EXTENDED REPORT

One year outcome of undifferentiated polyarthritis

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Objective: To identify variables that can predict a progressive outcome after one year of follow up in patients presenting with undifferentiated polyarthritis (UPA) at an early arthritis clinic.

Methods: New patients with arthritis in two or more joints of less than three years’ duration were categorised at entry as UPA or as rheumatoid arthritis (RA) based on the clinical diagnosis of the rheumatologist. Outcome variables after one year were radiographic damage (Sharp/van der Heijde score) and functional status (Health Assessment Questionnaire: HAQ score). A progressive disease at one year was defined as radiographic progression \( \geq 4 \), or one year radiographic damage \( \geq 10 \), or HAQ score \( \geq 1 \). The baseline variables of patients with UPA with a progressive or mild outcome were compared.

Results: 280 patients (70% women; median age 56 years (range 18–90), median duration of symptoms 3.5 months) were included. 203 (72%) patients were clinically diagnosed as having RA and 77 (27%) as having UPA. The group of patients with progressive UPA (n=32 (42%)) had a significantly higher mean age, prevalence of arthritis of the hands, and disease activity (DAS28) at the first visit compared with the patients of the mild UPA group (n=45 (58%)). The RA group had significantly more frequent serum IgM-RF positivity, higher mean disease activity (DAS28) and mean C reactive protein concentration, more frequent symmetric arthritis, and arthritis in more than three joint groups than the progressive UPA group. Six (19%) of the progressive UPA group versus eight (4%) of the RA group did not receive disease modifying antirheumatic drugs during the first year.

Conclusions: After one year of follow up, 32 (42%) of the patients with UPA had a progressive disease. A progressive outcome was associated with older age, higher disease activity, and arthritis of the hands at baseline. To avoid undertreatment of patients with UPA, treatment should be based on severity rather than on diagnosis.

Rheumatoid arthritis (RA) is a chronic inflammatory joint disease of unknown cause with a large variation in the severity of inflammation and joint destruction. Joint damage starts in an early phase of the disease. To prevent joint destruction, there is growing consensus that optimal management of RA requires both early diagnosis and early aggressive treatment. However, therapeutic decisions are often difficult to make because of the non-specific early clinical features of RA.

The 1987 revised criteria for RA of the American College of Rheumatology (ACR; formerly, the American Rheumatism Association) have a limited value (a low sensitivity and specificity) in early arthritis, because they were developed in patients with established disease. In contrast, an experienced rheumatologist can make a reliable clinical judgment on the presence or absence of RA after two weeks. Most patients with diseases such as psoriatic arthritis, sarcoidosis, or reactive arthritis can easily be separated from patients with RA. In the remaining patients with undifferentiated oligoarthritis and polyarthritis (UPA), defined as arthritis of more than one joint without fulfilment of the ACR criteria for RA, the diagnosis may change in the first years of follow up. Some patients will enter remission, some will progress to RA, and some to another rheumatic disease such as spondyloarthropathy. There is a scarcity of studies on the outcome and prognostic factors of patients with UPA in an early phase in comparison with studies of early RA. Also, patients with a diagnosis of UPA may be perceived as not in need of disease modifying antirheumatic drug (DMARD) treatment to the extent that patients with RA do.

The purpose of this study was (1) to find prognostic factors at entry for progressive UPA after one year; (2) to investigate the outcome after one year in patients with progressive UPA compared with the outcome after one year in patients with a clinical diagnosis of RA at entry.

PATIENTS AND METHODS

Patients

In a prospective study of patients with early arthritis seen at a large rheumatology outpatient clinic, the study comprised patients who were referred between September 1995 and April 1998, aged 18 years or older, with peripheral arthritis of at least two joints and less than three years of symptom duration. Patients who had previously been treated with a DMARD, patients with bacterial, psoriatic, or crystal induced arthritis, and patients with reactive arthritis, osteoarthritis, sarcoidosis, or systemic autoimmune diseases were excluded. Patients were diagnosed as having RA based on the clinical diagnosis of an experienced rheumatologist at the second visit to the clinic (one to three weeks after the first visit); the remainder of the patients were considered as having oligoarthritis or polyarthritis (UPA).

Disease variables

A trained investigator performed a structured interview and physical examination within two weeks of the first visit to the rheumatologist. Follow up assessments were performed at three, six, nine, and 12 months.

Variables assessed were demographic characteristics, time of onset of complaints (defined as persistent pain, swelling, or both), duration of morning stiffness, serum rheumatoid factor (IgM-RF), serum C reactive protein (CRP), disease activity...
Table 1 Comparison of baseline characteristics between mild UPA, progressive UPA, and the RA groups

<table>
<thead>
<tr>
<th>Baseline variables</th>
<th>Mild UPA† (n=45)</th>
<th>p Value mild v progressive UPA</th>
<th>Progressive UPA (n=32)</th>
<th>p Value progressive UPA v RA</th>
<th>RA‡ (n=203)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years) median (range)</td>
<td>45 (21–81)</td>
<td>**</td>
<td>53 (25–81)</td>
<td>**</td>
<td>61 (18–90)</td>
</tr>
<tr>
<td>Women [%]</td>
<td>58</td>
<td>72</td>
<td>69</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Disease duration (months) median (range)</td>
<td>3 (0–25)</td>
<td>*</td>
<td>4 (0–36)</td>
<td>*</td>
<td>4 (0–36)</td>
</tr>
<tr>
<td>Arthritis ≥3 joints [%]</td>
<td>47</td>
<td>*</td>
<td>72</td>
<td>**</td>
<td>92</td>
</tr>
<tr>
<td>Symmetric arthritis [%]</td>
<td>42</td>
<td>63</td>
<td>**</td>
<td>87</td>
<td></td>
</tr>
<tr>
<td>Arthritis of hands [%]</td>
<td>62</td>
<td>**</td>
<td>91</td>
<td>97</td>
<td></td>
</tr>
<tr>
<td>Arthritis of small joints (PIP, MCP, MTP [%])</td>
<td>78</td>
<td>81</td>
<td>94</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Arthritis of large joints (knee, ankle, shoulder, elbow [%])</td>
<td>97</td>
<td>77</td>
<td>87</td>
<td></td>
<td></td>
</tr>
<tr>
<td>IgMRF positive [%]</td>
<td>18</td>
<td>16</td>
<td>***</td>
<td>51</td>
<td></td>
</tr>
<tr>
<td>CRP (mg/ml) median (SD)</td>
<td>190 (270)</td>
<td>*</td>
<td>140 (160)</td>
<td>***</td>
<td>340 (400)</td>
</tr>
<tr>
<td>DAS28 score (median)</td>
<td>3.8 (1.5)</td>
<td>*</td>
<td>44 (9.9)</td>
<td>***</td>
<td>5.2 (1.2)</td>
</tr>
<tr>
<td>Sharp/van der Heijde score (median range)</td>
<td>2.0 (0–55)</td>
<td>*</td>
<td>8.0 (3–8)</td>
<td>***</td>
<td>5.0 (0–136)</td>
</tr>
<tr>
<td>HAQ score (mean range)</td>
<td>0.6 (0.6)</td>
<td>0.9 (0.6)</td>
<td>1.1 (0.8)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*p<0.05; **p<0.01; ***p<0.001 for differences between groups (Student’s t test).
†UPA, undifferentiated polyarthritis; ‡RA, rheumatoid arthritis.

were clinically diagnosed by the rheumatologist as RA and 77 (27%) as UPA at baseline. Forty five (58%) of these patients with UPA had polyarthritis and 32 (42%) had oligoarthritis.

The reasons for loss to follow up after one year were death (n=3), from malignancy (n=2), from renal failure (n=1), moving home (n=7), and non-compliance (n=27). Of the 37 (13%) patients who were lost to follow up, 22 (59%) were clinically diagnosed as RA and 15 (41%) as UPA. The baseline characteristics of the 15 patients with UPA lost to follow up were similar to those of the 77 completers (data not shown). However, the 22 patients with RA lost to follow up had a significantly higher mean baseline DAS28 score (p<0.05) than the 203 completers. The mean follow up of the UPA non-completers was 5.6 months and the mean of their last DAS score was 3.5 (the mean DAS score of the completers at six months was 2.9).

The first analysis was performed to determine prognostic factors for a progressive course of UPA. Of the patients qualified as “progressive UPA”, 21 (66%) fulfilled the criterion of radiographic damage ≥4 within one year.

The next step was to search for differences between the progressive UPA and RA groups. At baseline, the RA group showed a significantly higher mean HAQ score (p<0.01), and arthritis in more than three joint groups (p<0.05) than the progressive UPA group (table 1).

The radiographic damage and radiographic progression at one year was least in the mild UPA and highest in the RA group (fig 1). The mean one year HAQ score was highest in the progressive UPA group (fig 2).

To identify independent prognostic variables of progressive UPA, a stepwise logistic regression analysis was performed. Among the 77 patients with UPA, age (odds ratio=1.03; p<0.01; 95% confidence interval [95% CI] 1.01 to 1.09) and arthritis of the hands (odds ratio=4.2; p<0.05; 95% CI 1.04 to 4.2) were independently associated with a higher risk of developing progressive UPA.
patients with arthritis of the hands were 4.2-fold more likely to have a progressive disease than patients without arthritis. In other words, for every additional year of age there was a 1.05-fold higher risk of having a progressive disease and the proportion of patients that had ever used DMARDs during the first year was larger in the RA group than in the progressive UPA group (96% vs 81%, p<0.05; table 2). Moreover, the mean (SD) period of DMARD use was longer for the RA group than for the progressive UPA group (8.0 (3.2 ) vs 5.7 (4.3) months). After one year of follow up, patients with RA used sulfasalazine and methotrexate more often and hydroxychloroquine less often than patients with UPA (table 2).

DISCUSSION

In this study almost half of the patients with early UPA had a poor functional status, progressive radiographic damage, or both at one year. However, only 81% of these patients were treated with DMARDs. Patients with a progressive disease were older and had involvement of the hands more often than patients with mild disease.

The investigation of early arthritis is hampered by the lack of a clear definition of RA. The ACR criteria for RA have several shortcomings in early arthritis. For example, sensitivity and specificity are too low; synovitis has to be present for six weeks; and the criteria are no longer fulfilled when synovitis subsides in response to treatment. Also, individual criteria cannot be studied as predictors for the development of RA, because these factors are part of the criteria set. This disadvantage is not present when RA is defined as the clinical diagnosis of an experienced rheumatologist. Diagnosis of an experienced rheumatologist seems to be more subjective but has been found to be reliable. Because the final goal of classification of patients with early arthritis is to predict the prognosis, the focus should be shifted from classifying patients into RA and non-RA categories, towards different outcome categories.

The few available studies of the outcome of early undifferentiated arthritis and rheumatoid arthritis have used different outcome variables, such as the persistence of arthritis, functional ability, and radiographic damage. The comparison of these studies is further complicated by differences in the definition of UPA. In the present study, patients with progressive UPA could be differentiated from patients with a mild disease by an older age and more frequent involvement of the hands at baseline than in patients with a mild disease. This accords with a study of patients with palindrome rheumatism in which older age, early involvement of hand joints, and female sex and a positive IgM-RF predicted the subsequent development of RA. Others confirmed the finding that IgM-RF positivity was associated with the onset of RA and persistence of arthritis. Tunn and Bacon examined predictors for persistence of arthritis after one year in a cohort of 65 patients with early UPA. They showed that the combination of IgM-RF positivity with an increased erythrocyte sedimentation rate carried a high relative risk for persistent disease.

The inferiority of the ACR criteria for RA can be explained by the lack of a clear definition of RA. The ACR criteria for RA have several shortcomings in early arthritis. Because these factors are part of the criteria set, this disadvantage is not present when RA is defined as the clinical diagnosis of an experienced rheumatologist. Diagnosis of an experienced rheumatologist seems to be more subjective but has been found to be reliable. Because the final goal of classification of patients with early arthritis is to predict the prognosis, the focus should be shifted from classifying patients into RA and non-RA categories, towards different outcome categories.

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In a study by Wolfe et al,11 IgM-RF positivity also was the best predictor of development of RA. The association of a positive rheumatoid factor with progressive UPA was not found in our study (table 1).

Green et al12 studied predictors for disease outcome at six months in patients with early arthritis. They concluded that predictors for persistent disease were a longer disease duration, IgM-RF positivity, and the presence of the shared epitope. In the present study, it was also found that a short disease duration was associated with a mild disease outcome, but the correlation was not strong. Furthermore, Green et al concluded that a baseline CRP is not predictive in very early inflammatory arthritis, which accords with our finding for patients with UPA.

Harrison et al,13 focusing on the prediction of future disability in 277 patients with early inflammatory polyarthritis, found that female sex, longer disease duration, large joint involvement, and high baseline HAQ were associated with a high HAQ score after one year. In this cohort, HLA-DRB1 alleles had no influence on the likelihood of disease persistence and only a modest association with functional disability. In the subgroup of seronegative patients however, HLA-DRB1 did have an influence on the development of erosions.15 In the present study, genetic tests were not performed, which makes comparison difficult.

The study of UPA may seem irrelevant if it is assumed that patients with progressive UPA actually have RA. However, only 16% of the patients with progressive UPA were clinically classified as having RA by the rheumatologist at one year and only 31% fulfilled the ACR criteria. From the total UPA group, 10% were clinically classified as having RA by the rheumatologist at one year and 17% fulfilled the ACR criteria. There was no significant difference in outcome (radiographic damage) between patients changing after one year from the diagnosis of UPA into RA and patients retaining the diagnosis of UPA.

According to Hülsemann et al16 the outcome of patients with UPA seems to be less severe compared with those with RA. Complete remission occurred in 54% of the patients with UPA and only 7% of them developed RA after two years. A difference from our study was that 11% of their population had reactive arthritis, which was an exclusion criterion in our study. The same high percentage of patients reaching complete remission in UPA was found by Wolfe et al.11 In the present study more aggressive treatment was prescribed for patients with RA compared with patients with progressive UPA. Whereas 4% of the patients with RA did not receive a DMARD during the first year, 19% of the patients with progressive UPA were not treated with DMARDs, which suggests insufficient treatment.

In conclusion, in this study 42% of the patients with UPA showed a severe functional status, progressive radiographic damage, or both after one year which could be predicted by age, a high disease activity score, and arthritis of the hands at baseline. These patients developed a progressive disease that may not be recognised and treated adequately in time. A considerable number of the progressive UPA group were treated with a mild—that is, an antimalarial, drug—or were not treated with a DMARD at all. Therefore, treatment strategy should be focused more on disease severity than on diagnosis alone.

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REFERENCES

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