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The relation between cartilage biomarkers (C2C, C1,2C, CS846, and CPII) and the long-term outcome of rheumatoid arthritis patients within the CAMERA trial

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Abstract

Introduction: The aim of this study was to investigate whether serum biomarker levels of C2C, C1,2C, CS846, and CPII can predict the long-term course of disease activity and radiographic progression early in the disease course of rheumatoid arthritis (RA).

Methods: In patients in the CAMERA trial, levels of biomarkers were evaluated at baseline and after 1 year of treatment. Relations of (changes in) biomarker values with the mean yearly radiographic progression rate and mean disease activity over a 5-year period were evaluated by using regression analysis. The added predictive value of biomarkers over established predictors for long-term outcome was analyzed by multiple linear regression analysis.

Results: Of 133 patients, serum samples were available at baseline and after 1 year of treatment. In the regression analysis C1,2C at baseline, the change in C2C, C1,2C, and the sum of the standardized changes in C2C + C1,2C scores were statistically significantly associated with the mean yearly radiographic progression rate; the change in CPII was associated with the mean disease activity over 5 years of treatment. In the multiple linear regression analysis, only the change in C1,2C was of added predictive value ($P = 0.004$) for radiographic progression. Explained variances of models for radiographic progression and disease activity were low (0.28 and 0.34, respectively), and the biomarkers only marginally improved the explained variance.

Conclusions: The change in C1,2C in the first year after onset of RA has a small added predictive value for disease severity over a 5-year period, but the predictive value of this biomarker combined with current predictive factors is too small to be of use for individual patients.

Introduction

Biomarkers are molecules or fragments that are released into biologic fluids during the process of tissue turnover and, for rheumatoid arthritis (RA), are considered to be indicative of degradation or synthesis of cartilage, bone, and synovial tissue [1]. Several serum biomarkers are on the market, including those provided by IBEX (Montreal,

Quebec, Canada); C2C, C1,2C, CS846, and CPII [2-5]. These biomarkers might be good candidates because they directly reflect the bone and cartilage turnover rate in the (affected) joints of patients with RA. The two markers for collagen degradation originate from type II collagen (C2C) and from type I as well as type II collagen (C1,2C), reflecting cartilage and bone degradation. The marker for turnover originated from proteoglycan aggrecan (CS846) and the marker for synthesis of type II procollagen (CPII).

Earlier research with these biomarkers showed no consistent results regarding the predictive value for the

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long-term outcomes in (early) RA. Only six publications described the relation of (one of) these biomarkers with (long-term) radiographic (Table 1) or clinical (Table 2) outcome in RA [6-11]. The relation between these biomarker values and radiographic progression is inconsistent; some studies show a higher value in cases of higher radiographic progression [7,9,11], whereas others show a lower value in cases of higher radiographic progression [8] or show no association at all [7-11]. The same holds true for the relation between these biomarker values and disease activity over time [9].

Because of these conflicting results and the limited available literature on the association between these biomarkers and clinical and radiographic progression, the aim of this study was to investigate whether values of C2C, C1,2C, CS846, and CPII determined early in the disease can predict the long-term radiographic and/or clinical outcome in patients with early RA.

Materials and methods

Patients included in this study were participants in the 2-year randomized open-label prospective multicenter treatment strategy trial (Computer Assisted Management in Early Rheumatoid Arthritis, CAMERA) [12]. In the CAMERA study, patients were randomly assigned to either an intensive tightly controlled MTX-based

treatment strategy based on computer-guided monthly predefined response criteria or to a conventional MTX-based treatment strategy based on regular clinical practice with 3-monthly visits. All patients fulfilled the 1987 revised American College of Rheumatology (ACR) criteria for RA [13]. At study entry, all patients had a disease duration of less than 1 year and were DMARD and glucocorticoid naïve. The medical ethics committees of all participating hospitals approved the study, and all patients gave written informed consent before entering the trial.

From all available patients, serum samples were collected at baseline (before treatment) and 1 year after inclusion into the study. Serum samples were frozen as soon as possible after blood collection and stored at -20°C until analysis (analysis shortly after all 1-year samples were obtained). Because the trial was performed according to general clinical practice as much as possible, sample collection was not restricted to fasting conditions.

Biomarker analyses

For this study, only samples that had not been thawed before were used. For all biomarkers, enzyme-linked immunosorbent assays (ELISAs) were performed according to manufacturer's instructions (IBEX Montreal, Quebec, Canada).

Table 1 Overview of the literature on the (significant) relation between biomarker and radiographic progression

Author	Population	No.	Biomarker	Classification	Results
Syversen et al. ¹⁰	RA ≤4 yr	136	C2C (baseline serum)	SHS rapid >1 vs. slow <1 (radiographic progression per yr, progression change baseline to 5 or 10 yr)	NS
Mullan et al. ⁹	RA	45	C2C (baseline, 1, 3, 6, 9, 12-mo serum)		C2C ↑ at 1, 3 mo
	PsA	17	C1,2C	SHS rapid >1.5 vs. slow <1.5	C1,2C ↑ at 1, 3 mo
	(mean 11 yr, DAS28 >3.2)		CPII	(radiographic progression at 1 yr)	NS
Verstappen et al. ¹¹	RA ≤1 yr	87	ΔCOL (ΔC2C + ΔC1,2C + ΔCPII)		ΔCOL ↑ at 1, 3, 6, 9 mo
			C2C (1, 2, 3, 4-yr serum)		C2C ↑
			C1,2C CS846 CPII	66 th = SHS >7.4 vs. 33 rd percentile = SHS <2.3 (radiographic progression over 4-yr span)	C1,2C ↑ CS846 ↑ NS
Ishiguro et al. ⁷	RA (mean 10 yr)	63	C2C (knee SF)	Mild vs. moderate vs. severe RA	NS
			CS846	Mild vs. moderate RA	CS846 ↓
			CPII	Mild vs. moderate vs. severe RA (Larsen score: 0, 1 = mild; 2, 3 = moderate; 4, 5 = severe)	NS
Mansson et al. ⁸	RA ≤2 yr	18	CS846 (baseline serum)	Rapid vs. slow hip-joint radiographic progression	CS846 ↓
			CPII	(Larsen score: rapid = 46; slow = 4 at 2 yr)	NS

Number, number of patients investigated in the studies; DAS28, disease activity score based on 28 joints; mo, month; NS, not significant. PsA, psoriatic arthritis; RA, rheumatoid arthritis; SF, synovial fluid; SHS, SharpvanderHeijde score; yr, year.

Table 2 Overview of the literature on the (significant) relation between biomarker and the disease activity

Author	Population	No.	Biomarker	Classification	Results
Mullan et al. ⁹	RA	45	C2C (baseline, 1, 3, 6, 9, 12-mo serum)	DAS28 responders vs. nonresponders (at 3 mo)	C2C ↓
	PsA	17	C1,2C	(responder: ≥ 0.6 improvement and DAS28 ≤ 5.1 , nonresponder: < 0.6 improvement OR DAS28 > 5.1)	NS
	(mean 11 yr, DAS28 > 3.2)		CPII		NS
			Δ COL (Δ C2C + Δ C1,2C + Δ CPII)		Δ COL ↓
Mullan et al. ⁹	RA	45	C2C (baseline, 1, 3, 6, 9, 12-mo serum)		C2C ↓ at 1 mo
	PsA	17	C1,2C	Remission vs. no remission	C1,2C ↓ at 1 mo
	(mean, 11 yr, DAS28 > 3.2)		CPII	(remission = DAS28 < 2.6 at 6 mo)	NS
			Δ COL (Δ C2C + Δ C1,2C + Δ CPII)		Δ COL ↓ change 1 mo

DAS28, disease activity score based on 28 joints; mo, month; NS, not significant; number, number of patients investigated in the studies; PsA, psoriatic arthritis; RA, rheumatoid arthritis; SF, synovial fluid; yr, year.

The C2C serum ELISA detects a cartilage-specific collagen type II collagenase cleavage neoepitope [2]. The C1,2C ELISA detects a collagenase generated collagen type I and II cleavage neoepitope [3]. The CS846 ELISA detects an epitope on chondroitin sulfate of newly formed large aggrecan molecules [4]. The CPII ELISA recognizes epitopes of the propeptide of collagen type II reflecting synthesis [5].

Values of all four biomarkers were log transformed to obtain normal distributions. Additionally, seven extreme data point outliers derived from C2C, CS846, and CPII (based on visual inspection) were excluded for analysis.

(Long-term) outcome measurements

The long-term outcome of RA patients was determined by the radiographic joint progression and by the mean disease activity over a 5-year period of treatment. To assess radiographic joint progression, radiographs of hand and feet were made at baseline and every subsequent year. Radiographs were independently scored according to the Sharp-vanderHeijde score (SHS) [14] by two readers, blinded to clinical information. The mean yearly radiographic progression rate between baseline and 5 years was used as the outcome measure. For this rate, if 5-year radiographs were not available, the mean of the measurements between 4 and 6 years was used, or scores at 4 or 6 years, depending on the data available. Because scores were not normally distributed, the log-transformed progression rate (log rate +1) was used.

The mean disease activity over a 5-year period was determined by calculating a time-averaged value of the DAS28 [15] by using the area under the curve (AUC) from baseline until 5 years after treatment. If more than two yearly time points were missing, no time-averaged DAS28 could be calculated.

The early response has been shown to be a predictor for long-term outcome [16] and was therefore also taken into account in the analysis. The DAS28 at baseline and 6 months was used to calculate the early EULAR response. Patients were classified as good, moderate, or nonresponders based on their early (change in) disease activity. Good responders should have a DAS28 score ≤ 3.2 at 6 months and an improvement from baseline > 1.2 ; nonresponders a DAS28 score > 5.1 and an improvement between 0.6 and 1.2 or only an improvement of ≤ 0.6 . Patients with moderate response had a response in between the good responders and the nonresponders.

Statistical analyses

The change in biomarker values was calculated by subtracting the baseline biomarker value from the 1-year value for all biomarkers. Furthermore, sum scores of (changes in) markers representing synthesis (CS846 and CPII) and sum scores of (changes in) markers for degradation (C2C and C1,2C) were calculated. Finally, the ratio of (the sum scores of) synthesis and degradation markers were calculated. Because ranges of individual biomarker values differ, Z-scores (calculated by subtracting the average value from the individual value divided by the standard deviation) were used for the sum and ratio scores.

The relation between the individual (change, sum, and ratio of) biomarker values and long-term outcome (that is, mean yearly radiographic progression rate and time-averaged DAS28) was investigated by linear regression analysis, adjusting for the treatment strategy (that is, intensive tightly controlled or conventional MTX-based strategy).

Second, to investigate whether biomarker values were of additional value over already known baseline

predictors (rheumatoid factor (RF) and joint damage or disease activity at baseline, respectively), multiple linear regression analysis was used, adjusting for treatment strategy. The sum and ratio scores were considered in the analysis only when the individual biomarkers had a significant association with the outcome in the initial analysis. In the final model also, the early (6-month) EULAR response was added, by means of two dummy variables (good and moderate response, with nonresponse as the reference category).

The statistical software SPSS 15.0 was used for the analyses. A P value < 0.05 was considered statistically significant.

Results

Of 133 patients in the CAMERA trial, unfrozen serum samples were available at baseline and at 1 year of treatment. Of these patients, 75 had been treated according to an intensive, tightly controlled MTX-based strategy, and 58 patients according to a conventional MTX-based strategy. For five patients, no mean yearly radiographic progression rate could be calculated because of missing scores. For 11 patients, no time-averaged DAS28 could be calculated because more than two DAS28 scores were missing. Baseline characteristics of patients with missing data were not statistically significantly different from those of patients with complete data. Clinical characteristics and biomarker data of the patients are shown in Table 3. Note that radiographic progression is limited (median (IQR) radiographic progression rate over 5 years is 1.0 (0.0 to 3.4); mean (SD) value, 2.7 (4.5) SHS units).

In the analyses correcting for treatment strategy, C1,2C at baseline, the change in C2C and in C1,2C, and (consequently) the sum of the standardized changes in C2C + C1,2C levels were statistically significantly related to the mean yearly radiographic progression rate (all $P < 0.05$; Table 4). Only the change in CPII levels was related to time-averaged DAS28 ($P = 0.03$; Table 4).

In the multiple linear regression analyses, the change in C1,2C and the sum of the standardized changes in C2C + C1,2C levels were significantly related ($P = 0.004$ and $P = 0.02$, respectively) to mean yearly radiographic progression rate in addition to RF, baseline joint damage, and early (6-month) EULAR response. However, when including both changes in biomarkers values in the analysis, they were no longer statistically significant ($P = 0.13$ and $P = 0.94$, respectively). The change in C1,2C was chosen for the final model because this biomarker had the highest standardized beta, and the final model had the highest R^2 when compared with the sum of the standardized changes in C2C + C1,2C levels; furthermore, including only one biomarker instead of two is more efficient.

Table 3 Clinical and biomarker characteristics obtained at baseline and follow-up of all available patients

Characteristic	No. = 133	
Female gender (%)	87 (65)	
Age (years)	53 (14)	
RF positivity (%)	87 (65)	
Baseline DAS28	5.6 (1.0)	
Baseline joint damage	0.0 (0.0-0.0)	
EULAR good responders (%)	50 (38)	
EULAR moderate responders (%)	58 (44)	
EULAR no responders (%)	24 (18)	
Time-averaged DAS28	3.0 (0.9)	
Radiographic progression rate	1.0 (0.0-3.4)	
C2C (ng/ml)	Baseline	90 (71-124)
	1 yr	86 (70-109)
C1,2C (ng/ml)	Baseline	359 (286-427)
	1 yr	349 (269-415)
CS846 (ng/ml)	Baseline	100 (68-155)
	1 yr	113 (69-183)
CPII (ng/ml)	Baseline	335 (207-551)
	1 yr	436 (220-613)

Mean (SD) is shown for age and (baseline and time-averaged) DAS28; median (IQR) is shown for all other (non-normally distributed) continuous variables. For all categoric variables, number (%) of patients is shown. The EULAR response was determined after 6 months of treatment; time-averaged DAS28 and radiographic progression rate were calculated over 5 years of treatment; all other variables were determined at baseline unless otherwise stated. RF, rheumatoid factor; DAS28, disease activity score based on 28 joints.

The R^2 of the final model increased from 0.23 without biomarker to 0.28 including the change score of C1,2C (Table 5). When early response was not included, results were comparable, and the R^2 of the model changed from 0.20 to 0.27 if C1,2C was added. The standardized beta values showed that the influence of the biomarkers on prediction of the mean yearly radiographic progression rate was much smaller than, for instance, the predictive influence of baseline joint damage (standardized beta = -0.24 vs. 0.44, respectively; Table 5).

The change score of CPII was not statistically significantly related ($P = 0.18$) to time-averaged DAS28. The R^2 of the model increased marginally from 0.32 without biomarker to 0.34 including this biomarker (Table 6). When early response was not included in the model, the R^2 increased from 0.13 to 0.21 by adding the biomarker, but CPII was still not statistically significantly related to time-averaged DAS28. The standardized beta values also showed that the influence of the biomarkers was much smaller than those of RF, baseline disease activity, and early EULAR response (Table 6).

Discussion

The results show that some of the biomarkers have a small predictive value for long-term outcome in early RA, but clearly less, compared with established

Table 4 Association between biomarker values and the long-term outcome measures

Biomarker		Long-term outcome (5 years after treatment)					
		Yearly radiographic progression rate			Time-averaged DAS28		
		No.	B	95% CI	No.	B	95% CI
C2C	Baseline	126	0.08	-0.28 to 0.44	120	0.11	-0.25 to 0.47
	1 yr	126	-0.23	-0.67 to 0.20	120	0.18	-0.25 to 0.61
	Change	126	-0.59	-1.14 to -0.03	120	0.04	-0.52 to 0.60
C1,2C	Baseline	126	0.47	0.001 to 0.95	120	0.10	-0.40 to 0.60
	1 yr	127	0.14	-0.36 to 0.65	121	0.22	-0.29 to 0.73
	Change	126	-1.00	-1.80 to -0.20	120	0.33	-0.52 to 1.18
CS846	Baseline	127	-0.06	-0.27 to 0.16	121	-0.08	-0.29 to 0.14
	1 yr	126	-0.07	-0.29 to 0.15	120	-0.03	-0.26 to 0.20
	Change	126	-0.01	-0.23 to 0.21	120	0.05	-0.18 to 0.28
CPII	Baseline	124	0.14	-0.07 to 0.35	118	-0.05	-0.26 to 0.17
	1 yr	125	0.13	-0.09 to 0.34	119	0.10	-0.12 to 0.31
	Change	122	-0.07	-0.34 to 0.20	122	0.30	0.02 to 0.57
ZC2C + ZC1,2C	Baseline	125	0.07	-0.02 to 0.17			
	1 yr	126	0.01	-0.11 to 0.08			
	Change	126	-0.13	-0.22 to -0.04			

Biomarkers with B (95% confidence interval (CI)) values, which are shown in Bold type have a *P* value < 0.05 and have been included in the multiple regression analyses. Biomarkers values were determined at baseline, at 1 year, and the change between 1 year and baseline. Next are the sum and ratio scores (based on Z-values), determined when individual biomarkers had a significant association with the outcome in the initial analysis.

DAS28, disease activity score based on 28 joints; *n*, number of patients investigated; 95% CI, 95% confidence interval.

predictors. Only the change in C1,2C, the sum of the standardized changes in C2C + C1,2C levels, and the change in CPII were of added value for the mean yearly radiographic progression rate and the time-averaged DAS28, respectively. However, the explained variance of the final prediction models was low and therefore not useful for clinical practice, and both biomarkers increased the explained variance only marginally (and not statistically significantly for CPII).

Possible explanations for not finding a relation with all biomarkers are multiple. Importantly, it should be considered that blood for serum was collected during the day, which will influence the biomarker levels [17]. With respect to changes in biomarkers, it might have been worthwhile to evaluate changes in biomarkers within a shorter time span (for example, 3 or 6 months from baseline). However, no biologic samples were available at these time points. Also of relevance are the small

variances in outcome regarding the radiographic progression due to the low radiographic scores, despite the 5 years of follow-up. We compared other investigations of the four biomarkers (see Tables 1 and 2) with our own data; patients in the other studies had higher radiographic scores at baseline and had, on average, also higher disease durations (varying from 1 to 10 years RA). The available radiographic scores at baseline of the evaluated studies range from 6.8 to 60 for SHS (mean) and 2 to 7 for the Larsen score (median) compared with 0 SHS (median) in our study. Verstappen *et al.* [11] investigated the same biomarkers comparing fast (>7.3 SHS units/year) and slow progressors (<2.3 SHS units/year) and found significant differences in biomarkers values, except for CPII, in another cohort of patients with early RA. However, these slowest progressors (calculated over a 4-year period) in this previous study are comparable to the patients with the fastest progression (66th tertile >2.4

Table 5 Added predictive value of biomarkers over already known predictors for mean yearly radiographic progression rate over 5 years of treatment

Item	B	95% CI	Standardized beta	<i>P</i>	R ²
Intercept	0.54	0.12 to 0.96		0.013	
Treatment strategy	0.13	-0.16 to 0.43	0.08	0.375	0.000
RF positive	0.29	-0.02 to 0.60	0.15	0.063	0.029
Baseline joint damage	0.09	0.06 to 0.13	0.44	0.000	0.211
EULAR good response ^a	-0.36	-0.78 to 0.07	-0.20	0.100	
EULAR moderate response ^a	-0.13	-0.52 to 0.27	-0.07	0.534	0.229
C1,2C change from 1 yr to baseline	-1.11	-1.87 to -0.36	-0.24	0.004	0.283

^aEULAR nonresponse was used as reference category. 95% CI, 95% confidence interval; RF, rheumatoid factor.

Table 6 Added predictive value of biomarkers over already known predictors for time-averaged disease activity (DAS28) over 5 years of treatment

Item	B	95% CI	Standardized beta	P	R ²
Intercept	1.90	0.99 to 2.81		0.000	
Treatment strategy	-0.29	-0.62 to 0.03	-0.16	0.076	0.076
RF positive	0.35	0.02 to 0.68	0.18	0.040	0.098
Baseline disease activity	0.25	0.10 to 0.40	0.30	0.001	0.169
EULAR good response	-0.84	-1.30 to -0.37	-0.46	0.001	
EULAR moderate response	-0.19	-0.64 to 0.25	-0.11	0.393	0.322
CPII change 1 yr to baseline	0.18	-0.08 to 0.43	0.12	0.178	0.335

EULAR nonresponse was used as reference category. 95% CI, 95% confidence interval; RF, rheumatoid factor.

SHS units/year) in our present study (calculated over a 5-year period). Important to consider is that, because of improved treatment (strategies), the progression rate now in the Western community will hardly exceed the progression rate of the present cohort. This progress in treatment effectiveness and tight control strategies titrating treatment to the disease course of an individual patient might counterbalance the predictive value of biomarkers in prediction of disease outcome. However, it should not be ignored that also in the previous studies with higher radiographic progression rates, the relation of these biomarkers with outcome was not straightforward (see Table 1).

In a *post hoc* analysis evaluating all sum and ratio scores of synthesis and degradation markers (instead of only the ones when the individual biomarker had a significant association), no significant associations were seen with both the mean yearly radiographic progression rate and the time-averaged DAS28 over a 5-year period of treatment; this also applied for the multiple linear regression analysis (data not shown). The possible influence of age and gender on the biomarker values was also investigated with multiple linear regression analyses; adding these variables to the models did not change the results (data not shown). Using logistic regression analysis comparing progressors versus nonprogressors in radiographic progression (any radiographic damage at 5 years) also did not change the results (data not shown). When patients were selected with a minimum radiographic progression rate of 1 point per year (SHS of 5 units at 5 years), the relation with the biomarkers did not improve (data not shown). In case progression in joint space narrowing and erosions were taken apart, because biomarkers primarily represent cartilage turnover, no significant relations with biomarkers were found (data not shown).

Although glucocorticosteroid (GC) use was prohibited during the 2-year trial period; after 2 years, GC use was free. Only a limited number of patients used GC ($n = 13$). Because GC may influence joint damage significantly [18], analyses were performed in the group of

patients not using GC during the 5 years of treatment. In these *post hoc* analyses, no clear relations between radiographic joint damage and biomarkers were found.

The direction of the relation between the biomarkers and the mean yearly radiographic progression rate and time-averaged DAS28 was not anticipated. An increase in C1,2C during 1 year of treatment, which indicates more connective tissue degradation, led to lower mean yearly radiographic progression rate, whereas a higher time-averaged DAS28 was reached with an increase in cartilage collagen synthesis, as determined by an increase in CPII between baseline and 1 year of treatment. Conversely, *in vitro* data reveal that the neoepitope can increase when collagenase activity is blocked [19]. This is because collagenase can cleave the neoepitope that it generates [3]. In osteoarthritis (OA) serum, CPII increased with progression of OA (Poole *et al.*, unpublished data), similar to that in the present study on RA. As in general, contrasting relations have been found (Tables 1 and 2 and this study), clearly the nature, origin, and metabolism of these (and other) biomarkers require further investigation [20].

Based on the present results, the investigated markers are not the first choice in predicting long-term outcome in individual patients with early RA. The available studies together with the present results suggest that the role of these markers in predicting long-term outcome is at most modest. They might, conversely, be of value for other joint diseases or in distinguishing RA from other arthritis conditions. Significant differences in these biomarkers were reported when comparing RA with psoriatic arthritis [6], OA [6,7], and controls [8]. When we investigated the baseline biomarker values of the early RA patients of the CAMERA trial with controls, also significant differences were seen (all $P < 0.01$; data not shown). For assessment of progression in treatment with anti-TNF, these biomarkers appeared of use [9].

Biomarkers in general might be of value in prediction of the long-term outcome of RA. CTX-II [21-25], CTX-I [22,24], MMP-3 [25,26], COMP [27], calprotectin [28], RANKL [29,30], and IL-6 [31] all had a relation with

(long-term) radiographic progression and/or the disease-activity score. Of all these biomarkers, urinary CTX-II is at present the most frequently used and best-performing marker. Recently, a trial demonstrated urinary CTX-II and DAS28 almost equally effective when used to monitor disease activity and in treatment decisions aiming at remission of disease of RA [32]. Unfortunately, in our study, no urine samples were obtained.

Conclusions

In conclusion, the change in C1,2C and CPII in the first year after onset have a small added predictive value for radiographic progression and disease activity, respectively, over a 5-year period, although the predictive value is too small to be useful in daily clinical practice.

Abbreviations

ACR: American College of Rheumatology; AUC: area under the curve; CAMERA: Computer Assisted Management in Early Rheumatoid Arthritis; DAS28: disease activity score based on 28 joints; DMARD: disease modifying anti-rheumatic drug; MTX: methotrexate; OA: osteoarthritis; RA: rheumatoid arthritis; RF: rheumatoid factor; SHS: SharpvanderHeijde score.

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Authors' contributions

MB, SV, PW, JJ, JB, and FL contributed to the conception and design of the study. MB and PW contributed to the analysis of data. MB, SV, PW, JJ, JB, and FL contributed to the interpretation of data. NJ and MV provided study participants. Article drafts were written by MB and critically revised by all authors. The final version of the manuscript was approved by all authors.

Competing interests

The authors declare that they have no competing interests.

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