TEST:	Mumps Antibody IgG EIA
Synonym:	Anti-Mumps IgG; Mumps immunity test
Laboratory/Phone:	Vaccine Preventable Disease/443-681-3889
Specimen:	Serum
Specimen identification:	Label container with patient's last name, first Name, DOB,
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	specimen type, date and time of collection.
Specimen Volume Required:	1-2 ml
Specimen Volume Minimum:	1 ml
Collect:	Venipuncture; Red top vacuum tube, transfer serum to sterile tube
	with leak-proof cap.
Form:	DHMH 4677 (MMRV Immunity Screen; Mumps Immunity
	Screen)
Transport Conditions:	·
Transport Conditions: Packaging and Shipping:	2-8°C-Refer to serology test guideline. Follow packaging and shipping instructions.
Specimen Rejection:	Discrepancy between name on tube and name on form, unlabeled hemolytic; lipemic;
	gross bacterial contamination. Specimens collected > 7 days prior to submission. Refer
	to serology guideline.
Availability:	Service available only to government facilities Monday to Friday.
Results and Interpretation:	Negative: Indicates no detectable IgG antibody to Mumps virus. A negative results
	indicate no current or previous infection with Mumps. Virus. Such individuals are
	presumed to be susceptible to primary infection . However, specimen taken too early
	during a primary infection may not have detectable levels of IgG antibody. If primary
	infection is suspected, another specimen (convalescent) should be taken in 8-14 days
	and tested concurrently in the same assay with the original (acute) specimen to look for
	seroconversion. If acute specimen is negative and convalescent specimen is positive,
	seroconversion has taken place and a primary Mumps virus infection is indicated.
	Equivocal: Equivocal results are indeterminate. Patient may or may not have
	immunity to Mumps Virus. It is not acceptable proof of immunity.
	Positive: Indicates evidence of Mumps IgG antibodies and suggests past or current
	infection with Mumps virus. Antibodies obtained via acquired immunity or
Defended Devel	immunization and probable protection from clinical infection. (Immunity).
Reference Range:	Negative.
Additional Information:	For more information, see the CDC link at: http://www.cdc.gov/vaccines/pubs/pinkbook/downloads/mumps.pdf
Turnaround Time:	2-6 working days
Method:	ELISA
Purpose of Test:	For detection of IgG antibodies to Mumps virus, the test can be used to evaluate
	single sera for immune status.
Interfering Substances:	Test results in an immune compromised patients should be interpreted with caution.
Testing Site:	Central Laboratory 1770 Ashland Avenue, Baltimore, MD 21205.
Comment:	LIMITATIONS: A diagnosis should not be made on the basis of anti- Mumps results
	alone. Test results should be interpreted in conjunction with the clinical evaluation and
	the results of other diagnostic procedures. The antibody titer of a single serum
	specimen cannot be used to determine a recent infection. Paired samples (acute and
	convalescent) should be collected and tested concurrently to demonstrate
	seroconversion. Samples collected too early in the course of an infection may not have
	detectable levels of IgG. In such cases, a second sample may be collected after 2-7
	weeks and tested concurrently with the Original sample to look for seroconversion. A positive Mumps IgG test in neonates should be interpreted with caution since passively
	acquired maternal antibody can persist for up to 6 months.

TEST:	Mumps IgM Antibody IFA
Synonym:	Anti-Mumps IgM; antibody. Mumps IgM IFA
Laboratory/Phone:	Vaccine Preventable Disease/443-681-3889
Specimen:	Serum
Specimen identification:	Label container with patient's last name, first Name, DOB,
	specimen type, date and time of collection.
Specimen Volume Required:	1-2 ml
Specimen Volume Minimum:	1 ml
Collect:	Venipuncture; Red top vacuum tube, transfer serum to sterile tube
	with leak-proof cap.
Form:	DHMH 4677 (Other test) Prior authorization by Epidemiology is
	required (410 767-6628)
Transport Conditions:	2-8°C-Refer to serology test guideline.
Packaging and Shipping:	Follow packaging and shipping instructions.
Specimen Rejection:	Discrepancy between name on tube and name on form, unlabeled
	hemolytic; lipemic; gross bacterial contamination. Refer to serology
	guideline.
Availability:	Monday to Friday. Test available only to DHMH epidemiologist to
	investigate outbreak.
Results and Interpretation:	Negative: No significant level of Mumps IgM antibodies detected. A
•	negative result indicates no current infection with Mumps virus.
	However, specimens taken too early during a primary infection may
	not have detectable levels of IgM antibody. If a primary infection is
	suspected, another specimen should be taken within 7 days and
	tested concurrently in the same assay with the original specimen to
	look for seroconversion
	Positive: Evidence of Mumps IgM antibodies detected and indicative
	of current or recent infection.
Reference Range:	Negative.
Additional Information:	For more information, see the CDC link at:
	http://www.cdc.gov/vaccines/pubs/pinkbook/downloads/mumps.pdf
Turnaround Time:	1-2 working days
Method:	IFA
Purpose of Test:	Test available only to DHMH epidemiologist to investigate outbreak. For the detection of IgM antibodies to Mumps virus.
Interfering Substances:	Test results in an immune compromised patients should be interpreted with caution.
interrering substances.	IgM anticell antibodies, if present in the serum, may interfere with the Mumps IgM test.
Testing Site:	Central Laboratory 1770 Ashland Avenue, Baltimore, MD 21205.
Comment:	LIMITATIONS: Results of the Mumps IgM ELISA are not by themselves diagnostic and
	should be interpreted in light of the patient's clinical condition and results of other
	diagnostic procedures. Mumps virus specific IgG antibody may compete with IgM for binding sites and cause false negative results. Rheumatoid factor, if present along with
	specific IgG, will cause false positive results. The Sample diluent contains an absorbent
	which will remove IgG from the test specimen, and significantly reduce the possibility of
	false positive or negative results Heterotypic IgM antibody responses may occur in
	patients infected with Epstein-Barr virus. Samples taken too early during the course of
	a primary infection may not have detectable levels of mumps specific IgM. A negative result does not rule out a primary infection with mumps virus. False positive anti-mumps
	IgM results may be obtained from patients with autoimmune disease. The performance
	of the mumps IgM ELISA has not been validated using neonatal samples.