Correspondence on ‘Efficacy of a tight-control and treat-to-target strategy in axial spondyloarthritis: results of the open-label, pragmatic, cluster-randomised TICOSPA trial’

We read with great interest the article by Molto et al. The authors conducted a pragmatic, cluster trial evaluating the efficacy and cost-effectiveness of a tight-control/treat-to-target (TC/T2T) strategy versus usual care (UC) over 48 weeks in 160 patients with axial spondyloarthritis (axSpA). The results showed that the percentage of patients achieving ≥30% improvement on the Axial SpondyloArthritis International Society-Health Index (the primary outcome) was not statistically significantly different between the two groups (TC/T2T vs UC: 47.3% vs 36.1%, p=0.07). Despite this, 13 prespecified secondary outcomes performed better in the TC/T2T group at 48 weeks (5 of them reached statistical significance), and TC/T2T improved quality of life and reduced health-related cost compared with UC. This is an important trial showing the potential benefits of a TC/T2T strategy in clinical practice among patients with axSpA. However, there are some concerns that would better be clarified.

First, sample size calculation in this trial was based on a two-step method due to its cluster design, as indicated in the article. The designed sample size was 232 patients (116 per group), but only 160 patients (80 per group) were enrolled in the trial. The shortage of sample size was not discussed in the article throughout. While the primary outcome was not statistically significantly different between the two strategies, the authors acknowledged that this trial should not be simply interpreted as a negative trial and provided several good explanations for the lack of statistical significance. However, the main reason is more likely due to the much smaller sample size. Post-hoc conditional power analyses would help inform whether the trial was under-powered due to smaller sample size, had the study completed the preplanned enrolment size. Moreover, the 95% CIs around the point estimates for the primary and secondary outcomes, which would have provided more informative results of the trial, have not been displayed in the article. Therefore, whether the results were truly statistically non-significant and clinically meaningless needs further discussion.

Second, contamination bias is almost unavoidable in pragmatic trials and thus needs to be carefully controlled. For this study, it might be helpful to report the proportion of rheumatologists in each centre who have readily provided healthcare similar to the TC/T2T strategy. Despite centres were randomised to allocate to either treatment group, there were only 18 centres and assumably simple randomisation was used. Indeed, baseline imbalances between the two groups were observed.

Third, for some continuous secondary outcome measures, the authors reported between-group difference at week 48 rather than change from week 0 to week 48. It has been suggested in a clinical trial that analysis of change score with adjustment for baseline values of the outcome measure may be more appropriate, especially when there is a high correlation between baseline and follow-up measurements, which is generally the case in rheumatic diseases.

We respect the significant contributions of the authors and look forward to the follow-up results and interpretations of this study.

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