**Benefits of Anticitrullinated Peptides Examination in Rheumatoid Arthritis**

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**Background:** Anti-citrullinated peptides antibodies (ACPA) are specific for rheumatoid arthritis and have been implicated in disease pathogenesis. ACPA examination is a new component of ACR/EULAR 2010 classification criteria for rheumatoid arthritis. ACPA positivity predicts a more erosive disease course with severe joint damage and extra-articular manifestations. **Objectives:** To evaluate the benefits of ACPA examination in patients with early undifferentiated arthritis and patients with rheumatoid arthritis. **Methods:** We examined patients with arthritis and tested them for ACPA positivity. In every individual patient we evaluated if ACPA examination was necessary to establish the diagnosis of rheumatoid arthritis, or to change treatment, or if the diagnosis could have been established without ACPA examination (ACR/EULAR 2010 classification criteria was met without ACPA scoring). **Results and Conclusions:** The study was placed in Slovak Republic. We examined 833 patients with arthritis. There were 43 patients, or 62% of a subgroup of 69 who were ACPA positive whose ACPA examination was not needed—ACR/EULAR criteria was met without ACPA scoring. This number represents 5.1% of the total number examined. There were 15 patients, or 22% of the subgroup and 1.8% of the total whose diagnosis was revised to rheumatoid arthritis due to ACPA positivity—ACR/EULAR criteria were met solely with ACPA scoring. There were 11 patients (16% and 1.3%) whose medication was changed due to ACPA positivity. ACPA examination is useful in 3.1% of all examined patients. When we correlate data on ACPA positive patients, 38% of the patients profit from ACPA examination. Considering the relatively low price of ACPA testing, this examination should not be excluded.

**Key Words:** ACR/EULAR 2010 classification criteria, anti-citrullinated peptides antibodies, rheumatoid arthritis

**Introduction**

Rheumatoid arthritis is a complex disorder with many different aspects and many different forms at onset. It occurs in 1% of the population. It is characterized by chronic symmetric polyarthritis. Joint damage begins at the synovial membrane, and ongoing cartilage and bone destruction results in joint deformities.

A large number of patients also have extra-articular and systemic symptoms at the same time. Rheumatoid arthritis shortens life expectancy by approximately 10 years and it is a significant cause of disability and handicap in the population, ultimately leading to high economic loss in society.[1]

Most of the recommended medications that are used to treat rheumatoid arthritis have an immunosuppressive effect and they may have multiple side effects and adverse events. Risk-benefit ratio should be assessed individually in every patient.

Like most of the systemic connective tissue diseases, rheumatoid arthritis has a multi-systemic heterogeneous clinical presentation. That is the

**Access this article online**

Quick Response Code:

- **Website:** www.njcponline.com
- **DOI:** 10.4103/njcp.njcp_411_17

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reason why it may be difficult to establish a correct diagnosis in the early stages of the disease, and also why the “disease classification criteria” were formulated, consisting of clinical, histopathological, and laboratory features. Establishing the correct diagnosis early and starting treatment adjusted to the individual patient as soon as the diagnosis is established is essential and a cardinal principle in order to prevent the patient’s disability and handicap.[2]

The key task is to identify markers with the highest possible sensitivity and specificity for rheumatoid arthritis (diagnostic markers) and also markers with the highest possible range of prediction of the severity of the disease (prognostic markers). Consequently, patients with a suspected severe course of the disease should be treated with more aggressive medication immediately after establishing diagnosis, and patients with a suspected mild course of the disease would be treated with less aggressive treatment to avoid a possible adverse event.[3]

Anti-citrullinated peptides antibodies (ACPA) are autoantibodies that are highly specific for rheumatoid arthritis. Citrulline is a nonstandard amino acid created by deamination of the amino acid arginine by the enzyme peptidylarginine deiminase (PAD). Citrullination is a common biological process that occurs during inflammation, apoptosis, and cell differentiation. In the presence of certain genetic and environmental backgrounds, proteins are citrullinated excessively and they are not destroyed by the immune system, but contrariwise antibodies against citrullinated proteins are formatted and rheumatoid arthritis becomes manifested.[4]

There are many ACPA tests available with different sensitivity and specificity for rheumatoid arthritis. ACPA examination is a new component of ACR/EULAR 2010 (American College of Rheumatology/European League against Rheumatism) classification criteria for rheumatoid arthritis. Low positivity of ACPA scores 2 points, whereas high positivity scores 3 of a possible 10 points from the individual score.[4]

ACPA positivity predicts a more erosive disease course with severe joint damage and extra-articular manifestations. ACPA testing should be performed in individuals whose clinical signs suggest rheumatoid arthritis or who have already been diagnosed with undifferentiated arthritis. These antibodies also should be tested in patients who have already been diagnosed with rheumatoid arthritis to predict a possibly more progressive course of the disease.[5]

MATERIALS AND METHODS
At our workplace (out-patient clinic) we performed a clinical study whose target was to evaluate the benefits of ACPA examination in patients with early arthritis and patients with rheumatoid arthritis. We examined patients with arthritis and tested them for ACPA positivity during a period of 5 years. All patients were educated and informed about our study, they agreed to participate and signed an informed consent form. In the group of patients with ACPA positivity we measured levels of antibodies. For low-level positivity we scored 2 points, for high-level positivity (thrice higher than the upper limit of normal) we scored 3 points (according to ACR/EULAR 2010 classification criteria) of a possible 10 points of individual score.

We reviewed every individual patient and reconsidered whether ACPA examination was necessary to establish the diagnosis of rheumatoid arthritis or not. (Meaning the diagnosis could have been established without ACPA examination. The ACR/EULAR 2010 classification criteria were met without scoring ACPA positivity.) We also reviewed if the results of ACPA examination had led to changes in the patient’s medication in the group of patients with an already established diagnosis of rheumatoid arthritis.

RESULTS
In total we examined 833 patients with arthritis, of which 290 patients were diagnosed with rheumatoid arthritis according to ACR/EULAR 2010 classification criteria. There was a subgroup of 69 APCA positive patients, 63 with rheumatoid arthritis, 3 with osteoarthritis, and 3 with systemic lupus erythematosus [Table 1 and Figure 1]. Total EULAR score of patients without ACPA was 2.4 ± 0.62 and total EULAR score of patients with ACPA was 8.3 ± 0.34. Rheumatoid arthritis severity scale (RASS) in ACPA negative patients was 41, 36, and RASS in ACPA positive patients was 79.88. Within ACPA positive subgroups were 43 patients (62% of the

<table>
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<th>Table 1: Patients with arthritis and ACPA positivity according to the diagnosis</th>
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<td><strong>Diagnosis</strong></td>
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Nigerian Journal of Clinical Practice ¦ Volume 21 ¦ Issue 10 ¦ October 2018

subgroup, 5.1% of the total) whose ACPA examination was not needed—ACR/EULAR 2010 criteria was met without ACPA scoring/ACPA positivity. There were 15 patients (22% of the subgroup, 1.8% of the total), whose diagnosis was revised to rheumatoid arthritis due to ACPA positivity—ACR/EULAR 2010 criteria was met solely with ACPA positivity scoring. There were 11 patients (16%, of the subgroup, 1.3% of the total) whose medication was changed—augmented due to ACPA positivity or reduced due to ACPA negativity [Table 2 and Figure 2].

Discussion

ACPA examination is recommended by ACR and EULAR as it is a part of the classification criteria for rheumatoid arthritis. Many studies have confirmed its diagnostic and prognostic values, but there had been no study performed to evaluate the importance of this examination for a single individual patient.

We discovered that ACPA examination is helpful and useful in 3.1% of all examined patients with arthritis (those patients whose diagnosis was revised or whose medication was changed due to ACPA examination results). When we correlate our data to the subgroup of ACPA positive patients we see that 38% of the patients profit from ACPA examinations.

Table 2: Analysis of ACPA examination benefits in patients with arthritis and ACPA positivity (n=69)

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<th>No changes in diagnosis or treatment</th>
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<tr>
<td>Number of patients</td>
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Figure 1: Analysis of patients with arthritis and ACPA positivity according to the diagnosis

Figure 2: Analysis of ACPA examination benefits in patients with arthritis and ACPA positivity (n=69)

The price of ACPA examination kits is relatively low, with prices varying in local laboratories. Average price for a single ACPA examination is approximately 5 USD/Euros. It is definitely worth the price, especially for patients who can avoid disability and functional handicap, their families, communities, and society as a whole.

Conclusions

Rheumatoid arthritis is a chronic systemic disease that is shortening life-expectancy and possibly leading to functional disability and handicap. As the disease has heterogeneous clinical manifestation we use ACR/EULAR 2010 classification criteria to establish the diagnosis. There are diagnostic and prognostic markers of the disease, one of them is anti-citrullinated peptides antibodies. Patients with a suspected severe course of the disease should be treated immediately after establishing diagnosis with more aggressive drugs to avoid progression to joint deformities and disability. Patients with a suspected mild course of the disease should be treated with less aggressive drugs without serious adverse events.

Financial support and sponsorship

None.

Conflicts of interest

There are no conflicts of interest.

References

