a patient self-stratification methodology to identify those in need of shielding during COVID-19

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The logistical challenges of rapidly and accurately identifying those patients who needed to shield during the COVID-19 pandemic were unprecedented. We report our experiences of meeting this challenge for >9,000 patients with rheumatic and musculoskeletal disease at our centre, incorporating an element of guided patient self-stratification. Our results indicate that patients are able to stratify their own risk accurately using the BSR COVID-19 risk stratification guidance.

KEYWORDS: COVID-19, rheumatic and musculoskeletal disease, service evaluation, British Society for Rheumatology

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Introduction

Coronavirus 2019 (COVID-19) has presented a unique constellation of clinical, logistical and ethical challenges for rheumatology. There have been concerns that patients with rheumatic and musculoskeletal disease (RMD) receiving conventional synthetic (cs-), biological (b-) or targeted synthetic (ts-) disease modifying anti-rheumatic drugs (DMARDs) may have an increased risk of COVID-19, based upon previous experience in this population with infectious diseases.1,2 Furthermore, patients with multisystem diseases such as connective tissue diseases (CTDs) and vasculitis may be at particular risk in view of their potential co-existent respiratory or renal compromise.3–5 The need to urgently identify those at highest risk and provide them with guidance was supported by the publication of guidance and risk assessment tools by NHS England and the British Society for Rheumatology (BSR).6,7 These documents provided a framework to approach risk assessment and stratification, with those designated ‘high risk’ given advice on shielding. While early signals from Wuhan, China, and Italy had so far appeared to demonstrate reassuringly low numbers of cases of COVID-19 in those with immunodeficiency or on immunosuppressants, BSR produced clear guidance that these factors should be considered relevant in assessing patients’ risks of COVID-19 until further, unequivocal evidence demonstrated otherwise, a view held by many clinicians, and hence their stratification process was widely adopted.

Methods

The logistical challenges of undertaking this risk stratification rapidly in a large population were unprecedented.8 We report our experiences of meeting this challenge for >9,000 RMD patients managed at our centre, based upon the BSR guidance for identifying those at high risk of COVID-19 and in need of shielding. Our principal aim was to deliver accurate and timely risk stratification for COVID-19 precautionary measures for a large cohort of patients.

In order to complete this task quickly and efficiently, we took a stepwise approach:

> Undertake individualised assessment of those on bDMARDs and advise them of their personalised risk and related guidance.
> Contact patients with CTD and vasculitis, advising them to shield, accepting some low-moderate risk patients would be included.
> Contact patients with CTD-associated pulmonary arterial hypertension (PAH) and/or interstitial lung disease (ILD) with advice to shield.
> Devise a risk stratification tool to be sent out to all remaining RMD patients enabling them to score their own risk level accurately.

We initially prioritised assessment of those on biologics, identified through our biologics database, and their risk was scored by departmental clinicians. CTD and vasculitis cases were identified through electronic patient record (EPR) text mining and clinic lists, and shielding letters were sent accordingly.9 A total of 1,622 patients on bDMARDS and tsDMARDS and 474 patients with CTD/vasculitis were sent letters with their individualised risk stratification by 7 April 2020.

In order to reach other potentially at-risk patients from our cohort with RMD, a further 7,517 patients with RMD registered
under our care within the last 2 years were identified through EPR text mining of clinical correspondence. They were sent comprehensive information guiding them through a process of self-stratification based on BSR guidance. There were two parts to this: a paper scoring method (supplementary material S1) with instructions for the patient to work through and a link to an internet platform (supplementary material S2). Our online platform guided patients through the same risk stratification matrix to identify their risk group and also prompted patients to enter the score they had calculated using the paper risk matrix. The latter method enabled us to capture patients’ self-scoring and add them to the central list of shielding patients. To validate the reliability of the patient self-scoring method, 100 consecutive patients were contacted in telephone clinics and asked how they scored themselves. They were then re-scored by a rheumatologist.

Results
Of 100 patients contacted by telephone, 97 had received the letter and stratified their risk. Of these, 89 had estimated their risk correctly and 31 had not assessed themselves using the web portal. The reasons given for not using the web portal were that they felt sufficiently informed already or that it was too complex. Only 5% of this sample were required to shield, offering a basis on which to estimate the numbers needing to shield in our wider RMD population.

Of the 7,517 RMD patients who were sent guidance on self-scoring, 910 (13%) logged onto the web platform to complete the online process over the first 4 weeks. Table 1 shows how patients scored themselves using each method; 72% scored themselves consistently by both methods.

Discussion
We report a systematic approach to contacting our RMD patient cohort with personalised guidance to help them protect themselves during the COVID-19 pandemic, based upon the process produced by BSR. Our stepwise process enabled us to issue prompt guidance for those at the highest risk and adopt a more nuanced assessment for those with other risk factors. Our follow-up telemedicine reviews indicated that most patients felt they had been supplied with sufficient information via letter to safely manage their risk. Those using the web portal represented a small proportion of our patient cohort, with inherent selection bias. Limited patient engagement in this sort of initiative has also been reported by others.

Table 1. Results of patients’ self-scoring using both the paper and web based COVID19 scoring methods

<table>
<thead>
<tr>
<th>Scoring method</th>
<th>High risk (Shielding) n (%)</th>
<th>Moderate risk (Strict social distancing) n (%)</th>
<th>Low risk (Standard social distancing) n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Online risk matrix</td>
<td>99 (11%)</td>
<td>208 (23%)</td>
<td>603 (66%)</td>
</tr>
<tr>
<td>Paper risk matrix</td>
<td>165 (18%)</td>
<td>281 (31%)</td>
<td>464 (51%)</td>
</tr>
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</table>

Conclusion
We report the application of a multi-layered approach comprising individual case note review, rule-based methods for certain diseases (eg PAH and ILD), and postal/web-based patient self-stratification to promptly risk stratify >9,000 RMD patients from a single centre. Our results indicate that patients are able to stratify their own risk accurately using the BSR COVID-19 risk stratification guidance, enabling them to take precautionary measures to modify their risk of contracting COVID-19. While we cannot be sure that our efforts have led to complete coverage of those who need to shield, an analysis of data from the NHS spine has reassured us that, at the time of writing, no patient on bDMARDS or tsDMARDS from our cohort has died of COVID-19.

Supplementary material
Additional supplementary material may be found in the online version of this article at www.rcpjournals.org/clinmedicine:
S1 – Paper patient COVID-19 risk stratification letter
S2 – Online patient COVID-19 risk stratification form

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Conflicts of interest
Elizabeth Reilly reports research grants from Actelion and Celgene, outside of the submitted work. John Pauling reports research grants, personal fees and non-financial support from Actelion Pharmaceuticals. Dr Pauling also reports personal fees from...
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References


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