

Laboratory Investigation Report

Patient Name	Centre
Age/Gender	OP/IP No/UHID
MaxID/Lab ID	Collection Date/Time
Ref Doctor	Reporting Date/Time

Hematology Special


SIN No: MBB375017

Max ANA Profile Comprehensive (ANA by IFA & ANA LIA)

Test Name	Result	Unit	Bio Ref Interval
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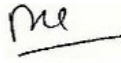
ANA (Anti-Nuclear Antibody) Immunofluorescence

Anti Nuclear Antibodies Immunofluorescence	Positive		Negative
Primary Dilution	1:40		
Primary Intensity on IF	4+		
Nuclear Pattern	Dense Fine Speckled (DFS)		
Cytoplasmic Pattern	Negative		
Mitotic Pattern	Negative		
End Point Titre	1:2560		
Advise :	ANA LIA		

Kindly correlate with clinical findings

***** End Of Report *****

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 Max Lab & Blood Bank Services


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Dr. Nitin Dayal, M.D.
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Laboratory Investigation Report

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Serology Special



SIN No: MBB375017

Max ANA Profile Comprehensive (ANA by IFA & ANA LIA)

Test Name	Result	Unit	Bio Ref Interval
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ANA - LIA, Serum

LIA

dsDNA	Negative
Nucleosome	Negative
Histones	Negative
SmD1	Negative
PCNA	Negative
PO (RPP)	Negative
SS-A/Ro60	Negative
SS-A/Ro52	Negative
SS-B/La	Negative
CENP-B	Negative
Sci70	Negative
U1-snRNP	Negative
AMA M2	Negative
Jo-1	Negative
PM-Sci	Negative
Mi-2	Negative
Ku	Negative
DFS70	Negative

Interpretation

1. The test provides a differential diagnosis using 17 different autoantibodies which is an additional diagnostic survey of autoimmune diseases like SLE , Mixed connective tissue diseases, Rheumatoid arthritis, Sjorgensyndrome, Progressive systemic sclerosis and CREST syndrome.
2. A definitive clinical diagnosis should not be made by result of a single test only, but should be made by taking clinical history and other laboratory findings in to account.

Kindly correlate with clinical findings

*** End Of Report ***



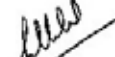
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The authenticity of the report can be verified by scanning the Q R Code on top of the page

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Conditions of Reporting: 1. The tests are carried out in the lab with the presumption that the specimen belongs to the patient name as identified in the bill/test request form. 2. They relate specifically to the sample received in the lab and are presumed to have been generated and transported per specific instructions given by the physicians/laboratory. 3. 1 results are for the information and interpretation by the referring doctor only. 4. Some tests are referred to other laboratories to provide a wider test menu to the customer. 5. Max shall in no event be liable for accidental damages loss, or destruction of specimen which is not attributable to any direct and mala fide act or omission of Max Healthcare or its Liability of Max Healthcare for deficiency of services, or other errors and omissions shall be limited to fee paid by the patient for the relevant laboratory services.



MC-2714