



Prenatal Screening and Diagnosis



testing at ARUP Laboratories



www.aruplab.com

ARUP LABORATORIES
500 Chipeta Way
Salt Lake City, UT 84108-1221
Phone: (800) 522-2787
Fax: (801) 583-2712

keyword: prenatal

*A nonprofit enterprise of the University of
Utah and its Department of Pathology*

© 2015 ARUP Laboratories
BD-CS-014, Rev 0, September 2015



[www.aruplab.com/
topics/prenatal](http://www.aruplab.com/topics/prenatal)



ARUP's comprehensive genetics lab allows both **prenatal screening and diagnostic testing** to be performed in the same location.

Prenatal screening and diagnosis refers to a group of tests used to assess the presence of fetal disease prior to birth.

Prenatal screening tests include maternal serum screening and non-invasive prenatal testing (NIPT). Prenatal diagnostic tests include amniocentesis and chorionic villus sampling, which are invasive procedures that allow the collection of material that can be used for biochemical testing, chromosome analysis, or DNA testing.

ARUP's comprehensive genetics lab allows both prenatal screening and diagnostic testing to be performed in the same location.

Diagnostic Testing	Sample type:		Performed in:		Methodology:		
	Amniocentesis	CVS	First trimester	Second trimester	Chromosome analysis	FISH	Microarray
Chromosome Analysis, Amniotic Fluid (2002293)	●			●	●		
Chromosome Analysis, Amniotic Fluid, with Reflex to Genomic Microarray (2008367)	●			●	●		●
Chromosome FISH, Amniotic Fluid with Reflex to Chromosome Analysis or Genomic Microarray (2011130)	●			●	●	●	●
Chromosome FISH, Prenatal (2002297)	●			●		●	
Cytogenomic SNP Microarray—Fetal (2002366)	●	●	●	●			●
Chromosome Analysis, Chorionic Villus (2002291)		●	●		●		
Chromosome FISH, Chorionic Villus with Reflex to Chromosome Analysis or Genomic Microarray (2011131)		●	●		●	●	●
Chorionic Villus, FISH (0040203)		●	●			●	

Prenatal Aneuploidy/NTD Screening	Performed in:		Screens for:			
	First trimester	Second trimester	Trisomy 21	Trisomy 18	Other chromosome abnormalities	Open neural tube defect
Non-Invasive Prenatal Testing for Fetal Aneuploidy (Panorama) (2007537)	●	●	●	●	Trisomy 13, triploidy, sex chromosome abnormalities	
Non-Invasive Prenatal Testing for Fetal Aneuploidy (Panorama) with Microdeletions (2010232)	●	●	●	●	Trisomy 13, triploidy, sex chromosome abnormalities, 5 microdeletion syndromes	
Maternal Serum Screen, First Trimester (0081150)	●		●	●		
Maternal Screening, Sequential, Specimen #1 (0081293)	●		●	●		
Maternal Screening, Sequential, Specimen #2 (0081294)	Result based on first and second trimester samples.		●	●		●
Maternal Serum Screening, Integrated, Specimen #1 (0081062)	●		●	●		
Maternal Serum Screening, Integrated, Specimen #2 (0081064)		●	●	●		●
Maternal Serum Screen, Alpha Fetoprotein, hCG, Estriol, and Inhibin A (0080269)		●	●	●		●
Maternal Serum Screen, Alpha Fetoprotein (Only) (0080434)		●				●

Products of Conception Testing
Chromosome Analysis, Products of Conception (2002288)
<ul style="list-style-type: none"> Order for standard chromosome analysis performed on fetal tissue or villi
Chromosome Analysis, Products of Conception, with Reflex to Genomic Microarray (2005762)
<ul style="list-style-type: none"> Useful to ensure the highest chance of obtaining meaningful results from POC specimens When tissue culture is unsuccessful or if the results of the chromosome analysis are normal, then testing reflexes to genomic microarray
Genomic SNP Microarray, Products of Conception (2005633)
<ul style="list-style-type: none"> For the detection of copy number alterations and loss of heterozygosity in unfixed fetal tissue or villi
Cytogenomic Molecular Inversion Probe Array, FFPE Tissue—Products of Conception (2010795)
<ul style="list-style-type: none"> For the detection of copy number alterations and loss of heterozygosity in FFPE fetal tissue or villi