

Client: Example Client ABC123 123 Test Drive Salt Lake City, UT 84108 UNITED STATES

Physician: Doctor, Example

Patient: Patient, Example

Unknown
Unknown
01234567890ABCD, 012345
01234567890ABCD
00/00/0000 00:00

Autoimmune Encephalopathy/Dementia Panel, Serum

ARUP test code 3006201

Neuronal Antibody (Amphiphysin)	Negative INTERPRETIVE INFORMATION: An	(Ref Interval: Negative) mphiphysin Antibody, IgG
	with stiff-person syndrome a of paraneoplastic neurologic	sent in about 5 percent of patients and is found variably in other causes cal syndrome (PNS). Amphiphysin ed with small-cell lung cancer and
	determined by ARUP Laborator approved by the US Food and	its performance characteristics ries. It has not been cleared or Drug Administration. This test was ed laboratory and is intended for
Purkinje Cell/Neuronal Nuclear IgG Scrn	None Detected	(Ref Interval: None Detected)
	ANNA-1, ANNA-2, PCCA-1 or PC No further testing will be p	CCA-Tr(DNER) antibodies not detected. Derformed.
	INTERPRETIVE INFORMATION: PU	urkinje Cell/Neuronal Nuclear IgG Scrn
	determined by ARUP Laborator approved by the US Food and	its performance characteristics ries. It has not been cleared or Drug Administration. This test was ed laboratory and is intended for
NMDA Receptor Ab IgG CBA-IFA, Serum	<1:10	(Ref Interval: <1:10)
	Antibodies to NMDA were not follow.	detected, no additional testing to

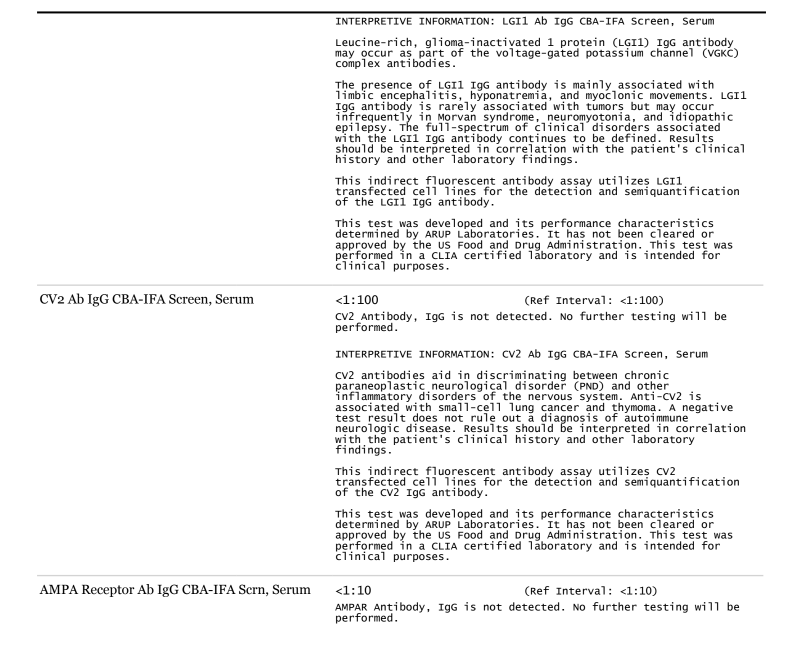
H=High, L=Low, *=Abnormal, C=Critical



INTERPRETIVE INFORMATION: NMDA Receptor Ab IgG CBA-IFA, Serum

	NMDA receptor antibody is found in a subset of patients with autoimmune limbic encephalitis and may occur with or without associated tumor. Decreasing antibody levels may be associated with therapeutic response. In addition, positive results have been reported in patients with non-autoimmune phenotypes. A negative test result does not rule out a diagnosis of autoimmune limbic encephalitis. Results should be interpreted in correlation with the patient's clinical history and other laboratory findings. Serum testing should be paired with CSF testing for improved diagnostic sensitivity. This indirect fluorescent antibody assay utilizes full-length GluN1 transfected cell lines for the detection and semiquantification of NMDA receptor IgG antibody. 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.
CASPR2 Ab IgG CBA-IFA Screen, Serum	<1:10 (Ref Interval: <1:10)
	CASPR2 Antibody, IgG is not detected. No further testing will be performed.
	INTERPRETIVE INFORMATION: CASPR2 Ab IgG CBA-IFA Screen,
	Serum Contactin-associated protein-2 (CASPR2) IgG antibody may occur as part of the voltage-gated potassium channel (VGKC) complex antibodies.
	The presence of CASPR2 IgG antibody is associated with a wide spectrum of clinical manifestations, including acquired neuromyotonia, limbic encephalitis, painful neuropathy, and Morvan syndrome. Tumors such as thymoma, small cell lung cancer, and other rarer tumors may occur. The full-spectrum of clinical disorders and tumors associated with the CASPR2 IgG antibody continues to be defined. Results should be interpreted in correlation with the patient's clinical history and other laboratory findings.
	This indirect fluorescent antibody assay utilizes CASPR2 transfected cell lines for the detection and semiquantification of the CASPR2 IgG antibody.
	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.
LGI1 Ab IgG CBA-IFA Screen, Serum	<1:10 (Ref Interval: <1:10)
	LGI1 Antibody, IgG is not detected. No further testing will be performed.

H=High, L=Low, *=Abnormal, C=Critical



H=High, L=Low, *=Abnormal, C=Critical

Unless otherwise indicated, testing performed at:

ARUP LABORATORIES | 800-522-2787 | aruplab.com 500 Chipeta Way, Salt Lake City, UT 84108-1221 Jonathan R. Genzen, MD, PhD, Laboratory Director Patient: Patient, Example ARUP Accession: 24-057-100498 Patient Identifiers: 01234567890ABCD, 012345 Visit Number (FIN): 01234567890ABCD Page 3 of 7 | Printed: 2/26/2024 11:13:06 AM 4848



	Alpha-amino-3-hydrox receptor (AMPAR) ant autoimmune limbic en associated tumor. De with therapeutic res out a diagnosis of a interpreted in corre and other laboratory This indirect fluore transfected cell lin of AMPAR IgG antibod	escent antibody assay utilizes AMPAR les for the detection and semiquantification
	determined by ARUP L approved by the US F	aboratories. It has not been cleared or ood and Drug Administration. This test was certified laboratory and is intended for
GABA-BR Ab IgG CBA-IFA Scrn, Ser	<1:10 GABA-BR Antibody, Ig be performed.	(Ref Interval: <1:10) G is not detected. No further testing will
	INTERPRETIVE INFORMA	TION: GABA-BR Ab IgG CBA-IFA Scrn, Ser
	found in a subset of autoimmune neurologi associated tumor. De with therapeutic res out a diagnosis of a	acid receptor, type B (GABA-BR) antibody is patients with autoimmune epilepsy and other c phenotypes; it may occur with or without creasing antibody levels may be associated ponse. A negative test result does not rule utoimmune neurologic disease. Results d in correlation with the patient's clinical boratory findings.
		escent antibody assay utilizes GABA-BR les for the detection and semiquantification loody.
	determined by ARUP L approved by the US F	pped and its performance characteristics aboratories. It has not been cleared or ood and Drug Administration. This test was certified laboratory and is intended for
SOX1 Antibody, IgG by Immunoblot, Serum	Negative	(Ref Interval: Negative)
	INTERPRETIVE INFORMA	TION: SOX1 Antibody, IgG by Immunoblot, Serum
	myasthenic syndrome cerebellar degenerat nonparaneoplastic ne small cell lung canc	ected in patients with Lambert-Eaton (LEMS) and in patients with paraneoplastic ion (PCD), paraneoplastic and suropathy. SOX1 antibody is associated with eer. A negative test result does not rule out or other causes of paraneoplastic
	determined by ARUP L approved by the U.S.	pped and its performance characteristics aboratories. It has not been cleared or Food and Drug Administration. This test was certified laboratory and is intended for
DPPX Ab IgG CBA-IFA Screen, Serum	<1:10	(Ref Interval: <1:10)

H=High, L=Low, *=Abnormal, C=Critical



DPPX Antibody, ${\tt IgG}$ is not detected. No further testing will be performed.

INTERPRETIVE INFORMATION: DPPX Ab IgG CBA-IFA Screen, Serum

DPPX antibody is found in a subset of patients with autoimmune encephalitis, and is often associated with prodromal gastrointestinal symptoms and unintentional weight loss. It may occur with or without associated tumor. Decreasing antibody levels may be associated with therapeutic response. 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 DPPX transfected cells for the detection and semiquantification of the DPPX 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.

IgLON5 Ab IgG CBA-IFA Screen, Serum

(Ref Interval: <1:10)

 $\tt IgLON5$ Antibody, $\tt IgG$ is not detected. No further testing will be performed.

INTERPRETIVE INFORMATION: IgLON5 Ab IgG CBA-IFA Screen, Serum

IgLON Family Member 5 (IgLON5) antibody is found in a subset of patients with autoimmune encephalitis or other autoimmune neurologic/neurodegenerative disorders and may occur with or without associated tumor. A negative test result does not rule out a diagnosis of an autoimmune neurologic disorder. Interpretation of any antineural antibody test requires clinical correlation.

This indirect fluorescent antibody assay utilizes IgLON5 transfected cell lines for detection and semi-quantification of IgLON5 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.

mGluR1 Ab IgG CBA-IFA Screen, Serum

<1:10 (Ref Interval: <1:10)
mGluR1 Antibody, IgG is not detected. No further testing will be
performed.</pre>

H=High, L=Low, *=Abnormal, C=Critical

<1:10



INTERPRETIVE INFORMATION: mg]uR1 Ab IgG CBA-IFA Screen,

Serum Metabotropic glutamate receptor 1 (mGluR1) antibody is found in a subset of patients with autoimmune cerebellar ataxia or

autoimmune encephalitis and may occur with or without associated tumor. A negative test result does not rule out a diagnosis of autoimmune cerebellar ataxia or limbic encephalitis. Interpretation of any antineural antibody test requires clinical correlation.

This indirect fluorescent antibody assay utilizes mGluR1 transfected cell lines for detection and semi-quantification of mGluR1 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.

Glutamic Acid Decarboxylase Antibody

<5.0 IU/mL (Ref Interval: 0.0-5.0) INTERPRETIVE INFORMATION: Glutamic Acid Decarboxylase Antibody

A value greater than 5.0 IU/mL is considered positive for Glutamic Acid Decarboxylase Antibody (GAD Ab). This assay is intended for the semi-quantitative determination of the GAD Ab in human serum. Results should be interpreted within the context of clinical symptoms.

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VERIFIED/REPORTED DATES						
Procedure	Accession	Collected	Received	Verified/Reported		
Neuronal Antibody (Amphiphysin)	24-057-100498	00/00/0000 00:00	00/00/0000 00:00	00/00/0000 00:00		
Purkinje Cell/Neuronal Nuclear IgG Scrn	24-057-100498	00/00/0000 00:00	00/00/0000 00:00	00/00/0000 00:00		
NMDA Receptor Ab IgG CBA-IFA, Serum	24-057-100498	00/00/0000 00:00	00/00/0000 00:00	00/00/0000 00:00		
CASPR2 Ab IgG CBA-IFA Screen, Serum	24-057-100498	00/00/0000 00:00	00/00/0000 00:00	00/00/0000 00:00		
LGI1 Ab IgG CBA-IFA Screen, Serum	24-057-100498	00/00/0000 00:00	00/00/0000 00:00	00/00/0000 00:00		
CV2 Ab IgG CBA-IFA Screen, Serum	24-057-100498	00/00/0000 00:00	00/00/0000 00:00	00/00/0000 00:00		
AMPA Receptor Ab IgG CBA-IFA Scrn, Serum	24-057-100498	00/00/0000 00:00	00/00/0000 00:00	00/00/0000 00:00		
GABA-BR Ab IgG CBA-IFA Scrn, Ser	24-057-100498	00/00/0000 00:00	00/00/0000 00:00	00/00/0000 00:00		
SOX1 Antibody, IgG by Immunoblot, Serum	24-057-100498	00/00/0000 00:00	00/00/0000 00:00	00/00/0000 00:00		
DPPX Ab IgG CBA-IFA Screen, Serum	24-057-100498	00/00/0000 00:00	00/00/0000 00:00	00/00/0000 00:00		
IgLON5 Ab IgG CBA-IFA Screen, Serum	24-057-100498	00/00/0000 00:00	00/00/0000 00:00	00/00/0000 00:00		
mGluR1 Ab IgG CBA-IFA Screen, Serum	24-057-100498	00/00/0000 00:00	00/00/0000 00:00	00/00/0000 00:00		
Glutamic Acid Decarboxylase Antibody	24-057-100498	00/00/0000 00:00	00/00/0000 00:00	00/00/0000 00:00		

END OF CHART

H=High, L=Low, *=Abnormal, C=Critical