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PATIENT REPORT

500 Chipeta Way, Salt Lake City, Utah 84108-1221			
phone: 801-583-2787, toll free: 800-522-2787		Patient Age/Sex:	Unknown
Jonathan R. Genzen, MD, PhD, Chief Medical Office	Pr	Tallent Age/Jex.	Unknown
Specimen Collected: 18-Dec-23 09:31			
Neuronal Nuclear Abs IgG, IB, CSF	Received: 18-Dec-23 0	9:40 Report/Ver	rified: 18-Dec-23 09:53
Procedure	Result	Units	Reference Interval
Neuronal Nuclear Ab (Hu) IgG, IH CSF	3, Positive * ⁱ¹		[Negative]
Neuronal Nuclear Ab (Ri) IgG, IA CSF	3, Positive * ⁱ²		[Negative]
Neuronal Nuclear Ab (Yo) IgG, I CSF	3, Positive * ⁱ³		[Negative]
Neuronal Nuclear Ab (TR/DNER) IqG,CSF	Positive * ⁱ⁴		[Negative]
Neuronal Nuclear Antibody Titer, IgG CSF	Received: 18-Dec-23 0	9:40 Report/Ver	rified: 18-Dec-23 09:53
Procedure	Result	Units	Reference Interval
Neuronal Nuclear Ab Titer,IgG CSF	1:40 * ¹⁵		[< 1:1]
Paraneoplastic Reflexive Panel, CSF	Received: 18-Dec-23 0	9:40 Report/Ver	rified: 18-Dec-23 09:53
Procedure	Result	Units	Reference Interval
Paraneoplastic Abs (PCCA/ANNA) IgG,CSF	ANNA Detected * fl i	6	[None Detected]
CV2 Ab IgG CBA-IFA Screen,CSF	Detected * t1 i7		[< 1:1]
SOX1 Antibody, IgG by Immunoblot CSF	, Positive * ⁱ⁸		[Negative]
Amphiphysin Antibody,CSF	High Positive ^{* i9}		[Negative]
CV2 Ab IgG Titer by CBA-IFA, CSF	Received: 18-Dec-23 0	9:40 Report/Ver	rified: 18-Dec-23 09:53
Procedure	Result	Units	Reference Interval
CV2 Ab IgG CBA-IFA Titer,CSF	1:40 * ⁱ¹⁰		[< 1:1]
Interpretive Textt1:18-Dec-23 09:31 (CV2 Ab IgG CBA-IFA Screen, CSF)CV2 Antibody, IgG is detected. Titer results to follow. Additional charges apply.			
Result Footnote f1: Paraneoplastic Abs (PCCA/ANNA) IgG, CSF			
Antibodies detected, therefore IFA titer and Immunoblot testing to be performed.			
Test Information i1: Neuronal Nuclear Ab (Hu) IgG, IB, CSF INTERPRETIVE INFORMATION: Neuronal Nuclear Ab (Hu) IgG, IB, CSF			
This test detects IgG and antigens.	cineuronal antibodie	es to Hu, Ri, and Yo	o and Tr (DNER)

Antineuronal antibodies serve as markers that aid in discriminating between a true paraneoplastic neurological disorder (PND) and other inflammatory disorders of the nervous system. Anti-Hu (antineuronal nuclear antibody, type I) is associated with

*=Abnormal, #=Corrected, C=Critical, f=Result Footnote, H-High, i-Test Information, L-Low, t-Interpretive Text, @=Performing lab

Unless otherwise indicated, testing performed at: **ARUP Laboratories** 500 Chipeta Way, Salt Lake City, UT 84108 Laboratory Director: Jonathan R. Genzen, MD, PhD

ARUP Accession: 23-352-900107 Report Request ID: 18510348 Printed: 20-Dec-23 12:37 Page 1 of 4

Patient Age/Sex:

Unknown

Test Information

i1: Neuronal Nuclear Ab (Hu) IgG, IB, CSF small cell lung cancer. Anti-Ri (antineuronal nuclear antibody, type II) is associated with neuroblastoma in children and with fallopian tube and breast cancer in adults. Anti-Yo (anti-Purkinje cell cytoplasmic antibody) is associated with ovarian and breast cancer. Anti-Tr(DNER) is associated with Hodgkin's lymphoma. The presence of one or more of these antineuronal antibodies supports a clinical diagnosis of PND and should lead to a focused search for the underlying neoplasm. This test was developed and its performance characteristics determined by ARUP Laboratories. It has not been cleared or approved by the US Food and Drug Administration. This test was performed in a CLIA certified laboratory and is intended for clinical purposes. i2: Neuronal Nuclear Ab (Ri) IgG, IB, CSF INTERPRETIVE INFORMATION: Neuronal Nuclear Ab (Ri) IgG, IB, CSF This test was developed and its performance characteristics determined by ARUP Laboratories. It has not been cleared or approved by the US Food and Drug Administration. This test was performed in a CLIA certified laboratory and is intended for clinical purposes. Neuronal Nuclear Ab (Yo) IgG, IB, CSF i3: INTERPRETIVE INFORMATION: Neuronal Nuclear Ab (Yo) IgG, IB, CSF This test was developed and its performance characteristics determined by ARUP Laboratories. It has not been cleared or approved by the US Food and Drug Administration. This test was performed in a CLIA certified laboratory and is intended for clinical purposes. Neuronal Nuclear Ab (TR/DNER) IgG, CSF i4: INTERPRETIVE INFORMATION: Neuronal Nuclear Ab (TR/DNER) IgG, CSF This test was developed and its performance characteristics determined by ARUP Laboratories. It has not been cleared or approved by the US Food and Drug Administration. This test was performed in a CLIA certified laboratory and is intended for clinical purposes. i5: Neuronal Nuclear Ab Titer, IgG CSF INTERPRETIVE INFORMATION: Neuronal Nuclear Ab Titer, IgG CSF This test was developed and its performance characteristics determined by ARUP Laboratories. It has not been cleared or approved by the US Food and Drug Administration. This test was performed in a CLIA certified laboratory and is intended for clinical purposes. i6: Paraneoplastic Abs (PCCA/ANNA) IgG, CSF INTERPRETIVE INFORMATION: Paraneoplastic Abs (PCCA/ANNA) IgG, CSF

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phone: 801-583-2787, toll free: 800-522-2787 Jonathan R. Genzen, MD, PhD, Chief Medical Officer

Patient Age/Sex:

Unknown

Test Information

i6: Paraneoplastic Abs (PCCA/ANNA) IgG, CSF
 This test was developed and its performance characteristics determined by ARUP
 Laboratories. It has not been cleared or approved by the US Food and Drug
 Administration. This test was performed in a CLIA certified laboratory and is
 intended for clinical purposes.
 i7: CV2 Ab IgG CBA-IFA Screen, CSF

INTERPRETIVE INFORMATION: CV2 Ab IgG CBA-IFA Screen, CSF

CV2 antibodies aid in discriminating between chronic paraneoplastic neurological disorder (PND) and other inflammatory disorders of the nervous system. Anti-CV2 is associated with small-cell lung cancer and thymoma. A negative test result does not rule out a diagnosis of autoimmune neurologic disease. Results should be interpreted in correlation with the patient's clinical history and other laboratory findings.

This indirect fluorescent antibody assay utilizes CV2 transfected cell lines for the detection and semiquantification of the CV2 IgG antibody.

This test was developed and its performance characteristics determined by ARUP Laboratories. It has not been cleared or approved by the U.S. Food and Drug Administration. This test was performed in a CLIA-certified laboratory and is intended for clinical purposes.

i8: SOX1 Antibody, IgG by Immunoblot, CSF INTERPRETIVE INFORMATION: SOX1 Antibody, IgG by Immunoblot, CSF

SOX1 antibody is detected in patients with Lambert-Eaton myasthenic syndrome (LEMS) and in patients with paraneoplastic cerebellar degeneration (PCD), paraneoplastic and nonparaneoplastic neuropathy. SOX1 antibody is associated with small cell lung cancer. A negative test result does not rule out a diagnosis of LEMS or other causes of paraneoplastic neurological syndrome.

This test was developed and its performance characteristics determined by ARUP Laboratories. It has not been cleared or approved by the U.S. Food and Drug Administration. This test was performed in a CLIA-certified laboratory and is intended for clinical purposes.

i9: Amphiphysin Antibody, CSF INTERPRETIVE INFORMATION: Amphiphysin Antibody IgG, CSF

Amphiphysin antibody is present in about 5 percent of patients with stiff-person syndrome and is found variably in other causes of paraneoplastic neurological syndrome (PNS). Amphiphysin antibody is mainly associated with small-cell lung cancer and breast tumors.

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Patient Age/Sex:

Unknown

Test Information

Amphiphysin Antibody, CSF
 Administration. This test was performed in a CLIA certified laboratory and is intended for clinical purposes.
 CV2 Ab IgG CBA-IFA Titer, CSF

INTERPRETIVE INFORMATION: CV2 Ab IgG CBA-IFA Titer, CSF

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