Predictive value of antinuclear antibodies in autoimmune diseases classified by clinical criteria: Analytical study in a specialized health institute, one year follow-up

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Abstract

Introduction: Determination of antinuclear antibodies (ANA) by indirect immunofluorescence (IIF) is usually the initial test for the diagnosis of systemic rheumatic diseases (SRD). Assigning predictive values to positive and negative results of the test is vital because of lack of knowledge about ANAs and their usefulness in classification criteria of SRD leads to inappropriate use. Methods: Retrospective study, ANA tests requested by different specialties, correlation to patients’ final diagnosis. Results: The prevalence of autoimmune disease was relatively low in our population yielding a low PPV and a high NPV for the ANA test. 40% of the patients had no clinical criteria applied prior to test. Coexistence of two or more autoimmune disorders affects prevalence and predictive values. Conclusion: Application of the test after careful evaluation for clinical criteria remarkably improves the positive likelihood ratio for the diagnosis.

1. Introduction

Immunological assays for the detection of antinuclear antibodies (ANA) are useful and important complementary tools for the diagnosis and follow-up of patients with autoimmune diseases [1]. The identification of the antigen–antibody coupling is the common end-point for all techniques; however, several differences exist as for the utility, sensitivity, specificity, and predictive values of each test [1,2].

In general, if a patient presents clinical manifestations of an autoimmune disease, the first test to be requested is ANA detection by indirect immunofluorescence using HEp-2 cells, due to its great sensitivity [1,3]. The different possible patterns, the intensity, and the titers obtained by consecutive dilutions must be carefully examined. Identification of the antigens recognized by the ANA is further evaluated by more specific tests such as ELISA, radioimmunoassay (RIA) or electroimmunotransference (EIT) [2,4].

The use of these tests requires knowledge of their fundamental aspects and also of the clinical classification criteria of each disorder in order to contribute to an appropriate diagnosis [5,6].

The usefulness of this testing has been evaluated in retrospective studies of patients with systemic rheumatic disease (SRD), and it has been proven that its positive predictive value is low due to the relatively large amount of false positive results. For specific rheumatic diseases, the ANA test yields a positive predictive value of 11%, a negative predictive value of 97%, and a sensitivity and specificity of 42% and 85% respectively [7].

Several physiological and pathological factors might favor the development of ANA in the non-rheumatic population, such as pregnancy, advanced age, family history of autoimmune disease, as well as infectious, cardiovascular or oncological diseases [8–12]. This situation conveys challenges such as interpretative standardization [13].

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A high percentage of patients with high autoantibodies titers do not manifest any clinical signs of autoimmune disease. This may be due to the existence of circulating antigens that are not routinely tested for, such as those resulting from infectious stimuli, from multifactorial synthesis or those naturally produced by CD5+ cells [14]. For this reason, clinicians should pay close attention to the titers in which the ANAs are reported, taking into account that in healthy individuals, antibodies should be negative or can be present in low titers, and that intermediate titers may be present in non-affected relatives of patients with autoimmune diseases or in elders with chronic infections or neoplasms [8,11,12,15].

In Mexico, ANA prevalence has been studied in healthy individuals and consensus has been reached as to consider positive a gross mottled pattern in dilutions over 1:160, while homogeneous, centromeric, peripheral or centriolar patterns should be considered positive even in dilutions as low as 1:40 [16]. Their presence can be, nevertheless, due to natural antigens [14,17,18].

In some instances the recognition of antibodies directed to known antigens cannot be achieved. This complicates the accurate measurement of the antibody’s predictive value [19,20].

The objective of the present study was to determine the predictive values (PPV, NPV) of ANA testing for suspected SDR by analyzing the pre-test assessment of rheumatologic clinical criteria as well as post-test diagnosis.

2. Methods

We analyzed samples for ANA studies requested to our lab during a twelve-month period. The tests were selected if they were performed by IIF in Hep-2 cells (INOVA Diagnostics INC San Diego USA) and if an initial positive result at a 1:40 dilution led to successive dilutions. An informed consent was obtained for each test form each patient.

Furthermore, the presence of specific auto-antibodies was evaluated by ELISA (ORGENTEC Diagnostica GmbH Carl-Seiss Mainz,Germany) using purified extractable nuclear antigens (ENA) for Sm, RNP/Sm, SSA/Ro, SSB/LA, Anti-Scl-70, and anti-centromere as well as crithidia luciliae substrate.

An ANA test was considered to be positive when titers were superior to the following dilutions: Nuclear pattern: homogeneous > 1:40, coarse speckled and fine speckled > 1:160, laminar and peripheral > 1:40. Cell cycle: nucleolar, centromeric, and centriolar > 1:40. Cytoplasmic > 1:80 and micotocondrial > 1:160.

Each patient’s clinical file was reviewed by a qualified rheumatologist to acknowledge, if the suspected diagnosis was confirmed or if there was an alternative final diagnosis. We confirmed form the records the evaluation made for the presence of diagnostic clinical criteria in each patient. Clinical criteria considered for each disease the following the guidelines for diagnosis.

2.1. Statistical analysis

Sample size was calculated by correlation as follows:

\[
  n = \frac{n'}{1 + n'N}
\]

\[
  n' = \frac{S^2}{\sigma^2}
\]

where \( N=1374 \), standard error, \( Se=0.025, p=0.18, S^2 \) the sample variance \( p(1-p)=(0.18)(1-0.18)(0.82)=0.1476 \), \( \sigma^2 \) is
In 364 (83%) samples, nuclear antibodies were found with dilutions 1:40 and no antibodies were found in 9 cases (2%). In 193 out of the 364 (52%) antibodies against specific antigens were found.

Frequency of test requests and use of clinical criteria by each department from our institution are shown in Table 1.

There was a total of 373 ANA tests performed. In 364 (98%) patients, antibodies were found in dilutions of 1:40. Out of these, SRD was confirmed in 213 (57%) cases and it could not be confirmed in 160 (42%).

From the 213 patients with confirmed SRD, in 187 (88%) clinical criteria were applied prior to the blood test, but only in 167 (78%) SRD was confirmed and it could not be confirmed in 160 (42%).

In 20 patients (9%), the clinical criteria were applied but no autoimmune disease could be diagnosed. 120 individuals (56%) were diagnosed with SRD by both clinical and laboratory criteria. In 47 cases (22%), clinical criteria were used prior to requesting the ANA test but no antigen-specific antibodies were found; 22 (10%) patients had no clinical criteria applied prior to requesting the test, but ANAs could be found in high dilutions and were positive against specific antigens. Therefore the physician concluded that the patients had an autoimmune disease.

Finally, there were 24 patients (11%) with no clinical criteria applied prior to soliciting the ANA test and no antigen-specific antibodies were found, however, they presented IIF patterns compatible with autoimmune disease on high titers.

No autoimmune disease was found in 160 patients. In 6 (4%) of them, clinical criteria were applied prior to soliciting the ANA test,
but antibodies were found only in dilutions of 1:40 with antigen-specific antibodies, and in 49 patients (31%) no clinical criteria were applied and the antibodies were found only in low ANA titers and antigen-specific antibodies. Also, from these 160 patients, 91 (57%) had not previously met the mentioned clinical criteria according to the suspected SRD.

These also corresponded mainly to patients with heart or kidney disease.

Dilutions that predominated were 1:40 although there was some percentage of antibodies in high titers. Even an antigen-specific antibody could be found in 8 cases; in these, the attending physicians applied their clinical judgment and discarded SRD. Regardless if they presented specificity towards an antigen or not, when the criteria were applied SRD was demonstrated in 167 patients (78%) while SRD could only be diagnosed in 46 (22%) when no criteria were used.

No SRD was found in 141 (88%) of the requests for ANA test in which no clinical criteria were applied compared to 19 (12%) in the group in which clinical criteria were used. This difference achieved statistical significance with an OR of 26 (95% CI 14–50, \( p < 0.0001 \)). This analysis strongly supports the application of clinical criteria prior to the request of antibody testing in patients in whom autoimmune disease is suspected.

The pre-test probability of the antigen-specific antibody test is of 57%. Whenever clinical criteria are applied, we observed an improvement of 32%. Nevertheless, when the antibody test was used to confirm SRD without the use of clinical criteria this value diminished to 15% as shown in Table 2.

Further analysis of the information revealed that when clinical criteria and specific antibodies are present, we achieve a sensitivity of 57% and a specificity of 96%; when one or both are negative we get an increase in sensitivity but a loss in specificity. We evaluated the effect of various combinations of these results in Tables 3 and 4.

The aforementioned improvement in the post-test likelihood ratio results should be observed every time considering both tools (clinical criteria and specific antinuclear antibody testing) are not independent for the diagnosis of autoimmune disease but usually sequential.

Pre-test probability is 0.57

\[
\frac{0.57}{1 - 0.57} = 1.3
\]

The likelihood ratio for stand-alone testing and their combined utility is shown in Table 5, in panel A we see the that pre-test likelihood for a patient with suspected autoimmune disease when a test for antigen-specific antibodies is requested is 1.94; the post-test probability with an antigen-specific antibody can be calculated as follows:

\[
\frac{1.94}{1.94 + 1} = 65\% \quad \text{Probability post – test}
\]

Calculating the post-test likelihood ratio when clinical criteria were used to classify the disease we obtain a result of 3.55 (Panel B) and a post-test probability of 78%.

\[
\frac{3.55}{3.55 + 1} = 78\% \quad \text{Probability post – test}
\]

Panel C shows the result from combining both assessments against the application of only one evaluation (clinical or serological).

As presented in Table 6, Panel A (patients with autoimmune disease were \( n = 213 \)), sensitivity was 72% when clinical criteria were used prior to antigen-specific antibodies (T-tasa FP) while it only was 48% when they were not applied and antibody testing was performed directly.
Both tests are independent (in Panel B) (patients without autoimmune disease = 160), the specificity for not having autoimmune disease was of 65% when the clinical criteria were negative or not used. However, when the test was negative and clinical disease was of 65% when the clinical criteria were applied. The likelihood ratio for each test.

This supposed concordance should be further evaluated in larger series.

As shown in Table 7 panel a, the likelihood ratio to find an autoimmune disease is of 14.97 when both clinical criteria and the ANA test are dependent upon the disease's prevalence, with false positive results being more common.

In this study we also described the frequency of ANA, specificity for specific antigens, and type of nuclear pattern and their relationship with SRD. These results are shown in Table 8.

Equally rare patterns such as pattern NuMA antibodies were found in 7 cases (2%), out of which 5 were women and 3 men, with a mean age of 33 (28–64) for women and 52 (21–57) for men. Two of them were diagnosed with secondary APS, one with SLE plus AS, another one with SLE plus limited systemic sclerosis, one with rheumatoid arthritis and Sjogren's syndrome, one with ANCA-associated vasculitis and pulmonary thromboembolism, and a last one with atrial tachycardia plus pulmonary arterial hypertension. Two females presented PCNA antibodies; one was diagnosed with Takayasu arteritis and the other one with SLE plus dilated myocardiopathy. Two cases were reported as having antibodies against proliferating cells.

4. Discussion

It is known that positive and negative predictive values for any test are dependent upon the disease's prevalence, with false positive results increasing in those samples in which the disease has a low prevalence, therefore decreasing the test's positive predictive value [7].

Healthy subject, people with non-autoimmune diseases and those with a family history of autoimmune disease present a high percentage of antibodies in low titers [21,22].

Thirty-five (22%) patients without autoimmune disease presented ANA in 1:40 dilutions, but none were observed in dilutions over 1:320.

In cases with SRD, ANA could be found in dilutions over 1:320 in 199 (57%) compared to those without autoimmune disease 44 (26%), OR 3.45 (CI 95%, 2.19–5.50), and in dilutions between 1:2560 and 5120 71 (39%) in SLE, scleroderma, mixed connective tissue disease, overlap, Sjogren's syndrome and rheumatoid arthritis plus systemic lupus erythematosus. Even though they were observed in non-autoimmune diseases, this percentage was lower: 12 (8%), OR 5.88 (CI 95%, 2.95–11.94) (Fig. 1a–h).

### Table 5

Likelihood ratio for each test.

| (a) Specific-antigen antibodies | With autoimmune disease | Without autoimmune disease | Likelihood ratio |
|---------------------------------|-------------------------|---------------------------|-----------------|
| Antibodies for specific antigen  | Number                  | Proportion                | Number          | Proportion    |                      |
| +                               | 142                     | 142/213 = 0.66            | 55              | 55/160 = 0.34 | 0.66/0.34 = 1.94   |
| −                               | 71                      | 71/230 = 0.30             | 105             | 105/160 = 0.65| 0.30/0.65 = 0.46   |
| Total                           | 213                     |                           | 160             |              |                      |

| (b) Clinical criteria           | With autoimmune disease | Without autoimmune disease | Likelihood ratio |
|---------------------------------|-------------------------|---------------------------|-----------------|
| Classification according to criteria | Number                  | Proportion                | Number          | Proportion    |                      |
| +                               | 167                     | 167/213 = 0.78            | 20              | 20/160 = 0.12 | 0.78/0.22 = 3.55  |
| −                               | 46                      | 46/213 = 0.22             | 140             | 140/160 = 0.88| 0.30/0.65 = 0.39  |
| Total                           | 213                     |                           | 160             |              |                      |

| (c) Effect of different combinations of specific antibodies and clinical criteria over the diagnostic workup for SRD | | |
|-----------------------------------------------------------------------------------------------------------------|------------------|------------------|
|                                                                                                                 | Number | Proportion | Likelihood ratio |
| Clinical criteria and specific antibodies positive                                                              | 120    | 120/213 = 0.56 | 0.56/0.037 = 15.02 |
| One or both negative                                                                                           | 93     | 93/213 = 0.44  | 0.44/0.96 = 0.45  |
| Total                                                                                                          | 213    |               |                  |
Table 6
Evaluation of the combination of positive results on specific antibodies and utilization of clinical criteria for the diagnosis of autoimmune disease.

Panel A: Patients with autoimmune disease

|                | Positive specific antigen antibodies | Sensitivity of specific antigen antibodies (false positive ratio) |
|----------------|---------------------------------------|---------------------------------------------------------------|
|                | Positive | Negative | With clinical criteria | Without clinical criteria | Total |
| With clinical criteria | 120      | 47       | 167                      | 120/167 = 72%            |
| Without clinical criteria | 22       | 24       | 46                       | 22/46 = 48%             |
| Total           | 142      | 71       | 213                      | 142/213 = 67%           |

Sensitivity of specific antigen antibodies (1-true positive ratio)

|                | When specific antibodies ( + ) | When specific antibodies ( - ) | Total |
|----------------|-------------------------------|-------------------------------|-------|
| With clinical criteria | 120/142 = 85%               | 47/71 = 66%                  | 167/213 = 78%          |
| Without clinical criteria | 49/55 = 89%                | 91/105 = 87%                 | 140/160 = 87%          |

Panel B: Without autoimmune disease

|                | Positive specific antigen antibodies | Sensitivity of specific antigen antibodies (false positive ratio) |
|----------------|---------------------------------------|---------------------------------------------------------------|
|                | Positive | Negative | With clinical criteria | Without clinical criteria | Total |
| With clinical criteria | 6       | 14       | 20                      | 14/20 = 70%            |
| Without clinical criteria | 49      | 91       | 140                     | 91/140 = 65%           |
| Total           | 55      | 105      | 160                     | 105/160 = 65%          |

Sensitivity of specific antigen antibodies (1-true positive ratio)

|                | When specific antibodies ( + ) | When specific antibodies ( - ) | Total |
|----------------|-------------------------------|-------------------------------|-------|
| With clinical criteria | 49/55 = 89%                | 91/105 = 87%                 | 140/160 = 87%          |
requesting the test [21]. Positive antibodies in low titers may lead to confusion when trying to establish a diagnosis, and can become problematic when they are found at higher titers. It is well known that any test such as antibodies against a specific antigen conveys false positive and false negative results. This can lead to diagnostic and therapeutic errors by utilizing measures when they are not required [23,24].

In this analysis, we see that using clinical criteria before requesting the test provides a considerable improvement in the diagnostic workup. Antibodies considered to be specific for SLE, such as double strand anti-DNA, have been reported as well in Sjogren’s syndrome, dermatomyositis and cutaneous sclerosis [25–29]. In our series the percentage of SLE patients with positive ANA was of 49%, with varying frequency ranges from 6% to 50%, when SLE coexisted with other diseases, and in 90% of patients with renal damage, a finding known to bear a worse prognosis [30–33]. In non-rheumatic diseases we found anti-DNA antibodies in frequencies similar to those previously reported in the literature, supporting the idea of the existence of an immunological alteration in cardiovascular and renal diseases, which might be explained by previous infections [15,34,35].

Antibodies directed towards ribonucleoproteins (SM, RNP, SSB) are usually detected in SLE, but not in discoid lupus. Our results concur with previous literature [31,36]. As for SM antibodies, there are reported presence of them in 15–40% of cases; we found that they are present in 30% of cases of SLE when not associated with other diseases, with ranges that vary from 15% to 50% when another SRD coexists with SLE or there is damage to a specific organ [37,38]. Quite remarkable, elevated ANA titers are important in the diagnostic of rheumatic diseases, but it is also very important to be familiar to each laboratory’s cut-off points. Also the type of pattern of antibodies was found in some cases in close correlation with the presence of some autoimmune diseases. It is known that antibodies directed against ribonucleoproteins are associated with connective tissue diseases [39]. An homogeneous pattern might be proof of reaction against native single or double stranded DNA and associated with SLE. The centromeric pattern is

Table 7
(a) Likelihood ratios when two converging tools for autoimmune disease are positive. (b) Likelihood ratios for SRD diagnosis.

| (a) | Autoimmune disease | Likelihood ratio |
|-----|---------------------|-----------------|
|     | Present             | Absent          |               |
| Both positive tool (clinical criteria CC and specific ANA) | 120 (56%) | 6 (3.7%) | 0.56/0.038=14.97 |
| One or both | 47 | 14 | +22/49 |
| - | 93 (44%) | 154 (96%) |
| Total | 213 (100%) | 160 (100%) |

(b) Positive Likelihood Ratio (L.R. +)

| Stand-alone test | Clinical criteria | AAN against specific antigen |
|------------------|-------------------|------------------------------|
| Positive Likelihood Ratio (L.R. +) | (1–0.78/1–0.87)=1.94 |
| Total | 1–0.78/1–0.87=1.94 |

| Disease | Antigen | Pattern |
|---------|---------|---------|
| SLE     | n SM, RNP, SSA, SSB, SCL70 | Anti-centromere, Homogeneous, Discrete speckled, Coarse speckled, NUMA, PCNA |
| RA+SLE  | 43     | 13 (31) |   |
| SLE+APS | 13     | 1 (5)  | 1 (2) |
| SLE+SS  | 2      | 0      | 1 (2) |
| Discoid lupus | 6     | 0      | 0.66/1–0.66=0.66 |
| SLE+hyperthyroidism | 2  | 0      | 1 (2) |
| SLE+hypothyroidism | 5     | 0      | 0.5 |
| RA      | 17     | 0      | 0.14 |
| RA+SS   | 10     | 1 (10) | 0.55 |
| RA+hypothyroidism | 1     | 0      | 0.14 |
| IJA     | 5      | 0      | 0.14 |
| Scleroderma | 16    | 2 (10)| 0.14 |
| SCL+SS  | 2      | 0      | 0.14 |
| MCTD    | 5      | 0      | 0.14 |
| Overlap | 8      | 1 (5)  | 0.14 |
| Polymyositis | 2   | 0      | 0.14 |
| Dermatomyositis | 1    | 0      | 0.14 |
| SS      | 8      | 0      | 0.14 |
| SS+hypothyroidism | 2    | 0      | 0.14 |
| Devic syndrome | 3    | 0      | 0.14 |
| PAPS    | 38     | 0      | 0.14 |
| Fibromyalgia | 7    | 0      | 0.14 |
| Cardiopathy | 94   | 1      | 0.14 |
| Hypothyroidism | 3    | 0      | 0.14 |
| Nephropathy | 24   | 0      | 0.14 |
| Cancer  | 2      | 0      | 0.14 |
| Takayasu’s arteritis | 0    | 0      | 0.14 |

APS=primary antiphospholipid syndrome, IJA=idiopathic juvenile arthritis, MCTD=primary antiphospholipid syndrome, RA=rheumatoid arthritis, SCL=scleroderma, SLE=systemic erythematous lupus, SS=systemic scleroderma.
Table 9
Sensitivity, specificity, and predictive values according to autoimmune disease and specificity of antibodies.

| Disease          | SM   | DNA  | SSA  | SSB  | RNP |
|------------------|------|------|------|------|-----|
| SLE N=37         | S=15 | S=62 | S=40 | S=25 | S=43|
| RA + SLE N=8     | S=25 | S=50 | S=50 | S=25 | S=38|
| Discoid lupus N=6| S=0  | S=20 | S=0  | S=0  | S=25|
| SLE + APS N=11   | S=10 | S=17 | S=33 | S=0  | S=15|
| SLE + SS N=3     | S=33 | S=33 | S=67 | S=67 | S=33|
| SLE + hypothyroidism N=4 | S=0 | S=25 | S=75 | S=33 | S=0  |
| Sjogren’s syndrome N=8 | S=0 | S=0  | S=0  | S=38 | S=0  |
| PAPS N=35        | S=0  | S=15 | S=5  | S=0  | S=0  |
| RA N=16          | S=0  | S=20 | S=13 | S=0  | S=0  |
| IJA N=5          | S=0  | S=20 | S=0  | S=0  | S=0  |
| Scleroderma N=12 | S=67 | S=13 | S=23 | S=0  | S=38|
| Crest N=5        | S=0  | S=0  | S=0  | S=0  | S=60|

APS—antiphospholipid syndrome, IJA—Idiopathic juvenile arthritis, MCTD—mixed connective tissue disease, PAPS—primary antiphospholipid syndrome, RA—rheumatoid arthritis, SCL—scleroderma, SLE—systemic lupus erythematosus, SS—systemic sclerosis, S—sensitivity, E—specificity, PPV—positive predictive value, NPV—negative predictive value.

characteristic of CREST syndrome and those against nuclear RNA are associated with SLE and systemic progressive sclerosis [40]. However, other unusual nuclear ANAs are those against the Nuclear mitotic apparatus (NuMA), which might or might not be reported accurately depending upon the laboratory’s experience [41]. Their positivity is associated with connective tissue disease, 45% corresponding to Sjogren’s syndrome and undifferentiated connective tissue disease as well as autoimmune diseases against specific organs in 17% even though up to 38% have been found in non-autoimmune diseases [42].

In this study 2% of patients had NuMA, and they were associated with primary AS, one of them with optic neuritis and a possible Devic syndrome. The prevalence of these antibodies and their clinical significance has been previously reported in the literature [43,44].

Antibodies against the nuclear antigen of proliferative cells were described over 30 years ago in patients with chronic hepatitis B or C, and they have only been found in about 5% of patients with SLE. Their clinical significance has been recently studied in a meta-analysis and they have also been detected in polymyositis, systemic sclerosis and even healthy individuals. However their prevalence has not surpassed 2% in any group [45]. In this study, they were detected in two cases, one of them a 64-year-old woman with SLE and end stage renal disease, and the second one in a 23-year-old female with Takayasu arteritis and systemic arterial hypertension.

The presence of specific antibodies against cellular components such as nuclear or cytoplasmic molecules are specific for some diseases [46,47] while some other might be completely non-specific [48,49]. Moreover other findings might depend upon a clinical characteristic of the disease, such as neuropsychiatric lupus in which anti-p ribosomal antibodies have a 10% prevalence and were observed in 2% of all patients with SLE [46]. Nevertheless, some antibodies are related with organ specific alterations and could be prognostic markers [50]. Commonly, when a non-rheumatologist specialist requests an ANA test in a patient it is due to the presence of inflammatory signs and symptoms that most physicians would not overlook. However antibody-testing results does not consider previous clinical details and specific diagnosis becomes quite difficult [51–54].

We were able to confirm the dispersion and utility that these results have depending upon the clinicians’ specialty, the use of clinical criteria, and indirectly, the knowledge of some recommendations from guidelines.

We believe that in some cases the severity of the clinical picture and diagnostic uncertainty may justly requesting for these tests, however a positive result might turn out to be a confusing factor and therefore require an interpretation that should into account, in first place, the clinical context.

The use of the test in patients with SRD and a positive result might lead to a second test. Several studies attempting to obtain an appropriate use of laboratory tests have been published with the fair purpose of reducing unnecessary testing [4,55,56].

A non-medical factor, knowledge of the techniques and standardized procedures, contributes to the optimal use of the test. Other variables could contribute to the variability of the results such as ethnicity, the use of clinical criteria, and the coexistence of several autoimmune diseases or presence of several other antigens.

On the other hand, when the prevalence of a disease in a sample is low, positive predictive value tends to be low as well dictating the need to confirm the result by using a second test.

We found a low prevalence of autoimmune diseases in the requests of these tests, which were solicited by several specialists with differing criteria.

From another view, the high percentage of false positives may be attributed to the fact that only a certain set of antibodies are routinely tested for, not including other recognized antibodies such as antihistone, antinucleosomes, CENP-B, CENP-A, CENP-C, Sp100 protein, PML or NDP53, which could increase the predictive value of the test [57–62].
Speciﬁcity can be increased when clinical criteria and diagnostic algorithms are applied. This reduces unnecessary ANA tests and correlating with a better analysis, utilization, and clinical judgment by the physicians [52,63,64].

5. Conclusion

Positive and negative predictive value for the ANA test is low, it is dependent on the clinical context of the patient and if the physician relies on clinical criteria for its request. The use of clinical criteria speciﬁc for each probable disease prior to antinuclear antibodies testing increases the likelihood ratio for the diagnosis of autoimmune diseases. This also depends upon the phenotype of the disease and the coexistence of two or more diseases or the presence of other antigens, which are not routinely tested for in all laboratories.

The proper use of laboratory tests, in accordance to knowledge and interdisciplinary communication, signiﬁcantly improves the diagnostic yield of specialized evaluations.

References

[1] Hernandez Ramirez, Cabiedes. Immunological techniques that support the diagnosis of the autoimmune diseases. Reumatol. Clin. 2010;6(3):173–7.
[2] Cabiedes, Nunez-Alvarez. Antinuclear antibodies. Reumatol. Clin. 2010;6(4):224–30.
[3] Briolay, Gioud, Monier, et al. Antinuclear antibodies detected by indirect immunofluorescence on Hep2 cells and by immunoblotting in patients with systemic sclerosis. Autoimmunity 1989;2(2):165–76.
[4] Sheldon. Laboratory testing in autoimmune rheumatic diseases. Best Pract. Res. Clin. Rheumatol. 2004;18(3):249–69.
[5] Binder. Antinuclear antibodies (ANA): immunologic and clinical signiﬁcance. Semin. Arthritis Rheumatol. 1976;6(2):83–124.
[6] Marin, Cardiel, Cornejo, Viveros. Prevalence of antinuclear antibodies in 3 groups of healthy individuals: blood donors, hospital personnel, and relatives of patients with autoimmune diseases. J. Clin. Rheumatol. 2009;15(7):325–9.
[7] George, Giburd, Shoenfeld. The emerging concept of pathogenic natural autoantibodies. Hum. Antibodies 1997;8(2):70–5.
[8] Baumgarth, Tung, Herzenberg. Inherent speciﬁcities in natural antibodies: a key to immune defense against pathogen invasion. Semin. Immunopathol. 2005;26(4):347–62.
[9] Hamaguchi. Autoantibody proﬁles in systemic sclerosis: predictive value for clinical evaluation and prognosis. J. Dermatol. 2010;37(1):42–53.
[10] Verstegen, Duyck, Meeus, Ravelingien, De. Detection and identiﬁcation of antinuclear antibodies (ANA) in a large community hospital. Acta Clin. Belg. 2009;64(4):317–23.
[11] Hayashi, Koshiba, Nishimura, et al. Prevalence of disease-speciﬁc antinuclear antibodies in general population: estimates from annual physical examinations of residents of a small town over a 5-year period. Mod. Rheumatol. 2008;18(2):153–60.
[12] Sebastian, Roy, Kin, Mullick. Correlation of antinuclear antibody immunofluorescence patterns with immune proﬁle using line immunosassay in the Indian scenario. Indian J. Pathol. Microbiol. 2010;53(3):427–32.
[13] Lora, Lurino, Becker, Monticello, Brenol, Xavier. Clinical diagnostic performance of different methods for the detection of antibodies to extractable nuclear antigens in connective tissue diseases: a cohort study. Clin. Lab. 2011;57(7–8):625–9.
[14] Pittock, Lennon, et al. Neutrophilic and neutrophil-specific autoimmunity. Arch. Neurol. 2008;65(1):78–83.
[15] Buzkola, Berezin, Gau, et al. Antinuclear autoantibody proﬁle in the sera of women with hyperprolactinemia. J. Autoimmun. 1995;8(3):415–24.
[16] Falanga, Medsger Jr. TA, Reichlin. Antinuclear and anti-single-stranded DNA antibodies in morphea and generalized morphea. Arch. Dermatol. 1987;123(3):350–3.
