

Patient Name	: Mr MR.DUMMY		
DOB/Age/Gender	: 23 Y/Male	Sample Collected	: Apr 26, 2024, 01:00 PM
Patient ID / UHID	: 8052668/RCL7249293	Report Date	: May 25, 2024, 12:11 PM.
Referred By	: Dr. Dr. X	Barcode No	: SI485025
Sample Type	: Serum	Report Status	: Final Report
Test Description	Value(s)	Unit(s)	Reference Range

Extractable Nuclear Antigen (ENA) Panel, Quantitative

U1RNP ANTIBODIES <i>EIA, SERUM</i>	2.32	AU/ml	NEGATIVE: <12 EQUIVOCAL: 12-18 POSITIVE: >18
SSA - A (Soluble Substance A) IgG <i>EIA, SERUM</i>	1.14	AU/ml	NEGATIVE: < 12 EQUIVOCAL: 12-18 POSITIVE: > 18
SSB - B Antibody <i>(Serum,EIA)</i>	1.16	AU/ml	NEGATIVE: < 12 EQUIVOCAL: 12-18 POSITIVE: > 18
Scl - 70 Antibody IgG <i>EIA, SERUM</i>	1.01	AU/ml	NEGATIVE: < 12 EQUIVOCAL: 12-18 POSITIVE: > 18
Jo-1 Antibody IgG <i>(Serum,EIA)</i>	1.32	AU/ml	NEGATIVE: < 12 EQUIVOCAL: 12-18 POSITIVE: > 18
Sm Antibody IgG <i>(Serum,EIA)</i>	1.11	AU/ml	NEGATIVE: < 12 EQUIVOCAL: 12-18 POSITIVE: > 18
Centromere Antibody IgG <i>(Serum,EIA)</i>	1.98	AU/ml	NEGATIVE: <12 EQUIVOCAL: 12-18 POSITIVE: >18
Anti Double Stranded DNA (dsDNA-IgG) <i>SERUM, EIA</i>	1.11	IU/ml	Negative: < 19.9 Equivocal: 20-29.9 Positive: > 30.0
Rheumatoid Factor IgM <i>EIA, SERUM</i>	1.02	AU/ml	NEGATIVE: < 12 EQUIVOCAL: 12-18 POSITIVE: > 18

*** End Of Report ***

Disclaimer: Method given in report are only indicative and can be changed depending upon type of machine and kit available at time of testing.

Not all tests at all locations are under NABL scope. Availability of tests under NABL scope varies from lab to lab.



Dr. Dummy



Booking Centre :- DEMO PARTNER CHENNAI, DEMO PARTNER CHENNAI
Processing Lab :-

928-909-0609

ccsupport@redcliffelabs.com

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All Lab results are subject to clinical interpretation by qualified medical professional and this report is not subject to use for any medico-legal purpose.

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5. The Customers assume full responsibility for apprising the Company of any factors that may impact the test finding. These factors, among others, includes dietary intake, alcohol, or medication / drug(s) consumption, or fasting. This list of factors is only representative and not exhaustive.

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