



PATIENT NAME (Last)	(FIRST)	(M.I.)
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PATIENT I.D. NUMBER	LAB I.D. NUMBER
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BIRTHDATE	SEX	REFERRING PHYSICIAN
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PHONE/FAX RESULTS	(Required) <input type="checkbox"/> PHONE <input type="checkbox"/> SECURE FAX
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SPECIMEN COLLECTION	
DATE: _____	TIME: _____

SPECIMEN TRANSPORT <input type="checkbox"/> Refrigerated	SPECIMEN TYPE <input type="checkbox"/> Serum
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*****ATTENTION CLIENTS*****

The maternal screens listed below have separate specimen requirements. For tests requiring ultrasound information the NT may be obtained when the CRL is between 39mm and 85mm.

If the patient's NT measurement cannot be obtained, the first trimester or sequential screens cannot be ordered.

First Trimester Screen Draw specimen between 11w 0d and 13w 6d gestation. (CRL=44-85mm)	Integrated Screen Specimen #1: Draw specimen between 10w 3d and 13w 6d gestation. (CRL=34-85mm)	Sequential Screen Specimen #1: Draw specimen between 11w 0d and 13w 6d gestation. (CRL=44-85mm)
Quad, Triple or AFP-Only Screens Draw specimen between 14w 0d and 24w 6d gestation	Specimen #2: Draw specimen between 15w 0d and 22w 6d gestation.	Specimen #2: Draw specimen between 15w 0d and 22w 6d gestation.

Maternal Serum Screening Assays (Reorder # 41130)

- | | | |
|----------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------|
| <input type="checkbox"/> 0081062 Integrated, Specimen #1
<input type="checkbox"/> 0081064 Integrated, Specimen #2 | <input type="checkbox"/> 0081293 Sequential, Specimen #1
<input type="checkbox"/> 0081294 Sequential, Specimen #2
<input type="checkbox"/> 0081150 First Trimester | <input type="checkbox"/> 0080434 Alpha Fetoprotein (Only)
<input type="checkbox"/> 0080269 Alpha Fetoprotein, hCG, Estriol, & Inhibin A (Quad) |
|----------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------|

REQUIRED PATIENT INFORMATION:

- A. Current weight _____ lbs. or _____ kg.
- B. Due date (EDD) _____
 Determined by:
 Last menstrual period, confirmed by US
 Ultrasound Last menstrual period
- C. Number of Fetus:
 Singleton Twins Unknown
 Check box if pregnancy is monochorionic
- D. Patient's race?
 Caucasian Black Hispanic Asian Other
- E. Was the patient diabetic at the time of conception?
 No Yes
- F. Is there a family history of neural tube defects (i.e., spina bifida, anencephaly, or encephalocele)?
 No Yes
 If yes, relationship of the affected individual to the fetus? _____
- G. Has the patient had a previous pregnancy with a chromosome abnormality (i.e., Down syndrome, Trisomy 18, or 13)?
 No Yes
 If yes, specify abnormality _____
- H. Is this an in vitro fertilization pregnancy using a donor egg?
 No Yes
 If yes, age of egg donor _____ years.
- I. Has the patient taken valproic acid or carbamazepine during this pregnancy?
 No Yes
 If yes, specify drug _____
- J. Is this a repeat sample?
 No Yes Unknown
- K. Does the patient currently smoke cigarettes?
 No Yes Unknown

ADDITIONAL PATIENT INFORMATION (required for the First Trimester, Integrated-1 or Sequential-1 screens only)

Date of Ultrasound _____ Sonographer Name _____ Certification # _____

NT (mm) _____ CRL (mm) _____ Reading MD name _____ Certification # _____

If twins: Twin B NT (mm) _____ CRL (mm) _____

NEW YORK CLIENTS ONLY

Informed consent must be obtained before testing for New York clients only.
 Maternal screening which includes NT, pregnancy associated plasma protein-A (PAPP-A), alpha fetoprotein (AFP, human chorionic gonadotropin (hCG), unconjugated estriol (uE3), and dimeric inhibin A (DIA) testing is offered to detect neural tube defects (spina bifida), chromosome disorders, incorrect dating, and adverse fetal outcomes. Neural tube defects occur when the brain and spine do not form properly. In 1 to 2 of 1000 pregnancies, conditions such as anencephaly or spina bifida occur. A normal test does not guarantee a normal baby. Most birth defects are not detected by maternal serum testing. An abnormal result means further testing is required. This may include a repeat screening test,

ultrasound or amniocentesis. Counseling regarding the results will be provided by your physician. Maternal serum may be abnormal for many reasons including miscalculation of due date, twin pregnancy, vaginal bleeding or for no apparent reason.

I, _____, certify that I have read this information and understand it. All of my questions have been answered satisfactorily. I freely and voluntarily consent to the test for NT/PAPP-A/AFP/hCG/uE3/DIA and give my permission to Dr. _____ to send a sample of my blood to ARUP for testing. I authorize ARUP to report the test results to the above-named physician.

Patient signature _____ Date _____ Physician signature _____ Date _____

THIS BOX FOR ARUP LABORATORIES' USE ONLY

NUMBER OF SPECIMENS SUBMITTED _____

TOTAL NUMBER OF TESTS ORDERED _____

QTY _____	RT _____	R _____	F _____	ID# _____
SER _____	PLA _____	WB _____	URINE _____	STOOL _____
TISSUE _____	SST _____	OTHER _____	CSF _____	S/P _____
				WRAPPED _____