Systemic Sclerosis Antibodies

Systemic sclerosis (SSc), also called scleroderma, is a chronic autoimmune disorder characterized by fibrosis of the skin and various organs. Early diagnosis and classification are important so that patients can be evaluated for organ involvement and/or damage. However, because SSc is a heterogeneous disease, clinical presentation and disease course vary, and manifestations may overlap with those of other rheumatic disorders, all of which can complicate diagnosis. The initial testing strategy includes CBC with platelet count and automated differential and nuclear antibody (ANA) by IFA, IgG. Confirmatory or secondary testing should be performed based on ANA IFA patterns (centromere, nucleolar, and speckled patterns), clinical presentation, and/or ethnicity.

Disease Overview

Prevalence
0.3-2.8/100,000 worldwide

Age of Onset
20-50 years of age

Sex
M:F, 1:3

Diagnostic Issues

Autoimmune connective tissue diseases may present with similar features, particularly early in disease, making diagnosis difficult. The following considerations may assist in determining a diagnosis of SSc:

- ANA IFA patterns may help define diagnostic pathways
  - Most patients with SSc will have either:
    - Centromere antibodies
    - Scl-70 antibodies
    - RNA polymerase III antibodies
  - These three antibody tests are required for the initial evaluation of SSc although the use of other autoantibody markers may improve diagnostic sensitivity for disease
- The presence of SSc-specific antibodies may help predict disease phenotypes, which include:
  - CREST (calcinosis, Raynaud phenomenon, esophageal dysmotility, sclerodactyly, and telangiectasia) syndrome: antibodies against centromere are most common
  - Diffuse cutaneous SSc: antibodies against Scl-70 and RNA polymerase III most common
- Antibody patterns may differ by ethnicity
  - Antibodies against U3-RNP are more common in African Americans
  - Th/To autoantibodies are more common in White individuals with limited cutaneous SSc

Test Interpretation

Clinical Sensitivity
- ANA by IFA for SSc: 90-95%
- Individual SSc-specific marker: may vary based on ethnicity
Results

- ANA patterns (including centromere) are reported
- If positive, pattern and titers are reported

Limitations

- Negative antibody test result does not exclude SSc (5-10% of patients with SSc are ANA IFA negative)²,⁶
- Panel does not include Th/To

References


Related Information

**Systemic Sclerosis - Scleroderma**

Related Tests

**Antinuclear Antibody (ANA) with HEp-2 Substrate, IgG by IFA 3000082**
Method: Semi-Quantitative Indirect Fluorescent Antibody

**Antinuclear Antibody (ANA) with HEp-2 Substrate, IgG by IFA with Reflex by Pattern 3000601**

**Centromere Antibody, IgG 0050714**
Method: Semi-Quantitative Multiplex Bead Assay

**Connective Tissue Disease First Line Panel with Reflex 3002463**
Method: Semi-Quantitative Enzyme-Linked Immunosorbent Assay/Semi-Quantitative Indirect Fluorescent Antibody (IFA)/Semi-Quantitative Multiplex Bead Assay

**Connective Tissue Diseases Profile 0051668**
Method: Semi-Quantitative Enzyme-Linked Immunosorbent Assay/Semi-Quantitative Multiplex Bead Assay

**Fibrillarin (U3 RNP) Antibody, IgG 2012173**
Method: Qualitative Immunoblot

**Extended Myositis Panel 3001781**
Method: Semi-Quantitative Enzyme-Linked Immunosorbent Assay/Qualitative Immunoprecipitation/Semi-Quantitative Multiplex Bead Assay/Qualitative Immunoblot

**PM/Scl-100 Antibody, IgG by Immunoblot 2003040**
Method: Qualitative Immunoblot

**RNA Polymerase III Antibody, IgG 2001601**
Method: Semi-Quantitative Enzyme-Linked Immunosorbent Assay

**Smith/RNP (ENA) Antibody, IgG 0050470**
Method: Semi-Quantitative Enzyme-Linked Immunosorbent Assay
**Scleroderma (Scl-70) (ENA) Antibody, IgG 0050599**

**Method:** Semi-Quantitative Multiplex Bead Assay

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