

ROUTINE TEST MANAGEMENT POLICY – 15.01.040

Biomarker Testing for Autoimmune Rheumatic Disease

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Policy Description

Systemic autoimmune rheumatic diseases (SARDs) are a diverse group of conditions that primarily affect the joints, bones, muscle, and connective tissue.¹ SARDs are characterized by dysregulated immunity and inflammatory responses, resulting in damage and destruction to joints, connective tissues, skin, blood elements, and other target organs; however, considerable diversity in clinical presentation, disease course, and treatment response exists.²

The diagnostic workup for SARDs may involve the antinuclear antibody (ANA) assay, which is used to detect autoantibodies (AAB) against intracellular antigens, originally known as antinuclear antibodies.³ Commonly used as part of the initial diagnostic workup to screen for evidence of systemic autoimmunity,⁴ detection and identification of AABs are important in the diagnosis of SARDs, such as systemic lupus erythematosus (SLE), Sjögren's syndrome (SjS), mixed connective tissue disease (MCTD), systemic sclerosis (SSc), and idiopathic inflammatory myopathies (IIMs).⁵ Extractable nuclear antigens or ENAs (a historical term from when the antigens were extracted from the cell into saline solution prior to testing) include Sm, U1 ribonucleoprotein (RNP), Ro, and La antigens, and are also useful for evaluating individuals with suspected connective tissue disease.⁶

Indications

1. For individuals with signs or symptoms of an autoimmune disease, screening for disease using antinuclear antibodies (ANA) is considered **reimbursable**:
 - a. Once during initial workup.
 - b. Up to two additional tests per lifetime if new or more severe signs or symptoms of an autoimmune disease develop.
2. For individuals with an abnormal, raised ANA titer and a clinical correlation with the appropriate autoimmune disorder, extractable nuclear antigens (ENA) panel testing of specific autoantibodies is considered **reimbursable**.
3. For individuals with an initial positive ANA test and a diagnosis of systemic autoimmune rheumatic disease, testing of dsDNA up to four (4) times per year is considered **reimbursable**.
4. For individuals with a negative or low positive ANA test, the following condition specific antibody testing is considered **reimbursable**:
 - a. Testing for anti-Jo-1 in a unique clinical subset of myositis.
 - b. Testing for anti-SSA in the setting of lupus or Sjögren's syndrome.
5. Monitoring of disease with ANA testing or ANA titers is not reimbursable.
6. For individuals without symptoms suggestive of an autoimmune disorder, ANA and/or ENA testing is **not reimbursable**.
7. For all other situations not described above, testing of specific antibodies in the absence of a positive ANA test is **not reimbursable**.
8. For asymptomatic individuals, testing of ANA and/or ENA during a wellness visit or a general exam without abnormal findings is **not reimbursable**.

The following does are **not reimbursable** due to a lack of available published scientific literature confirming that the test(s) is/are required and beneficial for the diagnosis and treatment of an individual's illness.

9. For the diagnosis of RA, testing for serum biomarkers not discussed above, alone or in a panel, is **not reimbursable**.

Coding

Code	Description
CPT	
86038	Antinuclear antibodies (ANA)
86039	Antinuclear antibodies (ANA); titer

Code	Description
86235	Extractable nuclear antigen, antibody to, any method (e.g., nRNP, SS-A, SS-B, Sm, RNP, Sc170, J01), each antibody
0039U	Deoxyribonucleic acid (DNA) antibody, double stranded, high avidity Proprietary test: Anti-dsDNA, High Salt/Avidity Lab/Manufacturer: University of Washington, Department of Laboratory Medicine/Bio-Rad

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Related Information

Table of Terminology

Term	Definition
AAB	Autoantibodies
AAP	American Academy of Pediatrics
ACL	Anticardiolipin
ACP	American College of Pathologists
ACPA	Anti-citrullinated peptide antibodies
ACR	American College of Rheumatology
AIH	Autoimmune hepatitis
AIIF	Automated indirect immunofluorescence
ANA	Antinuclear antibody
Anti La/SS-B	Anti La/Sjögren Syndrome-B
Anti-C1q	Autoantibodies against C1q
Anti-CCP	Anti-cyclic citrullinated peptide
Anti-dsDNA	Anti-double-stranded DNA
Anti-RNP	Antinuclear ribonucleoprotein
Anti-Ro/SS-A	Anti-Ro/Sjögren Syndrome related antigen A autoantibodies
Anti-Sm	Anti-Smith antibodies
APL	Antiphospholipid antibodies

Term	Definition
BC4d	B-lymphocyte-bound C4d
BSR	British Society for Rheumatology
CBC	Complete blood count
CB-CAPs	Cell-bound complement activation products
CCP	Cyclic citrullinated peptides
CDC	Centers for Disease Control and Prevention
CENP	Centromere protein B
CIA	Chemiluminescence immunoassay
CLIA '88	Clinical Laboratory Improvement Amendments of 1988
CMS	Centers for Medicare and Medicaid Services
CRP	C-reactive protein
CTD	Connective tissue diseases
CV	Coefficient of variation
ds	Double-stranded
dsDNA	Double-stranded DNA
EC4d	C4d bound to erythrocytes
eGFR	Estimated glomerular filtration rate
EIA	Enzyme immunoassay
ELISA	Enzyme-linked immunosorbent assay
ENA	Extractable nuclear antigens
ESPGHAN	European Society for Paediatric Gastroenterology Hepatology and Nutrition
ESR	Erythrocyte sedimentation rate
EULAR	European League Against Rheumatism
FEIA	Fluorescence enzyme immunoassay
HEp-2	Human epithelial type 2
ICAP	International Consensus on ANA staining Patterns
IFA	Immunofluorescence assay
IIF	Indirect immunofluorescence
IIMs	Idiopathic inflammatory myopathies
IQ	Interquartile

Term	Definition
ISLM	Italian Society of Laboratory Medicine
JIA	Juvenile idiopathic arthritis
Jo-1	Histidyl t-RNA synthetase
LAC	Lupus anticoagulant
LDT	Laboratory developed test
LE cell	Lupus erythematosus cell
LFA	Lupus Foundation of America
MAP	Multianalyte assay panel
MCTD	Mixed connective tissue disease
MIA	Multiplex immunoassay
MIIF	Manual indirect immunofluorescence
PC	Positive concordance
PMPM	Per member per month
PPPM	Per patient per month
RA	Rheumatoid arthritis
RF	Rheumatoid factor
RNP	Ribonucleoprotein
SARDs	Systemic autoimmune rheumatic diseases
SDI	SLICC damage index
SDLT	Standard diagnosis laboratory testing
SELENA	Safety of Estrogens in Lupus National Assessment
SjS	Sjögren's syndrome
SLE	Systemic lupus erythematosus
SLICC	Systemic Lupus International Collaborating Clinics
SRDs	Systemic rheumatic diseases
SS-B/La	Sjögren's syndrome Type-B
SSc	Systemic sclerosis

Scientific Background

Autoimmune diseases occur when an individual's immune system mistakenly attacks his or her own tissue. This can lead to a variety of conditions and diseases which vary in severity. Autoimmune diseases are estimated to affect five to ten percent of the industrial world population;⁷ autoimmune conditions are associated with increased morbidity and mortality, and are among the leading causes of death (under 65 years) and disability for women in the US.⁸

Systemic lupus erythematosus (SLE) is one of more than 80 recognized autoimmune disorders, affecting approximately 204,000 people in the United States.^{9,10} SLE can present with a wide range of clinical manifestations, typically related to connective-tissue disorders, and often mimics other illnesses.¹¹ This autoimmune disorder leads to inflammation and irreversible damage in one or more organs, including the joints, skin, nervous system, and kidneys.¹² The cause of SLE is not entirely understood, but it is predicted to manifest due to a combination of genetic and environmental factors, such as vitamin D deficiency, sunburn, and/or viral infections.¹³ SLE affects women more than men and is a challenging disease to diagnose because of a broad assortment of signs, symptoms, and serological abnormalities.¹² SLE morbidity can be attributed to both tissue damage, toxic treatments, and complications associated with treatments, such as immunosuppression, long-term organ damage due to corticosteroid therapy, and accelerated coronary artery disease.^{12,14} An early SLE diagnosis is particularly challenging as early-stage tests lack specificity; further, clinical signs and symptoms often only appear after organ damage has occurred, indicating later stages of the disease.¹⁵ SLE diagnoses are made based on lab findings, clinical manifestations, serology, and histology of impacted organs.¹⁵ However, current SLE screening tests are notoriously unreliable.¹⁶

Rheumatoid arthritis (RA) affects more than one million adults in the United States. RA is characterized by chronic inflammation of the synovial tissue of joints, cartilage, and bone.¹⁷⁻²¹ Pathological abnormalities in patients with RA includes chronic synovitis, which results in joint devastation.^{17,18,20} Cellular and humoral response aberrations result in autoimmunity; antibodies and rheumatoid factors against post-translational modified proteins (including modifications such as citrullination). As such, synthetic cyclic citrullinated peptides (CCP) have been developed for diagnostic use.²⁰

There is consensus to the value of serological testing for diagnostic purposes: both rheumatoid factor (RF) and anti-citrullinated peptide antibodies (ACPA) tests have diagnostic value in patients suspected of having RA (but not in asymptomatic patients as a general screen).²² Diagnostic testing with RF should be restricted to those with a moderate to high pretest probability of rheumatoid arthritis. RF testing should not occur in patients with joint pain in the absence of synovitis (e.g., nonspecific arthralgias, fibromyalgia, OA) because a positive test result is more likely to represent a false-positive result. ACPA testing is useful as a diagnostic test in

patients with a moderate to high pretest probability of rheumatoid arthritis, but similarly, should not be used in those with a low pre-test probability. For patients “with an inflammatory, small joint arthritis and with a moderate to high pretest probability of RA, the presence of ACPA testing confirms a diagnosis of RA.”²²

To date, the etiology of RA has not been fully elucidated, though recent studies have suggested that genetic, epigenetic, and environmental factors contribute to RA presentation.^{17,20} Due to the complexity of RA pathogenesis, there is no model drug to cure RA.

Biologic markers or “biomarkers” can provide objective measurements that reflect underlying pathophysiological processes, pathogenic processes, or responses to treatment. Most measures of monitoring disease and treatment progress rely on subjective measurements, such as joint evaluation, so biomarkers may be a useful complement in patient management.²³ Joint damage at the molecular level may be occurring before any clinical signs appear so identifying any indications of disease activity could allow clinical interventions to be taken earlier.²⁴ Markers such as erythrocyte sedimentation rate (ESR) and C-reactive protein (CRP) are part of clinical measures such as the DAS. However, these two biomarkers are nonspecific; abnormal amounts of these markers may be due to other reasons apart from RA and may be completely normal in patients with RA.^{25,26} This non-specificity is not limited to ESR and CRP. For example, antibodies (usually called rheumatoid factors or RF) produced against immunoglobulin G (IgG) are often tested to diagnose RA, but these antibodies may be produced in response to another rheumatic condition or a separate chronic infection.²⁷ Autoantibodies to citrullinated protein epitopes, such anti-cyclic citrullinated peptide (anti-CCP2), has also been a focus of biomarker research in RA. Both RF and anti-CCP2 have similar sensitivities for the diagnosis of RA, but anti-CCP2 is positive in 20%-30% of RA patients who are negative for RF.²⁸ RA is a heterogenous condition, and no single biomarker is a reliable predictor of RA disease activity.²⁴

Currently, conventional synthetic disease modifying anti-rheumatic drugs (csDMARDs) are the first line of RA therapy. Unfortunately, some RA patients do not respond to csDMARDs and clinical guidelines suggest use of alternative therapies such as biologic DMARDs (bDMARDs). bDMARDs are more specific to inflammatory factors than csDMARDs and more efficient in demonstrating remission and inducing low disease activity.²⁹ Several bDMARDs are available for RA management, and these include TNFis.^{17-19,30} TNFi treatment, however, is not without limitations. Unfortunately, the majority of patients fail to respond to TNFi treatment (measured by American College of Rheumatology (ACR)50-indicates 50% disease improvement) and only 10-25% achieve remission.^{17,19,21,31} Currently, there is no way to predict whether RA patients will respond to TNFi therapy, and approximately three months is needed to determine whether a patient is responding.^{17,19} Accordingly, there has been a push to create a personalized medicine approach to identify non-responders to enhance clinical outcomes.^{17,19}

The systems by which the immune system maintains tolerance to an individual's own antigens can be overcome by release of intracellular antigens following excessive cell death, ineffective clearance of apoptotic debris, inflammation-induced modification of self-antigens, or molecular mimicry, leading to the production of antibodies against self-antigens or autoantibodies (AAB).³² Autoantibodies mediate both systemic inflammation and tissue injury and may play a role in the pathogenesis of many autoimmune diseases.³² Generally, AAB development precedes the clinical onset of autoimmune disease and has predictive value;⁴ thus, AABs serve as good serological markers to screen for evidence of autoimmunity.^{33,34} Autoantibodies can target a variety of molecules (including nucleic acids, lipids, and proteins) from many cellular localizations—nucleus, cytoplasm, cell surface, extracellular organelles,³² and different specific AABs are associated with particular diagnoses, symptoms, unique syndromes, subsets of disease, and clinical activity.⁴ See Table 1 from Suurmond and Diamond (2015) below:

Table 1. Autoantibody recognition in systemic autoimmune disease

Antigen location	Antibody	Antigen	Disease	PRR recognition
Nuclear	Anti-Ro (SS-a)	Ro-RNP complex	SLE, Sjögren's syndrome	TLR7
	Anti-La (SS-B)	La antigen	SLE, Sjögren's syndrome	TLR7
	Anti-Sm	Small nuclear RNP	SLE	TLR7
	Anti-dsDNA	dsDNA	SLE	TLR9
	Anti-histone	Histones	SLE (drug-induced)	TLR2 and TLR4
	Anti-Scl-70	Topoisomerase I	Systemic sclerosis	
Cytoplasmic/ mitochondrial	Anti-centromere	Centromere	Systemic sclerosis/CREST syndrome	
	ANCA	Myeloperoxidase (p-ANCA) and proteinase 3 (c-ANCA)	Vasculitis, Wegener's granulomatosis	
Modified proteins	ACA	Cardiolipin	Antiphospholipid syndrome, SLE	NLRP3
	ACPA	Citrullinated proteins	RA	TLR4
	Anti-Carp	Carbamylated proteins	RA	
Extracellular	RF	RF (IgG)	RA	
	Lupus anticoagulant	Phospholipids	Antiphospholipid syndrome	TLR4?
	$\alpha 3$ Chain of basement membrane collagen (type IV collagen)	$\alpha 3$ Chain of basement membrane collagen (type IV collagen)	Goodpasture's syndrome	

ACA, anti-cardiolipin antibody.

However, serum AAB are present in 18.1% of the general population, and titers are higher in females and increase with age.³⁵ Additionally, only in a few cases does the antibody titer correlates with the severity of clinical manifestations or the response to treatment.³⁴ The use of ANA detection as a diagnostic test originated with the observation of the lupus erythematosus (LE) cell.³⁶ Since then, several tests have been developed to detect these antibodies.

The indirect immunofluorescence (IIF) test is the most widely used assay for the detection of AAB and remains the reference method of choice.³⁷ Detection of ANAs by the IIF technique demonstrates binding to specific intracellular structures within the cells, resulting in staining patterns reported using the consensus nomenclature and representative patterns defined by The International Consensus on ANA staining Patterns (ICAP) initiative and the degree of binding reflected by the fluorescence intensity or titer.^{5,38} The test takes advantage of a HEp-2 cell line, which have large, easy to visualize, nuclei and contain nearly all of the clinically important autoantigens, making these cells ideal for the detection of the corresponding AABs.³⁹ The ANA

IIF assay using HEp-2 slide has a high sensitivity for screening of SARDs and efforts to harmonize the nomenclatures for testing and reporting have made this a powerful screening tool.⁵ The frequency of ANA in SLE and SSc is 95–100%, 50–70% in SJS and 30–50% in rheumatoid arthritis (RA);⁴ however, their isolated finding in an otherwise healthy individual has a low positive predictive value which needs to be integrated with other laboratory parameters and patient risk factors.³⁵ Disadvantages of the indirect immunofluorescence test include its labor-intensiveness, significant training requirements for competence, and subjectivity in titer and pattern recognition; moreover, because the staining pattern usually does not identify the responsible autoantibody, additional testing may be required.^{5,39} Automated image analysis provides a viable option for distinguishing between positive and negative results although the ability to assign specific patterns is insufficient to replace manual microscopic interpretation.⁴⁰

The antinuclear antibody (ANA) test is commonly used in the evaluation of autoimmune disorders, as these antibodies are responsible for attacking healthy or normal cells. More than 95% of individuals with SLE will have a positive ANA test.¹⁶ However, ANAs are present in “a significant proportion of normal individuals and lacks specificity or prognostic value.”¹⁵ In particular, approximately only 11-13% of individuals with a positive ANA test will actually have SLE, and approximately 15% will be completely healthy.¹⁶ Other SLE diagnostic methods include the monitoring of anti-double-stranded DNA (anti-dsDNA), C3 and C4 complement levels, CH50 complement levels, erythrocyte sedimentation rate (ESR) and/or C-reactive protein (CRP) levels, antiphospholipid antibodies, and urine protein-to-creatinine ratios.⁴¹

If SLE is suspected based on the clinical picture following a positive ANA screen, the sera should be tested for antibodies to double-stranded DNA (dsDNA). Anti-dsDNA antibodies are present in two-thirds of patients with SLE, and they have a good association with disease activity and lupus nephritis. Serial monitoring of anti-dsDNA antibodies has modest correlation with disease activity.³³

A positive ANA screen should also be followed by identification of sub-specificities by screening for antibodies to extractable nuclear antigens (ENAs). ENAs were identified by using saline extract of nuclei as the antigen. Antibodies to ENA can be determined using double immunodiffusion, immunoblotting, enzyme-linked immunosorbent assays (ELISA), or bead-based assay using recombinant or affinity-purified antigens. Different ENAs have an association with different connective tissue diseases.³³

Reflex tests for positive ANA screens have been proposed to improve appropriateness in diagnosis of SARDs and avoid unnecessary second level testing. For specific autoantibodies responsible for certain fluorescent ANA patterns, such as homogeneous, speckled, fine grainy (Scl70-like), nucleolar, centromeric or speckled cytoplasmic, the identification of precise autoantibody markers is considered essential while for others it is not deemed to be necessary.⁴² See Table 1 from Tonutti, et al. (2016), below.

Table 1ANA-reflex test procedure with titres $\geq 1:160$ and typical patterns

ANA-IIF pattern on HEp-2 cells	Reflex test(s)
Nuclear homogeneous $\geq 1:160$	Antibodies to intracellular specific antigens (ENA) and to dsDNA/nucleosomes
Nuclear speckled $\geq 1:160$	Anti-dsDNA and antibodies to intracellular specific antigens (ENA), possibly including anti-RNA polymerase III
Nuclear Scl70-like $\geq 1:160$	Antibodies to intracellular specific antigens (ENA) (possibly including anti-PM/Scl)
Cytoplasmic speckled $\geq 1:160$	Antibodies to intracellular specific antigens (ENA), including anti-tRNA synthetases and anti-P ribosomal
Pleomorphic PCNA-like (any titre)	Anti-PCNA
Centromere	No confirmation necessary if high titres. Execute specific test for anti-CENP B only in dubious cases (low titre or centromeric pattern not clearly recognizable)

ENA includes SS-A/Ro52 and Ro60, SS-B/La, Sm, RNP, Jo-1, and Scl70

Proprietary Testing

A set of proprietary tests are available from Exagen, under the "AVISE" line. Their line of tests utilizes a two-tiered testing method and a novel algorithm that measures 10 SLE relevant markers to deliver an index calculation value suggestive of the presence or absence of SLE. This includes tests for prognosis (10 biomarkers including various autoantibodies such as anti-C1q and antiribosomal P), diagnosis (10 biomarkers, includes ENA panel), and monitoring (six biomarkers, includes anti-dsDNA and anti-C1q). AVISE CTD (standing for connective tissue disease) is intended to assist with the differential diagnosis of several autoimmune diseases and includes several ANA biomarkers, as well as an ENA panel. Other tests offered, such as AVISE Anti-CarP (evaluates autoantibodies to carbamylated proteins for rheumatoid patients) still include ANA components.⁴³

AVISE Lupus by Exagen is a laboratory developed test (LDT) designed to assist in SLE diagnoses. This LDT utilizes a two-tiered testing method and a novel algorithm that measures 10 SLE relevant markers to deliver an index calculation value suggestive of the presence or absence of SLE. The AVISE Lupus test also uses cell-bound complement activation products (CB-CAPs) to measure complement system activation.⁴⁴ The 10 SLE relevant markers in this test include anti-dsDNA, anti-Smith (anti-Sm) antibodies, erythrocyte-bound C4d or B-lymphocyte-bound C4d (BC4d), ANA, CB-CAPs, and autoantibody specificity components.⁴⁴ As noted on their website, "The AVISE Lupus test is an ideal test for ANA positive patients with a clinical suspicion of lupus."⁴⁴

PrismRA is a molecular signature test that predicts TNFi non-response prior to treatment initiation. PrismRA utilizes a 23-feature blood-based molecular signature response classifier (MSRC) which integrates next generation RNA sequencing data and clinical features (clinical metrics, demographic variables, C reactive protein (CRP) and anti-CCP antibodies) to predict patients' response to TNFi treatment.²¹ A high score is indicative of decreased likelihood of the RA patient to respond to TNFi therapies.

Vectra DA is a multi-biomarker disease activity (MBDA) blood test which combines the levels of 12 serum biomarkers into a single score from 1 to 100 to provide an objective measure of RA

disease activity. It is intended for use with existing symptom-based disease activity measures to improve long-term outcomes for RA patients.⁴⁵ While multi-biomarker panels are emerging as a potentially useful tool in the management of RA, there is not yet a consensus as to their clinical utility.²³

The proprietary test, aiSLE DX Disease Activity Index, is a blood test that measures the level of disease activity and supports clinicians in the measurement of lupus (SLE) and the assessment of treatments. This test examines a well-defined group of immune modulatory soluble mediators, such as cytokines, chemokines, and soluble receptors, which have been linked to disease activity in plasma. The panel of immune mediators assessed in this test are able to differentiate individuals with active clinical illness from those with low activity or quiescent disease in a simple blood test.⁴⁶

Seronegative Rheumatoid Arthritis Profile is a blood test which includes Rheumatoid Factor (RF) IgG, IgA, IgM, Antibodies to Cyclic Citrullinated Peptide (CCP)- IgG, and Scavenger Receptor A (SR-A) which helps to identify antinuclear antibodies. This test is seen as a more robust serological diagnostic panel for RA.⁴⁷

The Early Sjögren's Syndrome Profile from Immco Diagnostics Inc. is a blood test that combines traditional and novel markers for early diagnosis with higher sensitivity and specificity. These novel markers include SP-1, CA-6, and PSP which helps to increase the sensitivity to over 80% and enables diagnosis before the traditional markers, SS-A (Ro) and SS-B (La), appear.⁴⁸

Analytical Validity

A variety of manual or automated single or multiplex immunoassays have been introduced to make the process of detecting autoantibodies more efficient, including ELISA, fluorescent microsphere assays, and chemiluminescence immunoassays (CIA)—each with different performance characteristics.⁵ In these assays, a panel of purified native or recombinant autoantigens is prepared, and each antigen is immobilized on a solid surface (microtiter plate, fluorescent microsphere, or membrane) and incubated with diluted human serum.³⁹ The advantages of these alternative approaches to ANA IIF testing include their suitability for high-throughput testing, semi-quantification of test results, the lack of subjectivity, and the consolidation of ANA-related tests in a single platform as a positive test also provides identification of the responsible autoantibody.^{5,39} It has been estimated that solid phase assays may decrease the labor cost of ANA testing by as much as 95%.³⁹ In a recent study which evaluated the performance of an automated CIA and fluorescence enzyme immunoassay (FEIA) and compared their performance to that of IIFA, both FEIA and CIA screen significantly outperformed IIF, with a higher specificity for FEIA and higher sensitivity for CIA.⁴⁹ The use of solid phase assays as the initial test for the detection of ANA is concerning because the number of autoantigens that are included in solid phase assays is limited compared with the number

that are present in the Hep-2 cell substrate, thus limiting sensitivity.³⁹ Consequently, IIF remains the gold standard, and in cases of strong clinical suspicion of SARD and a negative screen from a solid phase assay, IIF should be performed.⁴⁹

Tipu and Bashir (2018) investigated the specificity and pattern for ANA in systemic rheumatic disease patients. A total of 4347 samples were sent, and 397 were positive for ANA. Of these 397, 96 were positive on the anti-ENA screen and tested for anti-ENA reactivity. Anti-SSA antibodies were found in 59 of these samples. The most common ANA patterns were "coarse" and "fine-speckled" (43 and 22 of 81 respectively). However, no specific ANA pattern was associated with anti-ENA reactivity.⁵⁰

Kim et al. (2019) performed a meta-analysis comparing ANA measurement by automated indirect immunofluorescence (AIIF) and manual indirect immunofluorescence (MIIF). A total of 22 studies including 6913 positive and 1818 negative samples of manual indirect immunofluorescence (MIIF) were included. Among this cohort, 524 samples with combined systemic rheumatic diseases (SRDs), 132 systemic lupus erythematosus (SLE) samples, and 104 systemic sclerosis (SSc) samples, and 520 controls were available. Positive concordance (PC) between AIIF and MIIF was 93.7%, although PC of total pattern and titer were lower. Clinical sensitivities of AIIF vs MIIF were 84.7% vs 78.2% for combined SRDs, 95.5% vs 93.9% for SLE, and 86.5% vs 83.7% for SSc. Clinical specificities of AIIF vs MIIF were 75.6% vs 79.6% for combined SRDs, 74.2% vs 83.3% for SLE, and 74.2% vs 83.3% for SSc. The authors concluded that the sensitivities did not differ between methods, but the specificities of SLE and SSc were statistically significant changes.⁵¹

Dervieux, et al. (2017) performed the analytical validation of Exagen's multianalyte panel test for SLE. This assay uses quantitative flow cytometry to assess the levels of the complement split product C4d bound to erythrocytes (EC4d) and B-lymphocytes (BC4d), in units of mean fluorescence intensity (MFI), and immunoassays to assay for antinuclear and anti-double stranded DNA antibodies (e.g. autoantibodies). The results were reported on a two-tiered index score as either positive or negative. The authors included specimens from both patients with SLE as well as individuals without SLE. Controls consisting of three-level C4 coated positive beads were run daily. The authors note that at ambient temperature both EC4d and BC4d are stable for two days and for four days if the samples are stored at four degrees Celsius. "Median intra-day and inter-day CV [coefficient of variation] range from 2.9% to 7.8% (n=30) and 7.3% to 12.4% (n=66), respectively. The 2-tiered index score is reproducible over 4 consecutive days upon storage of blood at 4°C. A total of 2,888 three-level quality control data were collected from six flow cytometers with an overall failure rate below 3%. Median EC4d level is six net MFI (Interquartile [IQ] range 4-9 net MFI) and median BC4d is 18 net MFI (IQ range 13-27 net MFI) among 86,852 specimens submitted for testing. The incidence of 2-tiered positive test results is 13.4%."⁵²

Putterman, et al. (2014) compared the performance of C4d CB-CAPs on erythrocyte and B cells with antibodies to dsDNA, C3, and C4 in patients with SLE. A total of 794 individuals participated in this study, which included 205 healthy controls, 304 patients with SLE, and 285 patients with other rheumatic diseases. Both erythrocytes and B cells were measured with flow cytometry, and antibodies, including anti-dsDNA, were measured with solid-phase immunoassays. SLE activity was determined using the SLE Disease Activity Index Safety of Estrogens in Lupus National Assessment (SELENA) Modification, and the two-tiered AVISE Lupus test was developed. Results showed that "The combination of EC4d and BC4d in multivariate testing methodology with anti-dsDNA and autoantibodies to cellular and citrullinated antigens yielded 80% sensitivity for SLE and specificity ranging from 70% (Sjogren's syndrome) to 92% (rheumatoid arthritis) (98%vs. normal)." ⁵³ Overall, the measurement of CB-CAPs was more sensitive for SLE diagnostic purposes than complement or anti-dsDNA measurements.

Ramsey-Goldman, et al. (2020) evaluated the use of CB-CAPs, using flow cytometry, or a multianalyte assay panel (MAP) that includes CB-CAPs (e.g., AVISE Lupus) on patients with suspected SLE (n = 92) who fulfilled three classification criteria of the American College of Rheumatology (ACR). They also compared the data with individuals with established SLE (n = 53). At the initial visit, the individuals with suspected SLE had statistically higher positive CB-CAP (28%) or MAP results (40%) than individuals with established SLE. "In probable SLE, MAP scores of >0.8 at enrollment predicted fulfillment of a fourth ACR criterion within 18 months (hazard ratio 3.11, P<0.01)." The authors, who did acknowledge compensation from Exagen, conclude that "[a] MAP score above 0.8 predicts transition to classifiable SLE according to ACR criteria." ⁵⁴

Clinical Utility and Validity

ANA, ENA, and SCLT

Oglesby, et al. (2014) performed a cost-savings impact analysis on when the diagnosis of SLE is made and how it affects the clinical and economic outcomes. Using a claims database of claims made between January 2000 and June 2010, the authors separated individuals into two groups (n = 4166 per group) —early diagnosis (within six months of onset of symptoms) and late diagnosis (6 or more months after the onset of symptoms)—based upon an algorithm using a patient's ICD-9 diagnosis code(s) on the claim(s) and when SLE medications were dispensed. Additional propensity scores were matched using data based on "age, gender, diagnosis year, region, health plan type, and comorbidities." Results show that the early diagnosis group had lower rates of mild, moderate, and severe flares as well as lower rates of hospitalization as compared to the late diagnosis group. Moreover, "[c]ompared with the late diagnosis patients, mean all-cause inpatient costs PPPM [per patient per month] were lower for the early diagnosis patients (US\$406 vs. US\$486; p = 0.016). Corresponding SLE-related hospitalization costs were also lower for early compared with late diagnosis patients (US\$71 vs US\$95; p = 0.013)." The

values are adjusted to 2010 US dollars. The authors note that the other resource use and cost categories were consistent, concluding "[p]atients diagnosed with SLE sooner may experience lower flare rates, less healthcare utilization, and lower costs from a commercially insured population perspective."⁵⁵

A study by Yeo, et al. (2020) demonstrates that there is little benefit to repeat ANA testing if the initial test was negative by evaluating the cost of repeat ANA testing. From 2011 to 2018, 36,715 ANA tests were performed for 28,840 patients at a total cost of \$675,029. Of these tests, 21.4% were repeats in which 54.9% of the patients initially tested negative. Of those who tested negative and repeated ANA testing, only 19% of the patients had a positive result when the test was repeated once in under two years, and this positive test did not lead to a change in diagnosis. Therefore, the authors conclude that "repeat ANA testing after a negative result has low utility and results in high cost."⁵⁶

Deng, et al. (2016) investigated the clinical utility of ANA testing through different assays to see which one was most appropriate for evaluating patients with CTD. With 1000 samples collected, they compared an enzyme immunoassay (EIA), immunofluorescence assay (IFA), and multiplex immunoassay (MIA) in terms of specificity and sensitivity of testing. The researchers found that through using weights to define a patient sample that reflected the intended testing population and a normalized specificity of 90% to standardize the comparison between tests, the MIA, EIA, and IFA had sensitivities of 67%, 67%, and 56%, respectively. However, with a varying clinical cutoff, the IFA could obtain a sensitivity of 94% and a corresponding specificity of only 43%. This demonstrated that the sensitivity and specificity could easily vary with predetermined cutoffs; but, there were "no statistically significant differences in the clinical utility of the IFA, EIA, or MIA."⁵⁷

Alsaed, et al. (2021) compared the performance of ANA testing via ELISA vs IIF for CTDs. From a sample of 1457 patients and 12,439 tests ordered in 2016, they found that with "cut-off ratio ≥ 1.0 for ANA-ELISA and a dilutional titre $\geq 1:80$ for ANA-IIF, the sensitivity of ANA-IIF and ANA-ELISA for all CTDs were 63.3% vs 74.8% respectively. For the SLE it was 64.3% vs 76.9%, Sjogren's Syndrome was 50% vs 76.9% respectively. The overall specificity of ANA-ELISA was 89.05%, which was slightly better than ANA-IIF 86.72%." This communicated the ELISA was slightly better than IIF in sensitivity and specificity, which could influence the convention of using IIF going forward if these findings are reflected in other cohort studies.⁵⁸

Biomarker analysis

Wallace, et al. (2019) performed a randomized prospective trial to assess the clinical utility of the AVISE lupus MAP test (MAP/CB-CAP) as compared to standard diagnosis laboratory testing (SDLT). A total of 145 patients with a history of positive antinuclear antibody status were randomly assigned to either an SDLT arm ($n = 73$) or the MAP/CB-CAP arm ($n = 72$) of the study.

Treatment changes were recorded based on either the SDLT or MAP/CB-CAP results. Even though the demographics between the two arms of the study were similar, the results were different. "Post-test likelihood of SLE resulting from randomisation in the MAP/CB-CAPs testing arm was significantly lower than that resulting from randomisation to SDLT arm on review of test results (-0.44 ± 0.10 points vs -0.19 ± 0.07 points) and at the 12-week follow-up visit (-0.61 ± 0.10 points vs -0.31 ± 0.10 points) ($p < 0.05$). Among patients randomised to the MAP/CB-CAPs testing arm, two-tiered positive test results associated significantly with initiation of prednisone ($p = 0.034$)."⁵⁹ The authors conclude that testing such as the AVISE Lupus test has clinical utility and does affect treatment decisions.

A longitudinal, retrospective study by Mossell, et al. (2016) of 46 patients who were anti-nuclear antibodies (ANA) positive but SLE-specific autoantibodies negative was conducted to evaluate the clinical utility of the AVISE Lupus test. There was a total of 23 patients in the "case" group (i.e. positive result based on the AVISE Lupus test), and 23 patients were in the "control" or negative results group. The charts of each individual were reviewed at two different times: T0 (or the initial time) and T1 (or approximately one year later). The case group was diagnosed with SLE at a higher rate than the control group (87% vs. 17%, respectively); moreover, the case group fulfilled four of the ACR classification criteria of SLE at a higher rate than the control group (43% vs 17%, respectively). The authors found that the sensitivity of the AVISE Lupus test (83%) is statistically significantly higher than the ACR score (42%, $p = 0.006$). Even at the initial baseline, individuals in the case group were prescribed anti-rheumatic medications more frequently (83% vs. 35%, $p = 0.002$) than the control group, indicating that a positive AVISE Lupus test may result in a more aggressive early treatment therapy.⁶⁰

Liang, et al. (2020) assayed the utility of the AVISE test in predicting lupus diagnosis and progression in 117 patients who previously did not have a diagnosis of SLE. The study assessed the patients at the time of the initial AVISE test ($t = 0$) and two years later ($t = 2$) using the SLE diagnosis criteria of the Systemic Lupus International Collaborating Clinics (SLICC) and ACR and the SLICC Damage Index (SDI) to measure SLE damage. After two years, patients who tested positive developed SLE at a significantly higher rate than those who tested negative using the AVISE test (65% vs 10.3%, $p < 0.0001$). AVISE-positive patients have more SLE damage after two years than AVISE-negative patients (1.9 ± 1.3 vs 1.03 ± 1.3 , $p = 0.01$). In particular, the authors note that the levels of BC4d "correlated with the number of SLICC criteria at $t=0$ ($r=0.33$, $p < 0.0001$) and $t=2$ ($r=0.34$, $p < 0.0001$), as well as SDI at $t=0$ ($r=0.25$, $p=0.003$) and $t=2$ ($r=0.26$, $p=0.002$)."⁶¹

Alexander, et al. (2021) further validated the clinical utility of the AVISE lupus test via a systematic review of medical records of ANA-positive patients with positive (>0.1) or negative (<-0.1) MAP scores. They found that the "odds of higher confidence in SLE diagnosis increased by 1.74-fold for every unit increase of the MAP score" with statistical significance, demonstrating

that the test still further solidifies a diagnosis of SLE and can help inform “appropriate treatment decisions.”⁶²

A study by Clarke, et al. (2020) demonstrates the cost-effective management of systemic lupus erythematosus (SLE) using a MAP rather than SDLTs. The higher specificity of MAP allows for an earlier SLE diagnosis, prompt initiation of the appropriate therapy, and fewer unnecessary and costly hospitalizations or investigations. Current SDLTs, such as ANA tests, have a high diagnostic sensitivity, but a high false-positive rate. MAP combines complement C4d activation products on erythrocytes and B cells with SDLTs, with antibodies to nuclear antigens, dsDNA IgG (with Crithidia confirmation), Smith, Sjogren’s syndrome type-B (SS-B/La), topoisomerase I (Scl-70), centromere protein B (CENP), histidyl t-RNA synthetase (Jo-1), and cyclic citrullinated peptides (CCP) to improve SLE diagnosis. MAP “yields improved overall diagnostic performance with a sensitivity and specificity of 80% and 86%, respectively, compared with a sensitivity and specificity of 83% and 76%, respectively, for SDLTs. Despite the lower sensitivity, the superior specificity of MAP (86%) over SDLTs (76%) results in a higher positive predictive value associated with MAP (36.75%) compared with SDLTs (26.02%).”⁶³ The improved specificity of MAP resulted in a cost savings of \$1,991,152 to a US commercial plan over a four year time horizon, which translates to \$0.04 in per member per month (PMPM) savings.⁶³

Clinical validation of PrismRA was conducted in the Comparative Effectiveness Registry to Study Therapies for Arthritis and Inflammatory Conditions (CERTAIN) study.^{30,64} The CERTAIN trial was conducted by the Consortium of Rheumatology Researchers of North America which consisted of 43 sites and 117 rheumatologists.⁶⁴ This prospective study analyzed baseline RNA sequencing and clinical assessments to determine the effectiveness of PrismRA to predict TNFi non-response. Evaluation of the clinical response to TNFi was performed at six months and was determined by ACR50. The CERTAIN study built and validated the biomarker panel used for MSRC analyses. The study found that PrismRA demonstrated a positive predictive value of 89.7%, a specificity of 86.8%, and a sensitivity of 50%.^{19,64}

Inadequate TNFi response predictions were further validated on integrated blood samples from CERTAIN and NETWORK-004 studies. NETWORK-004 was a 24-week blinded prospective study conducted at 73 sites to evaluate the ability of MSRC to identify TNFi non-responders at three and six months by ACR50 (evaluations were also conducted using other scales such as Disease Activity Score (DAS28)-CRP, and Clinical Disease Activity Index). CERTAIN samples were used for transcript biomarker feature selection (n=100) and cross validation of MSRC (n=245). In the NETWORK-004 cohort, MSRC validation was performed in samples from naïve (n=146) and TNFi exposed (n=113) patients. ACR50 of patients stratified by MSRC at six months according to prediction of an inadequate response to TNFi therapy had an odds ratio of 4.1 (95% CI 2.0–8.3; p value=0.0001). Patients with a non-response MSRC were 26 times less likely to achieve remission evaluated three months after TNFi therapy.²¹ Both studies found that PrismRA was able to

accurately predict TNFi non-responders according to multiple clinically validated measurement scales.^{21,64}

Bergman, et al. (2020) performed modeling of the projected improvements from PrismRA and determined that ACR50 improved in the stratified cohort (40%) compared to the unstratified patient cohort (30%) and decreased costs of ineffective treatment by 19%. Further, PrismRA was shown to be a better predictor of inadequate response to TNFi treatment than clinical metrics alone.³⁰ Pappas, et al. (2021) conducted a 32-question decision-impact survey involving 248 rheumatologists to determine whether predictive tests such as PrismRA appear to have clinical utility in RA patients' ability to respond to TNFi therapy. The study demonstrated that rheumatologists overwhelmingly supported the clinical need of predictive technologies to determine whether RA patients would respond to TNFi therapies and that payers should provide coverage of predictive technology.¹⁹

According to Curtis, et al. (2012), the MBDA algorithm (Vectra DA) was developed by screening 396 candidate biomarkers. An algorithm was then created to generate a composite score based on the 12 biomarkers most correlated to RA clinical disease activity which are as follows:

- Interleukin-6 [IL-6]
- Tumor necrosis factor receptor type I [TNFRI]
- Vascular cell adhesion molecule 1 [VCAM-1]
- Epidermal growth factor [EGF]
- Vascular endothelial growth factor A [VEGF-A]
- YKL-40
- Matrix metalloproteinase 1 [MMP-1]
- MMP-3
- CRP
- Serum amyloid A [SAA]
- Leptin
- Resistin

These biomarkers represent several processes related to RA, such as cartilage remodeling and cytokine signaling pathways. A score of ≤ 29 is considered "low" activity, between 29 and 44 is "moderate" activity, and > 44 is "high" activity. The MBDA is intended to provide separate information from a clinical evaluation of joints and should be used as a complement, not as a replacement.²⁵

This MBDA has been shown to correlate significantly ($r=0.72$; $p<0.001$) with a disease activity score based on the 28-joint Disease Activity Score based on CRP (DAS28-CRP) and has been validated for clinical use as a disease activity marker in RA.²⁵ Both Hirata, et al. (2013) and Bakker, et al. (2012) found the MBDA score to correlate well with disease activity and could

complement other existing measures of RA assessment. Remission based on the MBDA score was a significant predictor of radiographic non-progression, whereas both remission-defined DAS28-CRP and American College of Rheumatology/European League Against Rheumatism (ACR/EULAR) criteria was not. The MBDA test was also useful in assessing the risk of radiographic progression among patients who met clinical remission criteria. MBDA results may provide an important addition to clinical assessment, however, further studies are needed to confirm its clinical utility in the management of RA.⁴⁵

Li, et al. (2013) evaluated the impact of an MBDA blood test for rheumatoid arthritis (RA) on treatment decisions made by six health care providers (HCPs) in 101 patients. HCPs completed surveys before and after viewing the MBDA test result, recording dosage and frequency for all RA medications and assessment of disease activity. Frequency and changes in treatment plan that resulted from viewing the MBDA test result were determined. The MBDA test results were found to have changed 38% of patients' treatment plans. Furthermore, treatment plans were changed 63% of the time the MBDA test results were found to be "not consistent" or "somewhat consistent" with the clinical assessment of disease activity. However, any improvement in clinical outcomes caused was not reported, and the overall amount of drug use was not affected.⁶⁷

Another study by Li, et al. (2016) assessed the correlation between MBDA score and disease progression in 163 RA patients. The study found that low radiographic progression was associated with low MBDA scores, and higher scores were associated with more frequent and severe progression. Notably, MBDA scores correlated with progression even when a conventional measure such as the DAS28 indicated otherwise. For example, low risk of progression was associated with a low MBDA score, even when a concurrent DAS28 score was high. The authors concluded that MBDA may be a good complement for conventional measures, as well as provide information on changing treatment plans.⁶⁸

Curtis, et al. (2018) initially studied the influence of age, obesity, and other comorbidities on the MBDA test. A cross-sectional analysis of RA patients who have participated in an MBDA test was used (n=357). "Of 357 eligible patients, 76% (n = 273) had normal CRP (<10mg/L) with high (33%), moderate (45%), and low (22%) disease activity by MBDA. The MBDA score was significantly associated with BMI, age, CDAI [clinical disease activity index], and SJC [swollen joint count]."⁶⁹ Almost one third of participants had normal CRP scores but high MBDA scores. "In this real-world analysis, the MBDA score was associated with RA disease activity, obesity, and age, and was negligibly affected by common comorbidities."⁶⁹ The authors conclude by suggesting that an adjusted MBDA score may require development to account for BMI and age. Such a study was then published the following year. Curtis, et al. (2019) developed an MBDA test that will include additional factors such as sex, age and obesity in RA patients. Obesity, or adiposity, was measured using either BMI or serum leptin concentration. Two cohorts were studied, totaling 1736 patients. Overall, the authors have developed "a leptin-adjusted MBDA score that

has significantly improved [the] ability to predict clinical disease activity and radiographic progression."⁷⁰ It was suggested that this leptin-adjusted MBDA score "significantly adds information to DAS28-CRP and the original MBDA score in predicting radiographic progression. It may offer improved clinical utility for personalized management of RA."⁷⁰

A recent study analyzed the measurement of serum biomarkers at early RA disease onset in hopes to better predict disease progression.⁷¹ MBDA score and changes in this score were evaluated to predict DAS28-CRP remission. A total of 180 patients participated in this study and were treated with either methotrexate and adalimumab (n = 89) or methotrexate and placebo (n = 91) in addition to a glucocorticoid injection into swollen joints; results showed that "Early changes in MBDA score were associated with clinical remission based on DAS28-CRP at 6 months."⁷¹

In a study by Ma, et al. (2020), the MBDA test was used to explore the role of biomarkers in predicting remission of RA. Serum samples for 148 patients were assessed for MBDA score at three months, six months, and at one year. RA patients on greater than six months stable therapy in stable low disease activity were assessed every three months for one year. Patients not fulfilling any remission criteria at baseline were classified as 'low disease activity state' (LDAS). Patients not fulfilling any remission criteria over one year were classified as 'persistent disease activity' (PDA). Of the 148 patients, 27% were in the LDAS group and over one year and 9% of patients were classified as PDA. Baseline MBDA score and concentrations of IL-6, leptin, SAA and CRP were significantly lower in all baseline remission criteria groups in comparison to LDAS groups. The individual MBDA biomarkers (IL-6, leptin, SAA, CRP) and initial MBDA score was able to differentiate between remission at baseline and LDAS. The authors state that these findings highlight the potential value of repeated measurements of MBDA score to evaluate the stability of clinical disease activity over time.⁷²

In a combined analysis of the OPERA, SWEFOT, and BRASS studies in which a newer version of the MBDA score was validated, Curtis analyzed the prognostic value of the adjusted MBDA score for radiographic progression in RA. The new MBDA score, used in these three studies, adjusts for age, sex, and adiposity. Curtis evaluated associations of radiographic progression (Δ TSS) per year with the adjusted MBDA score, seropositivity, and clinical measures using linear and logistic regression. The adjusted MBDA score was validated in SWEFOT, compared with the other two cohorts, and used to generate curves for predicting risk of radiographic progression. The adjusted MBDA score was found to be the "strongest, independent predictor of radiographic progression (Δ TSS > 5) compared with seropositivity (rheumatoid factor and/or anti-CCP), baseline TSS, DAS28-CRP, CRP SJC, or CDAI. Its prognostic ability is not significantly improved by the addition of DAS28-CRP, CRP, SJC, or CDAI."⁷³

Fleischmann, et al. (2022) engaged in a multicenter, randomized, placebo-controlled trial of repository corticotropin injection (RCI) in patients with active RA. The utility of an MBDA score

was measured against the utility of the Disease Activity Score to assess disease activity in RA. Study participants received 80 units of RCI twice weekly, and those who had low disease activity at week 12 were given either 80 units of RIC or a placebo twice weekly. The changes in disease activity (measured by DAS28-ESR, CDAI, and MBDA scores) were analyzed, including correlations between MBDA scores and both DAS28-ESR and CDAI scores. Results showed “changes from baseline in DASw8-ESR and CDAI scores suggested the RCI therapy led to clinically meaningful improvements in disease activity, but improvements from baseline in MBDA scores were below the minimally important difference threshold.” The authors concluded that MBDA scores were not “sufficiently responsive” in the assessment of RA disease activity. The authors also said that MBDA should not be used as a preferred disease activity measure for RA patients.⁷⁴

Guidelines and Recommendations

American College of Rheumatology

Systemic Lupus Erythematosus (SLE)

In 1997, the Diagnostic and Therapeutic Criteria Committee of the ACR revised the 1982 criteria for SLE. Often referred to as the 1997 ACR criteria, these revisions included the addition of “[p]ositive finding of antiphospholipid antibodies based on 1) an abnormal serum level of IgG or IgM anticardiolipin antibodies, 2) a positive test result for lupus anticoagulant using a standard method, or 3) a false-positive serologic test for syphilis known to be positive for at least six months and confirmed by Treponema pallidum immobilization or fluorescent treponemal antibody absorption test.”⁷⁵ The 1997 ACR criteria consists of 11 possible different criterion and each criterion may have more than one definition. A minimum score of four out of 11 is indicative of SLE. According to the Centers for Disease Control and Prevention (CDC), rheumatologists can use these criteria “to classify SLE for research purposes.”⁷⁶ The 1997 ACR criteria in a study by Mosca, et al. (2019), using a cohort of 616 patients, has a reported accuracy of 75.5%, sensitivity of 66.1%, and specificity of 91.6%. The criteria are as follows:^{76,78}

- Malar Rash
- Discoid Rash
- Photosensitivity
- Oral Ulcers
- Nonerosive Arthritis
- Pleuritis or Pericarditis
- Renal Disorder
- Neurologic Disorder
- Hematologic Disorder
- Immunologic Disorder

- Positive Antinuclear Antibody

The ACR published a statement on the Methodology of Testing for Antinuclear Antibodies which states:³⁷

- The ACR supports the immunofluorescence antinuclear antibody (ANA) test using Human Epithelial type 2 (HEp-2) substrate, as the gold standard for ANA testing.
- Hospital and commercial laboratories using alternative bead-based multiplex platforms or other solid phase assays for detecting ANAs must provide data to ordering healthcare providers on request that the alternative assay has the same or improved sensitivity compared to IF ANA.
- In-house assays for detecting ANA as well as anti-DNA, anti-Sm (anti-Smith antibodies), anti-RNP (antinuclear ribonucleoprotein), anti-Ro/SS-A (anti-Ro/Sjögren Syndrome-A), anti-La/SS-B (anti-La/Sjögren Syndrome-B), etc., should be standardized according to national (e.g., CDC) and/or international (e.g., WHO, IUIS) standards.
- Laboratories should specify the methods utilized for detecting ANAs when reporting their results.

The above positions were reaffirmed in 2019.⁷⁹

The ACR, together with "Choosing Wisely" also developed a list of five tests, treatments or services that are commonly used in rheumatology practice, but their value should be questioned. The ANA testing was the first on the final top five items list with level of evidence Grade 1C. In their review, the Task Force considered recommendations currently published by American College of Pathologists (ACP), ACR, and Italian Society of Laboratory Medicine (ISLM). They have issued the following recommendation: "Do not test antinuclear antibody (ANA) subserologies without a positive ANA and clinical suspicion of immune-mediated disease."⁸⁰ For their list of five things to question for pediatric rheumatology, two points pertain to ANA testing.⁸¹ "Do not order autoantibody panels unless positive ANAs and evidence of rheumatic disease. There is no evidence that autoantibody testing (including ANA and autoantibody panels) enhances the diagnosis of children with musculoskeletal pain in the absence of evidence of rheumatic disease as determined by a careful history and physical examination." The latter recommendation also stated, "Do not repeat a confirmed positive ANA in patients with established JIA [juvenile idiopathic arthritis] or SLE."⁸¹ These guidelines were reviewed and reaffirmed in 2021.

Rheumatoid Arthritis

In 2021, the ACR released an updated guideline on the management of rheumatoid arthritis, including new recommendations for high-risk groups. Pertaining to disease management and the risk of hepatotoxicity associated with methotrexate therapy, the ACR notes that "the use of methotrexate should be restricted to patients with normal liver enzymes and liver function tests

without evidence of liver disease or liver fibrosis.” No multi-biomarker tests or disease activity tests (such as Vectra DA or PrismRA) were mentioned in the guideline for diagnostic or disease management indications.⁸²

European League Against Rheumatism/American College of Rheumatology (EULAR/ACR)

Systemic Lupus Erythematosus (SLE)

The EULAR/ACR published a joint guideline to develop new classification criteria for systemic lupus erythematosus (SLE). In it, they stated that antinuclear antibodies (ANA) “at a titer of $\geq 1:80$ on HEp-2 cells or an equivalent positive test” was to be an “entry criterion”: if absent, the condition is not SLE; if present, apply additive criteria such as leukopenia or oral ulcers. Antiphospholipid antibodies, complement proteins, and SLE-specific antibodies (anti-dsDNA antibodies, Anti-Smith antibodies) are all included as additive criteria for SLE diagnosis.⁸³

Rheumatoid Arthritis

In 2022, an international task force was formed to address the safety and efficacy of disease-modifying antirheumatic drugs (DMARDs) and glucocorticoids (GCs) in the treatment of Rheumatoid Arthritis. The guideline focuses on treatment concerns. Regarding “biomarkers” they caution that certain biomarkers – i.e., acute phase reactants (APRs) such as CRP and other biomarkers comprising APRs “may respond independently of clinical improvement when antibodies to the IL-6 receptors, JAK inhibitors and even TNF-inhibitors are used.” The guideline does not mention multi-biomarker and disease activity tests such as Vectra DA or PrismRA.⁸⁴

Systemic Lupus International Collaborating Clinics (SLICC)

The 2012 SLICC Classification Criteria for SLE splits the 17 criteria into two divisions—either clinical or immunologic. An individual scoring at least a four, including at least one clinical criterion and one immunologic criterion, is classified as having SLE. The criteria are cumulative and do not need to be concurrently expressed or present.⁸⁵ Mosca, et al. (2019) also analyzed the accuracy and validity of the SLICC classification criteria, using a cohort of 616 patients, reporting an accuracy of 83.1%, sensitivity of 83.5%, and specificity of 82.4%. The criteria include the following:⁸⁵

- a. Clinical Criteria
 - i. Acute cutaneous lupus, such as lupus malar rash or subacute cutaneous lupus
 - ii. Chronic cutaneous lupus, such as classic discoid rash or discoid lupus/lichen planus overlap
 - iii. Nonscarring alopecia

- iv. Oral or nasal ulcers
 - v. Joint disease
 - vi. Serositis
 - vii. Renal criteria, such as urine protein-to-creatinine ratio representing 500 mg protein/24 hours or red blood cell casts
 - viii. Neurologic criteria, such as seizures, psychosis, myelitis, and so on
 - ix. Hemolytic anemia
 - x. Leukopenia or lymphopenia
 - xi. Thrombocytopenia
- b. Immunologic Criteria
- i. ANA
 - ii. Anti-dsDNA
 - iii. Anti-Sm
 - iv. Antiphospholipid antibodies
 - v. Low complement (Low C3, Low C4, or Low CH50)
 - vi. Direct Coombs test in the absence of hemolytic anemia

British Columbia Rheumatoid Arthritis

The BC Rheumatoid Arthritis guideline includes a table of factors used in the diagnosis of rheumatoid arthritis. The C-Reactive Protein (CRP) or Erythrocyte Sedimentation Rate (ESR) test is noted as the "preferred test," CRP/ESR indicates only inflammatory process but the guideline notes "low specificity." For RF, "RF has low sensitivity and specificity for RA. Seropositive RA has a worse prognosis than seronegative RA." Regarding anti-CCP, they write, "Anti-cyclic citrullinated protein antibodies (Anti-CCP) may have some value."

For disease activity monitoring, "CRP is more sensitive to short term fluctuations" and "ESR elevated in many but not all with active inflammations." Concerning monitoring, Rheumatoid Factor Latex Test (RF), "RF has low sensitivity and specificity for RA. Seropositive RA has a worse prognosis than seronegative RA."⁸⁶

National Institute for Clinical Excellence (NICE)

In a section on referral, diagnosis and investigations, NICE recommends:

- "Refer for specialist opinion anyone with suspected persistent synovitis of undetermined cause. Refer urgently even if blood tests show a normal acute-phase response or negative rheumatoid factor and if:
 - The small joints of the hands or feet are affected
 - More than one joint is affected, or

- There has been a delay of three months or longer between symptom onset and seeking medical advice.

[Based on high and moderate quality observational studies of early prognosis and identification or diagnosis].”

- “Offer to test for rheumatoid factor in people with suspected rheumatoid arthritis who have synovitis. [Based on high and moderate quality early identification observational studies]
 - Consider measuring anticyclic citrullinated peptide antibodies in people with suspected rheumatoid arthritis if:
 - They are negative for rheumatoid factor, and
 - Combination therapy is being considered (see section on disease modifying antirheumatic drugs).⁸⁷

The Royal Australian College of General Practitioners (RACGP)

The RACGP provides a recommendation on diagnosing those with suspected rheumatoid arthritis: “RECOMMENDATION 4 – DIAGNOSTIC INVESTIGATIONS (Grade A)

For patients presenting with painful and swollen joints, GPs should support clinical examination with appropriate tests to exclude other forms of arthritis and other differential diagnoses, and to predict patients likely to progress to erosive disease. Base investigations should include:

- erythrocyte sedimentation rate (ESR) and/or C-reactive protein (CRP)
- rheumatoid factor (RhF) and anti-cyclic citrullinated peptide (anti-CCP) antibody levels.⁸⁸

US Food and Drug Administration (FDA)

Many labs have developed specific tests that they must validate and perform in house. These laboratory-developed tests (LDTs) are regulated by the Centers for Medicare and Medicaid (CMS) as high-complexity tests under the Clinical Laboratory Improvement Amendments of 1988 (CLIA '88). LDTs are not approved or cleared by the US Food and Drug Administration; however, FDA clearance or approval is not currently required for clinical use.

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History

Date	Comments
11/01/25	New policy, approved October 14, 2025, effective for dates of service on or after February 6, 2026, following 90-day provider notification. Add to Routine Test Management Policy section. Antinuclear antibodies (ANA) testing is reimbursable once during initial workup and up to two additional times for new or worsening symptoms; extractable nuclear antigens (ENA) panel for raised ANA with clinical correlation; dsDNA up to four times yearly for systemic autoimmune rheumatic disease; anti-Jo-1 and anti-SSA in specific contexts. ANA monitoring, testing in asymptomatic individuals, antibody testing without positive ANA, and rheumatoid arthritis biomarker panels not listed are not reimbursable.

Disclaimer: This policy for routine test management is a guide in evaluating the clinical appropriateness and reimbursement methodology for lab tests. The Company adopts policies after careful review of published peer-reviewed scientific literature, national guidelines and local standards of practice. Since medical technology is constantly changing, the Company reserves the right to review and update policies as appropriate. Member contracts differ in their benefits. Always consult the member benefit booklet or contact a member service representative to determine coverage for a specific medical service or supply. CPT codes, descriptions and materials are copyrighted by the American Medical Association (AMA). ©2025 Premera All Rights Reserved.

Scope: Medical policies for routine test management are systematically developed guidelines that serve as a resource for Company staff when determining coverage for specific medical procedures, drugs or devices and reimbursement methodology. Coverage and reimbursement for medical services is subject to the limits and conditions of the member benefit plan. Members and their providers should consult the member benefit booklet or contact a customer service representative to determine whether there are any benefit limitations applicable to this service or supply. This medical policy does not apply to Medicare Advantage.