

Automated Measurement of Joint Space Width in Small Joints of Patients with Rheumatoid Arthritis

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ABSTRACT. *Objective.* Comparison of performances of 5 (semi)automated methods in measuring joint space width (JSW) in rheumatoid arthritis.

Methods. Change in JSW was determined by 5 measurement methods on 4 radiographs per patient from 107 patients included in the COBRA trial (comparing sulfasalazine alone or in combination with methotrexate and corticosteroids). For each method the number of patients with sufficient available results was assessed (efficiency). An independent repeated measurement was carried out on a random sample of 30 patients' baseline and 1-year radiographs, to evaluate within-method reliability of change scores. Discriminatory ability (DA) of the measurement methods (between the 2 treatment arms) was compared with the DA of the Sharp-van der Heijde score (SHS) and its 2 components (erosion and JSW scores).

Results. The overall success rate varied widely between methods. Applying the chosen threshold of a minimum of 50% available joints with a change score per patient resulted in a success rate > 92% in 4/5 methods. Repeatability of measurements was good for most methods (intraclass correlation coefficient ≥ 0.80 in 4/5 methods). Almost all measurement methods in 3 followup periods (12/14) showed a lower mean loss of JSW in patients from the intensive treatment group, although this was rarely statistically significant, confirming the known difference in structural damage. JSW as measured by the (semi)automated systems often showed higher DA than the JSW score of the SHS, but was lower than the total SHS and erosion scores.

Conclusion. Although efficiency of the methods should be improved further, results already show good reliability and encouraging DA of most methods. Optimal information may be obtained with a combination of scoring of erosions and (semi)automated measurement of JSW. (First Release June 15 2008; J Rheumatol 2008;35:1288–93)

Key Indexing Terms:

RHEUMATOID ARTHRITIS RADIOGRAPH SCORING JOINT SPACE WIDTH

Rheumatoid arthritis (RA) is a chronic inflammatory disease leading to progressive destruction of joint structures, and thus functional disability. Besides inflammation (synovitis, acute-phase reactants, pain), a major outcome of clinical trials in RA is radiographic progression. However, current scoring methods, although widely applied, have several limitations such as limited generalizability due to the difficulty of standardizing scoring by different readers. The

use of an ordinal scale is also a theoretical limitation to measurement accuracy, which could be improved by an assessment of damage on a continuous metric scale, provided that the latter is reproducible. In this context, a subcommittee within the OMERACT imaging committee was formed after OMERACT 6 to test reliability, sensitivity to change, and feasibility of computer-based methods for measuring radiographic damage in the small joints of the

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hands and feet in patients with RA, and priority was given to assessment of joint space width (JSW)¹. Preliminary exercises with small numbers of patients were conducted and results, presented at OMERACT 7, were considered sufficiently encouraging to proceed with further studies including larger sets of images. In the present exercise, a metrological comparison between most currently available methods for (semi)automatically measuring JSW was performed using principles outlined by the OMERACT filter. Radiographs from a randomized controlled trial with proven efficacy on radiographic outcomes of an intensive therapeutic intervention in early RA (COBRA²) were digitized and used to evaluate the different systems.

Automated measuring systems

Main characteristics of 5 computer-based measurement systems of JSW that were evaluated in this exercise are summarized in Table 1. The tested methods differ in many aspects: the number of assessed joints across the 5 systems ranges from 8 [including only the 4 metacarpophalangeal (MCP) joints of each hand for the least comprehensive system] to 34 [including 4 MCP, 4 proximal interphalangeal (PIP) joints, 4 wrist joints, and 5 metatarsophalangeal (MTP) joints of each hand and foot for the most comprehensive systems]. The workflow of the programs is also diverse: the localization of the joints on the digitized radiographs can be performed by an algorithm or they have to be cropped, rotated, or centered for the measurement procedure by a technician. The same applies for the measurement process itself, which can be performed completely automatically by algorithms or with variable amounts of user input (e.g., presegmentation of contours). Finally, the absolute values obtained cannot be compared directly across meth-

ods, because they represent different entities (shortest or average distance, with different defined margins of the joint, etc.).

MATERIALS AND METHODS

Radiographics of wrists, hands, and feet from 107 patients included in the COBRA trial for whom all timepoints were available (baseline, 6, 12, and 18 months) were digitized at 20 pixels per mm (50 µm pixel size) in an 8-bit gray scale. Digitization was performed centrally, and a copy of the resulting batch of radiographs was sent to the developers of 5 methods for (semi)automated measurement of JSW. This clinical trial compared radiographic outcomes of patients with early RA who had been randomly allocated to one of 2 treatment regimens (sulfasalazine alone vs combination therapy of methotrexate, sulfasalazine, and temporary high-dose corticosteroids).

Treatment arm, patient identity, and timepoints were randomized and blinded to the person who applied the automated method. Because every method measures JSW differently, we focused on change over time. Change between baseline and a followup timepoint in a single joint is called change in a joint-pair. For the feasibility aspect we first determined the percentage of joints in which measurements were successful. Thereafter, we calculated the proportion of patients that could be measured based on different thresholds — 10%, 20%, 30%, etc. of successfully measured joints. Intramethod reliability (reproducibility) of measurement was evaluated by calculation of intraclass correlation coefficient (ICC; 2-way mixed, absolute agreement) based on 2 independent readings of baseline and 12-month radiographs in a random sample of 30 patients, and comparing the change in measurement over this time period. Because a correct interpretation of an ICC (“relative agreement”) would require comprehensive description of data distributions, we also present results of “absolute agreement” by means of smallest detectable change (SDC) values³. The discriminatory ability of the methods was assessed by a paired t-test comparing mean change of JSW over 3 time periods (baseline–6 months, baseline–12 months, and baseline–18 months) per treatment arm. Only patients with more than 50% evaluable joint-pairs as defined above were included in this latter analysis. Radiographs of Month 18 were not measured by method A, due to time constraints. Because the different methods measured a different kind and number of small joints of hands and feet, which could theoretically jeopardize a fair comparison of the 5 systems, the discrimina-

Table 1. Main characteristics of the 5 methods.

Method	Assessed Joints on Each Side	No. Joints	Location	Region of Measurement	Measurement	Method for Calculation of Mean Joint Space Width
A ^{5,6}	MCP 2–5 PIP 2–5 MTP 1–5 4 wrist joints	34	Manual: 3 points on each MCP, MTP, wrist joint	MCP: radian drawn from the perceived center of the arc PIP: full breadth of the joint	Automated	Distance between proximal and distal joint margins on vertical lines
B ^{7,8}	MCP 2–5 PIP 2–5 MTP 2–5	24	Automated	Fixed number of mm (dependent on joint type) centered within joint	Automated	Distance between proximal and distal joint margins on vertical lines
C ^{9,10}	MCP 2–5 PIP 2–5 MTP 1–5	26	Manual in COBRA set: 1 point in PIP 2–5, 1 point in MCP 2–5	Fixed number of mm (dependent on joint type) centered within joint	Automated	Averaged shortest distance at multiple locations
D ^{4,11}	MCP 2–5	8	Automated	Central part of the joint	Automated	Averaged shortest distance at multiple locations
E ^{1,12}	MCP 2–5 PIP 2–5 MTP 1–5 4 wrist joints	34	Manual: 1 marker on medial and lateral margins of joints	60% of joint span, from medial to lateral sides	Automated	Averaged shortest distance at multiple locations

MCP: metacarpophalangeal joints; MTP: metatarsophalangeal joints; PIP: proximal interphalangeal joints; wrist joints: capitatum-naviculare, capitatum-lunatum, radius-naviculare, radius-lunatum.

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tory ability was also calculated when taking into account only the second to fifth MCP joints, because these were assessed by all methods. This discriminatory ability was also compared with the original scoring method applied in the COBRA trial (Sharp-van der Heijde method, mean score of 2 independent readers), with additional separate comparisons of erosion and joint space narrowing scores.

All analyses were carried out using SPSS 12 and Excel 2002.

RESULTS

Feasibility of the 5 methods is illustrated in Figure 1. If only 50% measurable joint-pairs per patient was required, all methods except method D provided evaluable data. If a higher requirement (75% evaluable joint-pairs per patient) was applied, only 3 methods (A, B, and E) provided evaluable data. The yield of methods A, B, and E decreased only gradually by increasing requirements, but the yield of method C dropped sharply if more than 50% of evaluable joint-pairs was required. Method C seemed to have a pattern of randomly missing joints that could not be evaluated (e.g., third right MCP, second left PIP, etc.), as occurred in many patients, whereas methods A, B, and E were mainly unable to read an entire radiograph, although in a lower frequency.

Intrareader reliability (repeatability) of the methods is summarized in Table 2. In general, repeatability was very high for most methods: 4 of the 5 methods achieved an ICC > 0.8 (considered the cutoff for good reliability). Interestingly, when data were analyzed based on joint groups (e.g., only MCP joints), reliability of the measures appeared to be highly dependent on the region that was measured. In particular, the PIP joints (when measured) consistently yielded the worst reliability for each of the methods, while MCP and MTP joints achieved the most reliable measurements. The SDC were expressed as the percentage of the 99th observed percentile in values of change in the respective method in order to have a comparable measure across methods. The SDC ranged from 21% to 71%, with most methods around 50%.

If comparison was limited to the MCP joint measurements, the discriminatory ability of the 5 methods (the difference in change in MCP JSW between treatment groups) was as shown in Tables 3A, 3B, and 3C for all 3 evaluated time intervals. Only patients with at least 50% of evaluable joint-pairs were included in these analyses. This implies that

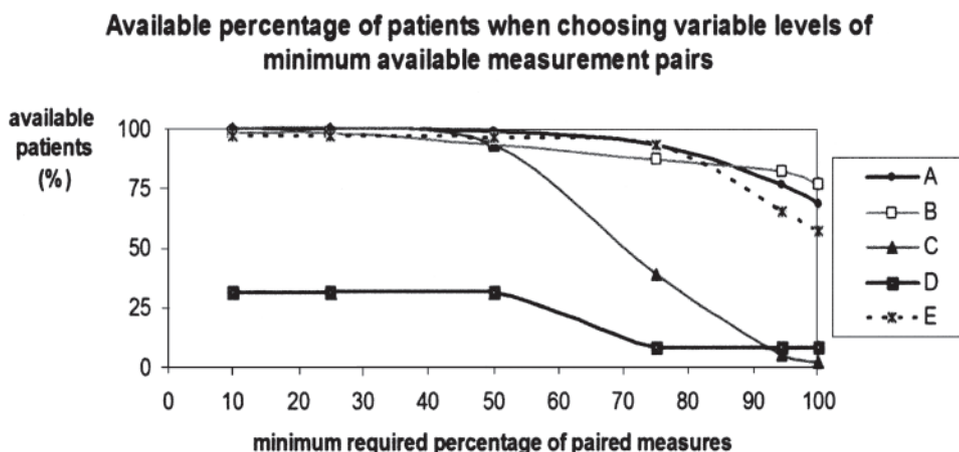


Figure 1. Efficiency of the 5 methods: available proportion of patients (Y axis) depending on various required successful reading rates (X axis) for the period from baseline to 12 months time period. Calculations are based on intentionally measured joints, as described by the respective methods.

Table 2. Intramethod reliability (independent repeated measurements) in assessing change in joint space over the baseline-to-12-month period on 2 independent readings in a random sample of 30 patients, per method (compared values are mean change per patient, patients with less than 50% of joints available being excluded).

Method	Based on All Measured Joints			Based on Separate Joint Groups (MCP, PIP, MTP, Wrists)		
	Valid Cases, %	ICC	SDC*, %	Range of Valid Cases, %	Range of ICC	Range of SDC*, %
A	100	0.98	21	100	0.96–0.98	20–30
B	100	0.96	41	100	0.78–0.97	22–43
C	96.7	0.80	57	42.4–96.7	0.60–0.80	41–79
D	23.3	0.81	59	23.3*	0.81*	59*
E	96.7	0.41	71	86.7–96.7	0.24–0.54	51–78

* Expressed as percentage of 99th observed percentile in values of change in the respective method. † Only MCP are measured in method D. ICC: intraclass correlation coefficient; SDC: smallest detectable change.

Table 3. Discriminatory ability of the 5 methods in measuring mean change of joint space width on last 4 metacarpophalangeal joints of both hands over the evaluated time intervals.

Table 3a. Baseline-6 months time interval.

Method	n*	Mean Change (SD)		t-test	p
		Monotherapy	COBRA		
A	105	-0.059 (0.095)	-0.045 (0.104)	-0.693	0.490
B	104	-0.055 (0.110)	-0.035 (0.122)	-0.884	0.379
C	80	-0.036 (0.078)	-0.013 (0.075)	-1.326	0.189
D	35	-0.072 (0.135)	-0.062 (0.146)	-0.198	0.844
E	102	-0.086 (0.272)	-0.087 (0.313)	-0.008	0.993

Table 3b. Baseline-12 months time interval.

Method	n*	Mean Change (SD)		t-test	p
		Monotherapy	COBRA		
A	106	-0.074 (0.120)	-0.051 (0.124)	-1.003	0.369
B	98	-0.064 (0.127)	-0.035 (0.127)	-1.136	0.259
C	82	-0.034 (0.087)	-0.031 (0.082)	-0.185	0.853
D	33	-0.021 (0.283)	-0.061 (0.165)	-0.507	0.616
E	104	-0.107 (0.308)	-0.060 (0.403)	-0.657	0.512

Table 3c. Baseline-18 months time interval.

Method	n*	Mean Change (SD)		t-test	p
		Monotherapy	COBRA		
A	NA	NA	NA	NA	NA
B	103	-0.067 (0.133)	-0.055 (0.132)	-0.445	0.657
C	80	-0.079 (0.132)	-0.041 (0.102)	-1.451	0.151
D	37	-0.082 (0.215)	-0.042 (0.199)	-0.579	0.566
E	104	-0.143 (0.349)	-0.060 (0.368)	-1.164	0.247

* Number of patients available for analysis with a 50%-available joint-pairs threshold. NA: not applicable.

a direct comparison of discriminatory ability across methods cannot be made, because different patients and joints were used depending on the various reading efficiencies. Again, one can get an impression of accuracy of each method in assessing the MCP by comparing the values given in the second columns of Table 3: compared to the other methods, method D had a consistently lower number of evaluable patients for all time periods. Although no significant difference was found when focusing on MCP only, the great majority of the comparisons (12/14) showed a treatment effect in the “expected direction,” that is, reflecting a lower loss of JSW in the patients from the intensively treated group, as compared to the monotherapy group. On the other hand, when using all available data (all measured joints for each method), results improved dramatically in methods measuring a substantial number of joints (methods A, B, C, and E), especially for the baseline to 6 months time period (Tables 4A, 4B, 4C). Additionally, the more comprehensive systems were able to discriminate between treatments better than the “default” Sharp-van der Heijde score for change in joint space narrowing using the average of 2 independent

readers’ scores. The erosion score and the aggregated erosion and joint space narrowing score of the Sharp-van der Heijde method, however, outperformed the (semi)automated methods with respect to discrimination.

DISCUSSION

Applying available automated methods for measuring JSW on a complete set of radiographs from a clinical trial in RA allowed us to compare “real-life” performance of a set of computer-based tools in terms of feasibility, effectiveness, reliability, and discriminatory ability.

A single set of radiographs was used to evaluate performances of the automated methods of joint space measurement, and thus a direct comparison of the different methods was possible for some aspects as long as the missing values were limited (especially when focusing on classical metrological features such as reliability and discrimination). Many aspects of the different systems are definitely distinct. There are important differences between the methods with regard to the time needed to measure a radiograph, which is due to different software, and differences in the time needed

Table 4. Discriminatory ability of the 5 methods in measuring mean change of joint space width on all measured joints over the evaluated time intervals.

Table 4a. Baseline-6 months time interval.

Method	n*	Mean Change (SD)		t-test	p
		Monotherapy	COBRA		
A	105	-0.068 (0.084)	-0.031 (0.073)	-2.466	0.015
B	104	-0.062 (0.077)	-0.011 (0.096)	-2.959	0.004
C	98	-0.024 (0.041)	0.001 (0.067)	-2.251	0.027
D	35	-0.072 (0.135)	-0.062 (0.146)	-0.198	0.844
E	102	-0.076 (0.148)	-0.028 (0.148)	-1.639	0.104
Erosion score	107	5.130 (5.405)	2.105 (3.323)	3.429	0.001
Joint space narrowing score	107	1.580 (3.208)	0.947 (1.929)	1.253	0.213
Total score	107	6.710 (7.147)	3.053 (4.460)	3.124	0.002

Table 4b. Baseline-12 months time interval.

Method	n*	Mean Change (SD)		t-test	p
		Monotherapy	COBRA		
A	107	-0.088 (0.110)	-0.054 (0.094)	-1.743	0.084
B	99	-0.035 (0.168)	-0.038 (0.091)	0.094	0.925
C	99	-0.024 (0.054)	-0.012 (0.065)	-0.927	0.356
D	33	-0.021 (0.283)	-0.061 (0.165)	-0.507	0.616
E	104	-0.096 (0.151)	-0.029 (0.152)	-2.239	0.027
Erosion score	107	7.878 (8.381)	4.061 (5.925)	2.666	0.009
Joint space narrowing score	107	3.122 (5.234)	2.605 (4.682)	0.537	0.592
Total score	107	11.000 (11.618)	6.667 (9.493)	2.113	0.037

Table 4c. Baseline-18 months time interval.

Method	n*	Mean Change (SD)		t-test	p
		Monotherapy	COBRA		
A	NA	NA	NA	NA	NA
B	103	-0.071 (0.100)	-0.032 (0.157)	-1.493	0.139
C	100	-0.037 (0.075)	-0.029 (0.078)	-0.521	0.603
D	37	-0.082 (0.215)	-0.042 (0.199)	-0.579	0.566
E	104	-0.111 (0.203)	-0.054 (0.175)	-1.548	0.125
Erosion score	107	10.918 (11.125)	6.439 (8.569)	2.294	0.024
Joint space narrowing score	107	4.827 (6.905)	4.026 (6.780)	0.601	0.549
Total score	107	15.745 (16.144)	10.465 (14.321)	1.784	0.077

* Number of patients available for analysis with a 50%-available joint-pairs threshold. NA: not applicable.

for a technician's interventions. While the fully automated systems virtually instantaneously localize and measure the joint of interest, the semiautomated methods require a substantial amount of the operator's time: an approximation of the total time required (to obtain measurements in one patient, i.e., from radiographs of 2 hands and 2 feet when applicable) was 15, 5, 2.5, 7, and 22 minutes for methods A, B, C, D, and E, respectively. The operator's intervention also introduces a possibly important source of systematic error in semiautomated methods. For example, if the margins of a joint are judged and marked in different ways from one operator to another or by one operator making 2 measurements at

different times, this may compromise reliability and generalizability. As a consequence, the interpretation of results is operator-dependent to some extent, which is similar for scoring systems such as the Sharp-van der Heijde score.

The SDC help to interpret how important a small difference in ICC (e.g., 0.96 vs 0.98) is when an absolute reliability cutoff is derived from the same data (e.g., the SDC dropping from 41% of the observed maximal change to 21%). The SDC is based on the limits of agreement between 2 measurements for a certain method, and gives a cutoff level of the method above which a change can be seen as a real change beyond measurement error.

Clearly, the efficiency of the methods in providing successful measurements for all joints that were presented for measurement needs to be improved before the computer-based methods can be used more widely. Excluding too many patients from the radiographic analysis of a clinical trial due to unevaluable joints per patient is unacceptable, in that it can cause a loss of important information and may yield potentially biased results (systematic selection of joints). Even if we applied a very lenient cutoff level for evaluability (more than 50% of joints with available paired measurements), a surprisingly low proportion of patients remained in the analysis with one of the methods, and a somewhat more stringent requirement also would have hindered a second system. As a comparison, the proportion of missing scores that was observed in the same trial for joint space narrowing score (Sharp-van der Heijde score) at a joint level was very low (< 1%), while the most effective automated method (method B) had a total of 4.4% of PIP and MCP joints considered “not measurable” in our set of radiographs (wrist joints were not measurable by method B). For radiographs produced with an imaging protocol consistent with the training set, however (i.e., beam geometry, hand positioning), 7.4% of MCP joints were “not measurable” with method D⁴.

However, despite these apparently low performances, even a modest requirement of 50% of comparable joints over time resulted in a consistently accurate discriminatory ability of most methods. The seemingly disappointing p-values should be compared with those from the scoring of joint spaces, namely, the Sharp-van der Heijde scoring system: the discriminatory ability of the scoring method is largely determined by the erosion score, while the joint space narrowing score contributes in a limited manner. Although this phenomenon is not specific for the COBRA trial, further evaluation in other trials will provide more insight into the real merits of the measurement methods.

There are limitations to this study, as it is an ad hoc comparison of systems that are at different stages of preclinical development, and results should therefore be interpreted carefully. Preliminary exclusions of anatomical regions or of sets of images may stem from decisions in software engineering rather than intrinsic limitations of particular methods. An additional limitation is that adjustments of existing algorithms of the automated methods to characteristics of the image set were almost completely precluded.

In view of these limitations, the remarkably better discrimination in most automated methods should be regarded as a promising sign of efficiency in accurately assessing change in JSW over time. This superiority of the semiautomated methods over the scoring with regard to their discriminatory abilities might be a consequence of the use of a continuous scale in the computer-based measurements of

JSW, while the Sharp-van der Heijde method (like all scoring systems) applies an ordinal scale, the latter being inherently less sensitive to change. In addition, a fully automated method for assessing all radiographic abnormalities in RA will require not only JSW but also erosion measurement. However, although such projects are under development, there is still a long way to go before they become available as a routine, and the general reliability and sensitivity of scoring makes the task especially difficult. An important consideration in this regard is to determine from a clinical point of view what is the optimal tradeoff between automated measurement and scoring. One could imagine that optimal discriminatory ability will be obtained with a combination of scoring erosions and automated measurement of joint space width by computer-driven techniques.

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