

Medical Laboratory, Sorrento Mesa 9535 Waples Street, Ste. 150 San Diego, CA 92121 CLIA# 05D1071362 CLIA Director: Amanda Haynes, DO

CHEMISTRY TESTS

Basic Metabolic Panel (BMP) – Na, K, Cl, CO2, Ca, ECREA, Glucose, BUN, (Calculated - GFR, Osmolality) Comprehensive Metabolic Panel (CMPN) – BMP plus ALPI, TBIL, AST, ALTI, Total Protein, (Calculated - GFR, Anion Gap, Osmolality)



Hepatic Function Panel/Liver Function Tests (HPF/LFT) – Albumin, ALPI, TBIL, DBIL, AST, ALTI, Total Protein Lipid Panel (LPC) – Chol, Trig, HDL, LDL (calculated). LDL Direct performed when requested.

Renal Function Panel (RFP) - BMP plus Albumin, Phosphorous (Calculated - GFR, Osmolality)

+ During downtime, calculate using the Excel spreadsheet available on <u>https://www.testmenu.com/scripps</u> under Downtime Resources or scan the QR code

Analyte, Blood	Reference range (Expected Values)	
A1 Antitruncin a1AT	O to uppropriate votes 00, 200 mg/dl	
	>19 yrs: 5 - 55 U/L	
ASI	>19 yrs: 5 - 34 U/L	
Albumin	19yrs to 60yrs: 3.5 to 5.0 g/dL	
	> 60 years: 3.4 to 4.8 g/dL	
Alkaline Phosphatase	Male: >19yrs: 40-150 U/L	
	Female: >19yrs: 40-150 U/L	
Amylase	Male & Female: (U/L)	
	0 -15days: 4-10	
	15d –13wks: 4-22	
	13wks - 1yr: 4-50	
	1yr -19yr: 25-101	
	>19- 70: 25 to 125 U/L	
· · ·	> 70 years: 20 to 160 U/L	
Ammonia	0 to 29 days: 18 – 72 µmol/L	
	>29 days: 18 – 72 µmol/L	
Anion Gap, Calculated	6-14 mmol/L	
Apolipoprotein B	0-1yr	
	Male: 16 - 124 mg/dL	
	Female: 17 - 120 mg/dL	
	1yr-12yrs	
	Male: 48 - 125 mg/dL	
	Female: 51 - 126 mg/dL	
	> 12 to 60 years	
	Male: 49 – 173 mg/dL	
	Female: 53 – 182 mg/dL	
	> 60 years	
	Male: 54 – 163 mg/dL	
	Female: 64 – 182 mg/dL	
ASO Streptolysin-O	50 - 200 IU/mL	
β2-Microglobulin	1.0 – 2.3 mg/L	
Bilirubin, Direct	Adult > 19 years	
	0.1 to 0.5 mg/dL	
Bilirubin, Total	0 day to 15 days: 0.2-15.0	
	15 days to 28 days: 0.1-0.7	
	28 days to 1 year: 0.1-0.7	
	1 year to 9 years: 0.1-0.4	
	9 years to 12 years: 0.1-0.6	
	12 yrs to 15 yrs: 0.1-0.7	
	15 yrs to 19 yrs: 0.1-0.8	
	Adult >19 years: 0.2 to 1.4 mg/dL	
Blood Urea Nitrogen (BUN)	Adult, Male	
	19- 50 years 9 to 21	
	>50 years 8 to 26	
	Adult, Female	
	19- 50 years 7 to 19	
	> 50 years 10 to 20	
Calcium	Newborn, 0 to 10 days 7.6 to 10.4 mg/dl	
	Newborn, 10 to 20 days 9.0 to 11.0 mg/dL	
	20 days to 12 years 8.8 to 10.8 mg/dL	

CORE LABORATORY REFERENCE RANGES

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Analyte, Blood	Reference range (Expected Values)
	For Pediatric values, see Caliper studies
	Adult 12 yrs and older 8.3 to 10.3 mg/dL
Carbamazepine	4 to 12 ug/mL
Carbon Dioxide	M/F 0 up to 15 day: 5-20 mmol/L
	M/F 15 days up to 1 year: 10-24 mmol/L
	M/F 1 year up to 5 years: 14-24 mmol/L
	M/F 5 years up to 15 years: 17-26 mmol/L
	Female 15 years to 19 years: 17-26 mmol/L
	Male 15 years to 19 years: 18-28 mmol/L
	Adult ≥19 years: 21-29 mmol/L
Cholesterol	Child: 0-12 yrs: 7-170 mg/dL
	Desirable <180 mg/dl
	Borderline 170-199 mg/dl
	High >=200 mg/dl
	Adult >12 yrs: 7-200
	Desirable <200 mg/dl
	Borderline 200-239 mg/dl
	High >=240 mg/dl
Chloride	0-30 days: 98 to 113 mmol/L
	30 days to 90 years: 98 to 107 mmol/L
	>90 years 98 to 111 mmol/L
Complement C3	M/F 0-15davs: 50-121
	M/F 15d – 1vr: 51-160
	1 to 14 years
	Male: 80 to 170 mg/dL
	Female: 82 to 173 mg/dL
	> 14 to 80 years
	Male: 82 to 185 mg/dL
	Female: 83 to 193 mg/dL
Complement C4	M/F 0-1vr: 7-30
	1 to 14 years
	Male: 14 to 44 mg/dL
	Female: 13 to 46 mg/dL
	> 14 to 80 years
	Male: 15 to 53 mg/dL
	Female: 15 to 57 mg/dL
Creatine Kinase	Male: 30 to 200 U/L
	Female: 29 to 168 U/L
Creatinine (Enzymatic)	>19 year Male: 0.56 to 1.40 mg/dL
	>19 year Female: 0.52 to 1.07 mg/dL
C-reactive protein CRP	0.4 – 5.0 mg/L
C-reactive protein CRP, high sensitivity	0.2 – 5.0 mg/L
Ceruloplasmin	0-2 months Male: 7.3-23.6
	0-2 months Female: 7.4-23.7
	M/F 2mos-6mos: 13.5-32.9
	M/F 6mos- 1yr: 13.7-38.9
	M/F 1-8yr: 21.7-43.3
	M/F 8-14yr: 20.5-40.2
	14-19yrs Male: 17-34.8
	14-19yrs Female: 20.8-43.2
	M/F >19yrs: 20- 60 mg/dL
Cyclosporine	Patient assessment required.
	Methodology: Chemiluminescent microparticle
	immunoassay (CMIA) performed on the Abbott Architect
	i1000sr/i2000sr
Digoxin	<1.0 ng/mL (therapeutic range)
Gamma-Glutamyl Transferase (GGT)	M/F 0-15days: 23-219
,	M/F 15d- 1yr: 8-127
	M/F 1yr -11 yr: 6-16
	M/F 11-19yr: 7-21
	Male 12 to 64 U/L
	Female 9 to 36 U/L

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Analyte, Blood	Reference range (Expected Values)
	For Pediatric values, see Caliper studies
Glucose	ADA decision limits for fasting glucose:
	Normal 70-99 mg/dL
	Impaired 100-125 mg/dL
	Diabetes >125 mg/dL
	0-29 days: 40-90mg/dL
	>29 days: 70-125
Glucose Screen, Pregnancy, 50gms 1hr-Gestational	50-139 mg/dl
Glucose-3hr-Tolerance Test, 100-gm (Gestational)	Fasting: 70-94 mg/dl
	1 hour: 70-179 mg/dl
	2 hour: 70-154 mg/dl
	3 hour: 70-139 mg/dl
Glucose, 2-hour (Non-Gestational) 75-gm	70-139 mg/dl
Haptoglobin	Units (mg/dL)
	0 to 1 year
	Male 0 to 300
	Female 0 to 235
	> 1 to 12 years
	Male 3 to 270
	Female 11 to 220
	> 12 to 60 years
	Male 14 to 258
	Female 35 to 250
	> 60 years
	Male 40 to 268
	Female 63 to 273
Hemoglobin A1c (Glycosylated hemoglobin)	Normal: < 5.7%
	Increased risk for diabetes: 5.7-6.4%
	Diabetes: >6.4%
High Density Lipoprotein (HDL), Ultra	Male 30-70 mg/dL
	Female 30-85 mg/dL
Immunoglobulin A (IgA)	Units (mg/dL)
	0 to 3 months
	Male 5 to 34
	Female 5 to 34
	> 3 months to 1 year
	Male 8 to 91
	Female 8 to 91
	> 1 to 12 years
	Male 21 to 291
	Female 21 to 282
	> 12 to 60 years
	Male 63 to 484
	Female 65 to 421
	> 60 years
	Male 101 to 645
	Female 69 to 517
Immunoglobulin G (IgG)	Units (mg/dL)
	0 to 1 month
	Male 39/ to 1/65
	Female 391 to 1/3/
	> 1 month to 1 year
	Iviale 205 to 948
	Female 203 to 934
	> 1 to 2 years
	Wale 4/5 to 1210
	Female 483 to 1226
	> 2 to 80 years
	Iviale 540 to 1822
In many un a state ution MA (IncMA)	
Immunogiodulin IVI (IgMI)	Units (mg/aL)
	Newporn U-28 days
	28 days to 1 year

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Analyte, Blood	F	Reference range	e (Expected Val	lues)	
	F	For Pediatric va	lues, see Calip	er studies	
	Ν	Aale 17 to 143	_		
	F	emale 17 to 150)		
	>	1 to 12 years			
		/lale 41 to 183	_		
	F	emale 47 to 240)		
		• 12 years			
		/iale 22 to 240	,		
Immunadabulin E (IgE), anagifia	r	emale 33 to 29.	5		
ininiunoglobulin E (igE), specific		Specific IgE t	raditional class	ification]
		kU/L	Class	Interpretation	
		<0.10	0	None detected	
		0.10 – 0.34	0/1	Low	
		0.35 – 0.69	1	Low	
		0.70 -3.49	2	Moderate	
		3.50 – 17.49	3	High	
		17.5 – 49.9	4	Very High	
		50.0 - 99.9	5	Very High	
		>100	6	Very High	
Immunoglobulin E (IgE), Total		Age Range	Refere	nce range (kU/L)]
		0-3 months	<9		
		3-6 mos.	<1	7	
		6-9 mos	<3	0	
		9mos- 1vr	<3	9	
		1 vr - 2 vr	<5	3	
		2vr - 3vr	<9	3	
		3vr - 4vr	<1	28	
		4yr - 5yr	<11	60	
		5yr -6 yr	<1	00 02	-
		6yr - 7yr	<1	2 <u>4</u>	
		7yr - 8yr	<2	24 /8	-
		Pyr Oyr	-2	+0 80	-
		Oyr = 3yr	-20	00	-
		3yr = 10yr	0	* ~11/	-
		> 1091 - < 1	O After the pea	< 1 14 k at the age of 10	
		yıs	Arter the pea	total laE lovals decline	
			years, seruin		
		۸ مار باغ		25	
			<	14	
Iron		-14yrs Male & F	emale: 16-128		
		Fomale	· 20-162		
		19vrs Male: 65	to 175 mca/dl		
		19vrs Female: 5	50 to 170 mca/dl		
Iron Binding Capacity (TIBC)	2	250 - 450 mca/d		_	
Iron Saturation%	2	200 - 50 %			
Lactate Dehvdrogenase (LDH)	4	Adult \$19 years: 125 to 220 U/			
Lactic Acid	1	Venous 0 50 to 2 20 mmol/l			
Lipase	Ċ)-19 yrs Male &	Female: 4-39 U/	L	
		19 years: Male	& Female: 4-60	U/L	
Lipoprotein (a)	3	3.1 - 30 ma/dL			
Lithium	1	2 hr post dose (trough) concent	ration	
	1	.0 to 1.2 mmol/L	-		
Low Density Lipoprotein (LDL), Calculated	<	<130 mg/dl			
Low Density Lipoprotein (LDL), Direct	1	– 100 mg/dL			
	0	Dptimal <100 mg	j/dl		
	١	lear or above op	otimal 100-129 n	ng/dl	
	E	Borderline high 1	30-159 mg/dl		
	ŀ	ligh 160-189 mg	g/dl		

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Analyte, Blood	Reference range (Expected Values)
	For Pediatric values, see Caliper studies
	Very High >= 190 mg/dl
Magnesium	0 day to 5 months 1.5 to 2.2 mg/dL
	5 months to 6 years: 1.7 to 2.3 mg/dL
	6 to 12 years: 1.7 to 2.1 mg/dL
	12 to 18 years: 1.7 to 2.2 mg/dL
	Adult >18 years: 1.6 to 2.6 mg/dL
	NOTE: Higher values can be expected in females during
	menses
Osmolality, calculated	280-305 mOsm/Kg H ₂ 0
PBNP	<75 yr = 125 pg/mL</td
	=/>75 yr = 450 pg/mL</td
Phenytoin	10 to 20 mcg/mL
Phosphorus	M/F 0day to 15days: 5.6-10.5
	M/F 15d – 1yr: 4.8-8.4
	M/F 1-5yr: 4.3-6.8
	M/F 5-13yrs: 4.1-5.9
	13-16yrs Male: 3.5-6.2
	13-16yrs Female: 3.2-5.5
	M/F 13y – 19yrs: 2.9-5.0
	M/F >19yrs: 2.3 to 4.7 mg/dL
Potassium (K+)	Newborn 0- 7 days: 3.7 to 5.9
	Newborn 7-28 days: 3.7 to 5.9
	Infant 28 d -2 years: 4.1 to 5.3
	Child 2 yrs -12 yrs: 3.4 to 4.7
	Adult >12 years: 3.5 to 5.1
Prealbumin	0 to 1 year
	Male 7-25 mg/dL
	Female 8 -25 mg/dL
	> 1 to 12 years
	Male 11-34 mg/dL
	Female 12-30 mg/dL
	> 12 to 60 years
	Male 18-45 mg/dL
	Female 16-38 mg/dL
	> 60 years
	Male 10-42 mg/dL
Drotain Tatal (Blaama)	
Phoumatoid Factor	Addit >19 years. 0.4 to 0.5
SARS COV-2 Igo Nucleocapsiu	Negative
SARS COV-2 IgG Spike Antibody	Neuhorn 0.29doug 122 146 mmg//
Sodium (Na)	Newborn 0-200ays 133 - 146 mmol/L
	26 days to 2 years. $139 - 146 mmol/L$
	2 10 12 years. $136 - 145 mmol/L$
	12 10 90 years $130 - 145 mmol/L$
Theophylling	>90 years. 132 - 140 minol/L
Transforrin	M/E 0 day to 0wke: 104 224
	M/F 0 udy to 9wKs. 104-224 M/F 0 wks to 1/m 107.224
	1 to 14 years
	Malo 196 to 299
	Female 180 to 301
	14 to 60 years
	Male 174 to 364
	Female 180 to 382
	60 to 80 years
	Male 163 to 344
	Female 173 to 360
	>80 years:
	"The reference range for transferrin was not specifically
	verified in patients over 80 years of age"
Triolyceride	Normal < 150 mg/dl
	Borderline High 150 to 199
	High 200 to 499

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Analyte, Blood	Reference range (Expected Values)
	For Pediatric values, see Caliper studies
	Very High ≥ 500
Troponin	< 0.034 pg/mL
Tryptase	< 11.4 ug/L
	In patients with systemic mastocytosis levels of tryptase
	are, in general, persistently elevated above 20 µg/l
Uric Acid	Alinity Reference Ranges in mg/dL
	M/F 0-15days: 2.8-12.7
	M/F 15d-1yr: 1.6-6.3
	M/F 1yr-12yrs: 1.8-4.9
	Male 12-19yrs: 2.6-7.6
	Female 12-19yrs: 2.6-5.9
	>19yrs Male 3.5 to 7.2
	>19yrs Female 2.6 to 6.0
Valproic Acid	50-100 mcg/mL
Vancomycin, Random	Reference range not established
Vancomycin, Peak	25.0-40.0 mcg/mL
Vancomycin, Trough	0 to ≤ 29 days
	5.0 – 10.0 mcg/ml
	29 days & older
	5.0 – 20.0 mcg/mL

Analyte, Body Fluid	Reference range (Expected Values)
Body Fluid Albumin	Not established
Body Fluid Amylase	Not established
Body Fluid Creatinine	Not established
Body Fluid Cholesterol	Not established
Body Fluid Glucose	Not established
Body Fluid Lactate Dehydrogenase	Not established
Body Fluid Total Protein	Not established
Body Fluid Triglycerides	Not established
Synovial Fluid Total Protein	Not established

Analyte, Urine	Urine Chemistry Reference Ranges
Urine Amylase	Random: No established range
	Timed: No established range (IU/hour)
Urine Calcium	Random: No established range
	Timed: 0-300 mg/24hr
Urine Chloride	Random: No established range
	Timed: (mmol/day)
	0-30 days: 2-10
	30d – 60 years: 110-250
	> 60 years: 95 to 195
Urine Creatinine	Random
	Adult Male:
	58 to 161 mg/dL
	Adult Female:
	45 to 106mg/dL
	Timed:
	Adult Male:
	0.87 – 2.41 g/day
	Adult Female:
	0.67-1.59 g/day
	Uncorrected & Corrected Creatinine Clearance:
	Adult Male: 61 to 147 mL/min/1.73 m2 BSA
	Adult Female: 59 to 151 mL/min/1.73 m2 BSA
Urine Glucose	Random 1 to 15 mg/dL
	Timed: < 0.5 g/24hr
Urine Potassium	Random: No established range
	Timed:
	Range (mmol/day)

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Analyte, Urine	Urine Chemistry Reference Ranges
	0 day-6yrs: Not established
	6 to 10 years Male: 17 to 54
	Female: 8 to 37
	10 to 14 years Male: 22 to 57
	Female: 18 to 58
	>14 years: 25 to 125
Urine Magnesium	Random: No established reference range
	Timed: No established reference range
Urine Sodium	Random: Not established
	Timed: Range (mmol/day)
	0 day – 6 years:
	Not established
	6 to 10 years
	Male 41 to 115
	Female 20 to 69
	10 to 14 years
	Male 63 to 177
	Female 48 to 168
	Adult >14 years
	Male 40 to 220
	Female 27 to 287
Urine Phosphorus	Random:
	No established for reference range
	Timed:
	0.4 to 1.3 g/24hr
Urine	Random: Not established
Lirea Nitrogen	Timed: Not established
Urine Protein	Random:
Since Protein	7-14 mg/dl
	Timed:
	<300 mg/24br (see comment append to 24br urine
	Microalbumin
	Protein/Creatinine ratio random: <200 mg/g
	Protein/Creatinine ratio, timed: <200 mg/g
Lirine Microalbumin	Random:
	Microalhumin Quant
	5-20 mg/l
	Microalhumin/Creatining Ratio:
	0-30 mg/g
	Timed:
	Microalhumin Quant:
	$5_20 \text{ ma}/24 \text{ hr}$
	The following comment appends to Urine Microalbumin
	"Eversice within 24 hours infection fover conceptive heart
	failure marked hyperglycomia and marked hypertension
	manure, markeu nypergrycenna, and markeu nypertension
	may elevate unnary albumin excretion over baseline
	values.

+ During downtime, calculate using the 24 hour/timed urine value Excel spreadsheet available on https://www.testmenu.com/scripps under Downtime Resources

Scan the QR code:



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IMMUNOASSAYS AND ENDOCRINOLOGY

Analyte	Reference Range	Comments
AFP	2.0 - 8.78 ng/mL	This result was obtained using Abbott Alinity Chemiluminescent microparticle immunoassay (CMIA). Values obtained with different assay methods cannot be used interchangeably.
Anti-CCP	<5.0 U/mL	Alinity i Anti-CCP results should not be used interchangeably with other manufacturers' methods for anti-CCP determinations.
Anti-TPO	<5.61 IU/mL	
B-hCG, quantitative	0 - 5 mIU/mL	The Alinity i Total β -hCG assay is cleared for use in the early detection of pregnancy only. It is not approved for any other uses such as tumor marker screening, tumor marker monitoring, etc. and it should not be performed for any other uses.
B-hCG, qualitative, serum pregnancy screen	No established reference range	
B-hCG, qualitative, urine pregnancy screen	No established reference range	
CA 125 II	0 - 35 U/mL	This result was obtained using Abbott Alinity Chemiluminescent microparticle immunoassay (CMIA). Values obtained with different assay methods cannot be used interchangeably.
CA 19-9	0 - 37 U/mL	This result was obtained using Abbott Alinity Chemiluminescent microparticle immunoassay (CMIA). Values obtained with different assay methods cannot be used interchangeably.
CA 27.29 (BR Assay)	< 38.6 U/mL	This result was obtained using Siemens direct chemiluminometric technology. Values obtained with different assay methods cannot be used interchangeably.
CEA	0 - 5 ng/mL	This result was obtained using Abbott Alinity Chemiluminescent microparticle immunoassay (CMIA). Values obtained with different assay methods cannot be used interchangeably.
CK-MB	Male: 0 - 7.2 ng/mL Female: 0 - 3.4 ng/mL	
C-Peptide	0.78 - 5.19 ng/mL	
Cortisol, Random	2.9 - 19.4 mcg/dL	Patients receiving fludrocortisone, prednisolone or prednisone (which is converted to prednisolone in vivo) may show artificially elevated cortisol values due to cross-reactivity.
Cortisol AM	3.7 - 19.4 mcg/dL	Patients receiving fludrocortisone, prednisolone or prednisone (which is converted to prednisolone in vivo) may show artificially elevated cortisol values due to cross-reactivity.
Cortisol PM	2.9 - 17.3 mcg/dL	Patients receiving fludrocortisone, prednisolone or prednisone (which is converted to prednisolone in vivo) may show artificially elevated cortisol values due to cross-reactivity.
Cortisol Baseline	2.9 - 19.4 mcg/dL	Patients receiving fludrocortisone, prednisolone or prednisone may show artificially elevated cortisol values due to cross- reactivity.
Cortisol 30 min post cortrosyn	>14.0 mcg/dL	Patients receiving fludrocortisone, prednisolone or prednisone may show artificially elevated cortisol values due to cross-reactivity.
Cortisol 45 min post cortrosyn	>14.0 mcg/dL	Patients receiving fludrocortisone, prednisolone or prednisone may show artificially elevated cortisol values due to cross-reactivity.

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Analyte	Reference Range	Comments
Cortisol 60 min post cortrosyn	>14.0 mcg/dL	Patients receiving fludrocortisone, prednisolone or prednisone may show artificially elevated cortisol values due to cross- reactivity
Cortisol Post Dexamethasone	<5.0 mcg/dL	Patients receiving fludrocortisone, prednisolone or prednisone may show artificially elevated cortisol values due to cross- reactivity.
Cortisol Post Stimulation	>14.0 mcg/dL	Patients receiving fludrocortisone, prednisolone or prednisone may show artificially elevated cortisol values due to cross- reactivity.
DHEA-S	FEMALES: Age in years: Units: mcg/dL 0-11 Not established 11-15 8.6-169.8 15-20 61.2-493.6 20-25 134.2-407.4 25-35 95.8-511.7 35-45 74.8-410.2 45-55 56.2-282.9 55-65 29.7-182.2 65-70 33.6-78.9 70 and older Not established MALES: Age: Age: Units: mcg/dL 0-11 Not established 11-15 16.6-242.7 15-20 45.1-385.0 20-25 238.4-539.3 25-35 167.9-591.9 35-45 139.7-484.4 45-55 136.2-447.6 55-65 48.6-361.8 65-70 228.5-283.6 70 and older Not established	
Estradiol	Category/Phase: Units: pg/mL Normal menstruating Females: Follicular 21-251 Midcycle 38-649 Luteal 21-312 Post-menopausal Females: Not on HRT <10-28 On HRT <10-144	
Ferritin	Males: 24-44 Males: 21.81 – 274.66 ng/mL	
Fetal Fibronectin	Negative	
Folate	7.0 - 31.4 ng/mL	
FSH	Category/Phase: Units: mIU/mL Normal menstruating Females: Follicular 3.03-8.08 Midcycle 2.55-16.69 Luteal 1.38-5.47 Post-menopausal Females: 26.72-133.41 Males: 0.95-11.95	
Homocysteine	M: 5.46 - 16.20 mcmol/L	The following drugs may elevate levels of homocysteine:
(HCY)	F: 4.44 - 13.56 mcmol/L	methotrexate, carbamazepine, phenytoin, nitrous oxide, anticonvulsants and 6-azauridine triacetate. S-adenosyl- methionine is an antidepressant may interfere with the Alinity i

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Analyte	Reference Range	Comments
		Homocysteine assay
Insulin	2 – 28.0 mcgU/mL	
LH	Category/Phase: Units: mIU/mL	
	Follicular 1.80-11.78	
	Midcycle 7.59-89.08	
	Post-menopausal Females:	
	Without HRT 5.16-61.99	
Occult Blood, fecal	Males: 0.57-12.07	
(FOB)	Negative	
Parathyroid	22.4-88.2 pg/mL	
(iPTH)		
Progesterone	Category/Phase: Units: ng/mL	
	Follicular <0.1-0.3	
	Luteal 1.2-15.9	
	Post-menopausal Females: <0.1-0.2	
	Pregnant Females:	
	First trimester 2.8-147.3 Second trimester 22 5-95 3	
	Third trimester 27.9-242.5	
	Males: <0.1-0.2	
Prolactin	Males: 3.46 - 19.40 ng/mL Females: 5.18 – 26.53 ng/mL	
PSA, Total (Advia	Males:	Patients under treatment with anti-androgens, LHRH agonists,
Centaury	50 – 59 yrs 0.0 – 3.5 ng/mL	markedly reduced levels of PSA. Specimens obtained from
	60 – 69 yrs 0.0 – 4.5 ng/mL	patients undergoing prostate manipulation, especially needle
	Females: < 4.0 ng/mL	results. Care should be taken that PSA samples are drawn
		before these procedures are performed. To see full Limitations of PSA, please refer to SML menu at
		https://www.testmenu.com/scripps/Tests/329306
		chemiluminometric technology. Values obtained with different
	CE 0 III/ml Negative for IgC	assay methods cannot be used interchangeably.
Rubella, IgG	antibodies to Rubella virus	Rubella G assay, an IgG antibody capture microparticle direct
	≥ 5.0 IU/mL and ≤9.9 Equivocal	chemiluminometric immunoassay. Values obtained with
	≥10.0 IU/mL Positive for IgG antibodies to Rubella virus	different assay methods cannot be used interchangeably.
SHBG	M: 11.2 - 78.1 nmol/L	
Sirolimus	See comment	Patient assessment required. Therapeutic range dependent
		upon the type of transplant, time the dose was administered,
		and other drugs administered to the patient. Methodology: Chemiluminescent microparticle immunoassay
		(CMIA). Platform: Abbott Alinity
Tacrolimus	See comment	Patient assessment required. Therapeutic range dependent
		and other drugs administered to the patient.
		Methodology: Chemiluminescent microparticle immunoassay

CORE LABORATORY REFERENCE RANGES

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Analyte	Reference Range	Comments
Testosterone	Males (21-49 yrs): 240.24 – 870.68 Males (≥50 yrs): 220.91 – 715.81	A strong interaction with D-(-) Norgestrel (1000 ng/mL), 19- nortestosterone (Nandrolone), Ethisterone, 11b- Hydroxytestosterone, and 11-Ketotestosterone may interfere with the Abbott Alinity testosterone assay. Do not use samples from patients receiving these compounds.
	Females (21-49 yrs): 13.84 – 53.35 Females (≥50 yrs): 12.40 – 35.76	
T3, Free	1.58 - 3.91 pg/mL	
T3, Total	0.40 - 1.93 ng/mL	
T4, Free	0.70 - 1.48 ng/dL	
T4, Total	4.87 - 11.72 mcg/mL	
TSH	0.35 to 4.94 mcIU/mL	
Vitamin B12	213 - 816 pg/mL	
Vit D 250H	0-18 years:	0-18 years old:
	20-80 ng/mL	Deficiency< 20 ng/mLOptimal level≥ 20 ng/mL
	30-90 ng/mL	Potential toxicity> 100ng/mLWagner CL et al. Pediatrics 2008; 122: 1142-5218 years and older:Deficiency< 20 ng/mL
		Patients whose predominant form of vitamin D is D2, such as those receiving vitamin D2 supplementation, results that are subtherapeutic should be confirmed with another method, such as LC-MS/MS.

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INFECTIOUS DISEASE:

Assay	Reference Range	Comments	Resulting
Henatitis A	Non-Reactive		S/CO - Interpretation < 1.0 Non-Reactive
IgG			≥ 1.0 Reactive
Hepatitis A IgM	Non-Reactive	Non-Reactive: A negative test result does not exclude the possibility of exposure to hepatitis A virus. Levels of IgM anti-HAV may be below the cutoff in early infection and late acute infection.	< 0.80 Non-Reactive 0.80 – 1.20 Equivocal ≥ 1.21 Reactive*
		Equivocal: Patients exhibiting equivocal results should be closely monitored by redrawing and retesting at approximately one- week intervals.	
		*Reactive: The positive anti-HAV IgM test is consistent with recent or current Hepatitis A infection. A reactive IgM anti-HAV result does not necessarily rule out other hepatitis infections.	
Hepatitis B core Total	Non-Reactive	A nonreactive test result does not exclude the possibility of exposure to or infection with HBV.	 < 0.80 Non-Reactive 0.80 – 1.20 Equivocal <u>Retest sample in duplicate</u> <u>2 of 3 Result:</u> < 0.80 Non-Reactive 0.80-1.20 Equivocal ≥ 1.21 Reactive ≥ 1.21 Reactive ≥ 1.21 Reactive <u>2 of 3 Result:</u> < 0.80 Non-Reactive 0.80-1.20 Equivocal ≥ 1.21 Reactive 2 1.21 Reactive 2 1.21 Reactive 2 1.21 Reactive (0.80 Non-Reactive 0.80-1.20 Equivocal ≥ 1.21 Reactive (0.80 Non-Reactive 0.80-1.20 Equivocal ≥ 1.21 Reactive
Hepatitis B core IgM	Non-Reactive	Non-Reactive: A nonreactive test result does not exclude the possibility of exposure to or infection with HBV. Equivocal: Patients exhibiting equivocal results should be closely monitored by redrawing and retesting at approximately one- wook intervale	< 0.80 Non-Reactive 0.80 - 1.20 Equivocal ≥ 1.21 Reactive*

CORE LABORATORY REFERENCE RANGES

Resulting

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Assay
Reference Range
Comments

			S/CO – Interpretation
Hepatitis C Ab	Non-Reactive	Non-Reactive: A nonreactive test result does notexclude the possibility of exposure to or infection with HCV. Immunocompromised patients who have HCV may produce levels of antibody below the sensitivity of this assay and may not be detected as positive.	0.0 - 0.79 Non-Reactive 0.80 - 0.99 Equivocal <u>Retest sample in duplicate</u> <u>2 of 3 Result:</u> 0.0 - 0.79 Non-Reactive 0.80 - 0.99 Equivocal ≥ 1.0 Reactive
		Equivocal for Hep C Ab: Supplemental testing is recommended for Equivocal result. Options include: retest in 2-4 weeks or PCR Hepatitis C testing (requires new sample). Equivocal for Hep C Ab with reflex to RNA quant PCR: For Equivocal result, supplemental testing is recommended. A reflex PCR test is being performed.For Reactive for Hep C Ab: A reactive result may indicate current HCV infection, past resolved HCV infection, or a false positive result. HCV RNA testing is recommended. Reactive for Hep C Ab with reflex to RNA quant PCR: A reactive result may indicate current HCV infection, past	≥ 1.0 Reactive*
Honotitic P		false positive result. A reflex PCR test is being performed. See separate report.	< 8.0 Non Immuno to HP)/
surface Ab		The immune status of the individual should be further assessed by considering other factors, such as clinical status, follow-up testing, associated risk factors, and the use of additional diagnostic information.	 ≥ 8.0 – 11.99 Equivocal Retest sample in duplicate 2 of 3 Result: < 8.0 Non-Immune to HBV ≥ 8.0 – 11.99 Equivocal ≥ 12.0 Immune to HBV
Hepatitis B surface Ag	Non-Reactive		< 1.0 Non-Reactive ≥ 1.0 Reactive Retest sample in duplicate 2 of 3 Result: < 1.0 Non-Reactive ≥ 1.0 Reactive (Reflexes to Hep B surface Ag Confirmatory)

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Assay	Reference Range	Comments		Resulting S/CO – Interpretation
Hepatitis B surface Ag, Confirmatory	Negative	Positive results will be reported to the San Diego Department of Health Services as required by Title 17, California Code of regulations, Section 2505.		* See table below
	* Hep B surface Ag Cont	firmatory Resul	ting Table	
	DILUTION	HBsAg Qu C2 S/CO	% Neutralization	FINAL INTERPRETATION
	NEAT (UNDILUTE	< 0.70 D)	Not applicable	Not confirmed/Negative
		< 10.0	< 50 %	Not confirmed/Negative
		≥ 0.70	≥ 50 %	Confirmed Positive
		≥ 10.0	< 50 %	Repeat test using a 1:500 dilution
	1:500	< 0.70	Not applicable	Not confirmed/Negative
		≥ 0.70	≥ 50 %	Confirmed Positive
		≥ 0.70	< 50 %	Repeat test using a 1:20 000 dilution
	1:20,000	< 0.70	Not applicable	Not confirmed/Negative
		≥ 0.70	≥ 50 %	Confirmed Positive
		≥ 0.70	< 50 %	Not confirmed/Negative
HIV Ag/Ab Combo	Non-Reactive			< 1.0 Non-Reactive ≥ 1.0 Reactive* Retest sample in duplicate 2 of 3 Result: < 1.0 Non-Reactive ≥ 1.0 Reactive (Reflexes to HIVMS)
Syphilis	Non-Reactive	Non-reactive: No serologica Syphilis. If rea suspected, ref	I evidence of cent exposure test in 2-4 weeks.	< 1.0 Non-Reactive ≥ 1.0 Reactive*

*Reactive or positive results will be reported to the San Diego Department of Health Services as required by Title 17, California Code of regulations, Section 2505.

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PEDIATRIC REFERENCE RANGES (CALIPER STUDY)

AFP (ng/mL)		
AGE	FEMALE/ MALE	
0 - 1 month	>2000	
1 - 6 months	10 - 1359	
6 months – 1 year	0 - 103	
1 - 19 years	2 - 35	

FERRITIN (ng/mL)

AGE	FEMALE	AGE	MALE
0 – 14 days	99.6 – 717	0 – 14 days	99.6 – 717
15 days – 6 months	14 – 647.2	15 days – 6 months	14 – 647.2
6 months – 1 year	8.4 – 181.9	6 months – 1 year	8.4 – 181.9
1 – 5 years	5.3 – 99.9	1 – 5 years	5.3 – 99.9
5 – 14 years	13.7 – 78.8	5 – 14 years	13.7 – 78.8
14 – 19 years	5.5 – 67.4	14 – 16 years	12.7 – 82.8
		16 – 19 years	11.1 – 171.9

FREE T3 (pg/mL)

AGE	FEMALE	MALE
0 – 1 year	2.32 – 4.87	2.32 – 4.87
1 – 12 years	2.79 – 4.42	2.79 – 4.42
12 – 15 years	2.5 – 3.95	2.89 – 4.33
15 – 19 years	2.31 – 3.71	2.25 – 3.85

FSH (mIU/mL)

AGE	FEMALE	AGE	MALE
0 - 30 days	Not established	0-30 days	Not established
30 days – 1 year	0.4 - 10.4	30 days – 1 year	0.11 – 2.4
1 – 9 years	0.4 – 5.5	1 – 5 years	≤ 0.9
9 – 11 years	0.4 - 4.2	5 – 10 years	≤ 1.6
11 – 19 years	0.3 – 7.8	10 – 13 years	0.4 - 3.9
		13 – 19 years	0.8 - 5.1

LH (mIU/mL)

AGE	FEMALE	MALE
0 – 4 days	Not established	Not established
4 days – 3 months	≤ 2.4	0.2 – 3.8
3 months – 1 year	≤ 1.2	≤ 2.9
1 – 10 years	≤ 0.3	≤ 0.3
10 – 13 years	≤ 4.3	≤ 4.3
13 – 15 years	0.4 - 6.5	≤ 4.1
15 – 17 years	≤ 13.1	0.8 – 4.8
17 – 19 years	≤ 8.4	0.9 – 7.1

PROGESTERONE (ng/mL)

AGE	FEMALE	MALE
0 – 4 days	Not established	Not established
4 days - 1 year	≤ 1.32	≤ 0.66
1 - 10 years	≤ 0.35	≤ 0.35
10 – 15 years	0.5 - 0.85	0.5 - 0.85
15 - <19 years	0.5 - 10.26	0.5 - 0.57

TOTAL T3 (ng/mL)

AGE	FEMALE	MALE
0 – 4 days	Not established	Not established
4 days - 1 year	0.85 - 2.34	0.85 - 2.34

CORE LABORATORY REFERENCE RANGES

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AGE	FEMALE	MALE
1 - 12 years	1.13 - 1.89	1.13 - 1.89
12 – 15 years	0.98 - 1.76	0.98 - 1.76
15 - 17 years	0.92 - 1.42	0.94 - 1.56
17 - 19 years	0.90 - 1.68	0.90 - 1.68

TOTAL T4 (mcg/dL)

AGE	FEMALE	MALE	COMMENT
0 – 7 days	Not established	Not established	Reference range not established for this Abbott Alinity assay for 0- 7 days of age. The reference range for 7d-1yr is: 5.9-13.7 mcg/dL
7 days – 1 year	5.9 – 13.7	5.9 – 13.7	
1 – 9 years	6.2 – 10.3	6.2 – 10.3	
9 – 12 years	5.5 – 9.3	5.5 – 9.3	
12 – 14 years	5.1 - 8.3	5.0 - 8.3	
14 – 19 years	5.5 – 13.0	4.7 – 8.6	

TESTOSTERONE (ng/dL)

AGE	FEMALE	AGE	MALE
0 – 4 days	Not established	0 – 4 days	Not established
4 days - 9 years	4.33 - 62.0	4 days - 6 months	4.33 – 299
9 - 13 years	≤ 28.2	6 months - 9 years	≤ 36
13 – 15 years	10.4 - 44.4	9 – 11 years	≤ 23
15 - 19 years	14.1 - 49.0	11 - 14 years	≤ 444
		14 – 16 years	36 - 632
		16 - 19 years	148 - 794

TSH (mcIU/mL)

AGE	FEMALE/ MALE	COMMENT
0 – 4 days	Not established	Reference range not established for the Abbott Alinity assay for 0- 4 days of age. The reference range for 4 days to < 6 months is: 0.73-4.77 mclU/mL
4 days – 28 days	0.73 – 4.77	
28 days – 6 months	0.73 – 4.77	
6 months – 14 years	0.7 – 4.17	
14 – 19 years	0.47 – 3.41	

VITAMIN B12 (pg/mL)

AGE	FEMALE/ MALE
0 – 1 year	259 – 1576
1 – 9 years	283 – 1613
9 – 14 years	252 – 1125
14 – 17 years	244 – 888
17 – 19 years	203 – 811

COMMENTS APPENDED TO RESULTS FROM PEDIATRIC PATIENTS:

- A. Pediatric reference ranges not validated *for this Alinity assay*. For discussion of pediatric reference ranges, see Critical reviews in clinical laboratory science (2017), 54:6,358-413, <u>https://doi.org/10.1080/10408363.2017.1379945</u>
 - CA125
 - CA 19-9
 - CEA
 - FOLATE
 - INSULIN
 - Intact PTH
 - Cortisol, Random
 - ATPO

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- **B.** Pediatric reference ranges not validated for this Alinity assay. For discussion of pediatric reference ranges, see:
 - 1. Critical reviews in clinical laboratory science (2017), 54:6,358-413, https://doi.org/10.1080/10408363.2017.1379945
 - 2. Clin Chem Lab Med (2021), 59(10):1680-87, https://doi.org/10.1515/cclm-2021-0337
 - FREE T4

• Scripps

- PROLACTIN
- **C.** Pediatric reference ranges not well-established for the Abbott Alinity assay. See references (1,2).
 - 1. Critical reviews in clinical laboratory science (2017), 54:6,358-413, https://doi.org/10.1080/10408363.2017.1379945
 - 2. Clin Chem Lab Med (2021), 59(10):1680-87, https://doi.org/10.1515/cclm-2021-0337
 - HOMOCYSTEINE
 - SHBG

CORE LABORATORY REFERENCE RANGES

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IMMUNOLOGY TESTS

TEST	REFERENCE RANGE	
Borrelia burgdorferi (Lyme), IgG/IgM LYM	=/<0.90 OD RatioNegative0.91 to 1.09 OD RatioEquivocal=/>1.10 OD RatioPositiveNote: Positive results will reflex to Western Blotsupplemental assay and are reported to SDPH asrequired by Title 17, CCR Sec.2505	
Anticardiolipin antibodies, IgG and IgM ACLP	Cardiolipin IgM<20	
Cytomegalovirus, IgG CMVG	=/<0.90 OD Ratio Negative 0.91 to 1.09 OD Ratio Equivocal =/>1.10 OD Ratio Positive	
Cytomegalovirus, IgM CMVM	=/<0.90 OD RatioNegative0.91 to 1.09 OD RatioEquivocal=/>1.10 OD RatioPositive	
Epstein-Barr Virus Ab Panel without Early Antigen EBVPL	No detectable antibody to EBV IgG, EBV IgM, EBV EBNA IgG Index Value (IV)	
Includes: Viral Capsid Antigen IgG Viral Capsid Antigen IgM Nuclear Ag Antibodies	=/<0.90 IV Negative 0.91 to 1.09 IV Equivocal =/>1.10 IV Positive	
HerpeSelect1 ELISA IgG by Focus Technologies HSV1GG	Index Value (IV) =/<0.90 IV Negative No IgG antibodies to HSV-1 0.91 to 1.09 IV Equivocal =/>1.10 IV Positive Presumptive for the presence of IgG antibodies to HSV-1	
HerpeSelect2 ELISA IgG by Focus Technologies HSV2GG	Index Value (IV)=/<0.90 IV	
Measles (Rubeola) IgG RUBO	=/<0.90 OD Ratio Negative 0.91 to 1.09 OD Ratio Equivocal =/>1.10 OD Ratio Positive	
Mumps IgG MUMPSG	=/<0.90 OD RatioNegative0.91 to 1.09 OD RatioEquivocal=/>1.10 OD RatioPositiveIndicates past or current infection with Mumps Virusor prior vaccination against Mumps Virus.	



CORE LABORATORY REFERENCE RANGES

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TEST	REFERENCE RANGE
Varicella-Zoster Virus IgG VRCZ	=/<0.90 OD Ratio Negative for IgG antibodies to VZV. Indicates no current or previous infection with VZV. Non-Immune 0.91-1.09 OD Ratio Equivocal. Should be retested. =/>1.10 OD Ratio Positive for IgG antibodies to VZV. Indicates past or current VZV infection. Immune.
QuantiFERON (QTB)	Interpretation <0.35 IU/ml Negative = >0.35 IU/ml Positive CD4 Lymphocyte Reactivity (TB1-NIL) 0.0 to 0.34 IU/ml CD4 and CD8 Lymphocyte Reactivity (TB2-NIL) 0.0 to 0.34 IU/ml Positive results are reported to San Diego Public Health Department as required by Title 17, CCR Sec.2505
Rapid Plasma Reagin	Non-Reactive
	Interpretative comments will be printed on report.
	Reactive RPR results are reported to San Diego Public Health Department as required by Title 17, CCR Sec.2505
Free light chain/ratio Kappa Quantitative Free Light Chain	3.30 – 19.40 mg/L
Lambda Quantitative Free Light Chain	5.71 – 26.30 mg/L
Kappa/Lambda Free Light Chain Ratio (calculated)+	0.26 – 1.65
	Normal = titer of 1:32 or less
Ther at 4 C, 22 C, 37 C	Elevated = 1:64 or greater
Cryoglobulin	Negative
Infectious Mononucleosis (Mono test)	Negative
HIV GEENIUS Supplemental Assay HIV1-Ab	Non-Reactive
HIV2-Ab	Non-Reactive
	Note: Reactive results are reported to SDPH as required by Title 17, CCR Sec.2505
ANA Ab by EIA	
Antinuclear Antibodies Note: If positive, will reflex to ANA titer and pattern by Immunofluorescent antibody method (ANAH)	Negative

CORE LABORATORY REFERENCE RANGES

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TEST	REFERENCE RANGE	
SMRNP Ab If positive, Smith Antibody will be repeated. RNP antibody will be sent out to ARUP (see RNPAB)	Negative	
Smith Ab	Negative	
Sjogren's Ab (SSA Ab/SSB Ab)	Negative	
Scleroderma Ab SCL-70	Negative	
ANA Hep2 Cell, Quant	Negative at 1:40	
Pattern	Pattern not present	
DNA Ab	Negative at 1:10	
Liver Kidney Microsomal Ab (LKMA)	Negative at 1:20	
Antimitochondrial Ab (AMITA)	Negative at 1:20	
Anti-smooth muscle Ab (ASMA)	Negative at 1:20	
Anti-parietal cell Ab (APARC)	Negative at 1:20	
	Total Protein 6.3 - 8.2 g/dL	
	Albumin 3.3 - 4.4 g/dL	
Protein Electrophoresis, Serum	Alpha 1 0.1 - 0.3 g/dL	
	Alpha 2 0.4 - 1.0 g/dL	
(Abnormal scan will reflex to Immunofixation)	Beta 0.8 - 1.3 g/dL	
	Gamma 0.8 - 1.7 g/dL	
	See pathologist interpretation	
Protein Electrophoresis, Urine (Abnormal scan will reflex to Immunofixation)	See pathologist interpretation	



CORE LABORATORY REFERENCE RANGES

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HEMATOLOGY

COMPLETE BLOOD COUNT - Adult Reference Ranges

For reference ranges for other age groups or gender not specified, go to <u>https://www.testmenu.com/scripps</u> and click on Downtime Resources or scan the QR Code for CBC Ranges

To obtain absolute cell count, multiply the WBC count by the % of differentiated cell (ex. Absolute Neutrophil count = WBC count x % neutrophil), or access the downtime calculation spreadsheet for Calculations.





CBC PARAMETER	UNIT	MALE	FEMALE
WBC	10 ³ /ul	3.40 -11.0	3.40 -11.0
RBC	10 ⁶ /ul	4.46 - 5.85	3.98 -5.25
HGB	g/dl	13.0 – 17.1	11.9 – 15.3
НСТ	%	39.8 – 51.5	37.3 - 46.7
MCV	fl	81.0 -	100.0
MCH	pg	26.0	-33.0
MCHC	g/dl	31.0	-36.0
RDW-CV	%	11.7 -	- 14.9
Platelets	10 ³ /ul	150	- 425
MPV	10 ³ /ul	9.0 -	12.8
% Neutrophils	%	38 -	- 74
% Lymphocytes	%	16	- 48
% Monocytes	%	4.9 -	12.5
% Eosinophils	%	0.4 – 9.5	
% Basophils	%	0.2 – 1.6	
% Band	%	0 - 6	
% Immature Granulocytes (IG)	%	0.0 – 1.2	
NRBC	%	0.0	
Reticulocyte %	%	0.9 – 2.4	
Reticulocyte	10 ⁶ /ul	0.044 – 0.115	
Immature Retic Fraction (IRF) %	%	2.7 - 13.5	
Ret-He	pg	30.1 – 37.3	
Immature Platelet Fraction (IPF) %	%	0.9 – 9.7	
IPF	10 ³ /ul	2.8 – 19.5	
Absolute Neutrophil Count*	10 ³ /ul	1.5 – 7.4	
Absolute Lymphocyte Count*	10 ³ /ul	0.9 – 3.1	
Absolute Monocytes Count*	10 ³ /ul	0.26 – 0.87	
Absolute Eosinophils Count*	10 ³ /ul	0.03 - 0.51	
Absolute Basophils Count*	10 ³ /ul	0.01 – 0.09	
Absolute Band Count*	10 ³ /ul	0.0 - 6.0	
Absolute IG Count*	10 ³ /ul	0.00 - 0.10	

HEMATOLOGY MANUAL TESTS

TEST	REFERENCE RANGE	TEST	REFERENCE RANGE
ERYTHROCYTE SED RATE (ESR)	AGE MALE FEMALE <13 0-10 0-10 mm/hr 14-50 0-15 0-20 mm/hr >50Y 0-20 0-30 mm/hr	Viscosity, Serum (ratio)	1.4-1.8
Urine Eosinophil	NONE SEEN		

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ROUTINE AND SPECIAL COAGULATION

TEST	REFERENCE RANGE	TEST	REFERENCE RANGE
PROTHROMBIN TIME (Seconds)	10.0-13.1 (>= 18Yr) 8.8-12.5 (6 mos to < 18 yr) 8.8-14.7 (0-6 mos)	Activated Protein C (FV Leiden)	Greater than 2.2 ratio
INR	Therapeutic: 2.0 – 3.0 conventional anticoagulation 2.5-3.5 intensive anticoagulation	Anti-thrombin	83 – 128%
ACTIVATED PTT (Seconds)	26-38 (>= 18Yr) 25-39 (<18yr) Therapeutic: 53-87 seconds	Protein C Activity	70 – 140%
FIBRINOGEN (mg/dl)	187-416 mg / dL (>= 18Yr) 150-400 mg/dL (<18yr)	Factor II Activity	79 – 131%
INHIBITOR SCREEN (PT or PTT mixing studies)	PT or PTT – same ranges as above See pathologist interpretation on separate report	Factor V Activity	62 – 139%
D-DIMER (DDQ)	< 500 ng/mL FEU (>= 18Yr) <= 570 ng/mL FEU (<18Yr) Manufacturer studies indicate a D- Dimer value <500 ng/mL FEU has a high negative predictive value for DVT or PE in clinically low risk ambulatory patients. A value ≥500 ng/mL FEU warrants further studies to exclude DVT or PE.	Factor VII Activity	50 – 129%
PLATELET FUNCTION	EPI: 73-190 seconds EPI result >170 second will reflex to ADP	Factor VIII Activity	50 – 150%
ASSAY (PFA)	ADP: 65-118 seconds	Factor IX Activity	65 – 150%
	See pathologist interpretation	Factor X Activity	77 – 131%
		Factor XI Activity	65 – 150 %
		Factor XII Activity	50 – 150%
		Factor XIII Screen	Stable



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URINE TESTS

URINALYSIS	REFERENCE
	RANGE
Clarity	Clear
Color	Yellow
Glucose	Negative (mg/dL)
Ketones	Negative (mg/dL)
Bilirubin	Negative
Blood	Negative
Protein	Negative
Nitrite	Negative
Specific Gravity	1.005 – 1.030
рН	5.0 - 8.5
Urobilinogen	<2.0 mg/dL
Leukocyte Esterase	Negative
WBC	0 – 2 / HPF
RBC	0 – 2 / HPF
EPITHELIAL CELLS	None/LPF
CASTS	None/LPF
MUCUS	None/LPF
BACTERIA	None/HPF
CRYSTALS	None/LPF

URINE TOXICOLOGY DRUGS OF ABUSE SCREEN	Reference Range	Negative Threshold (cutoff concentration)
AMPHETAMINE	Negative	500 ng/ml
BARBITURATE	Negative	200 ng/ml
BENZODIAZEPINE	Negative	150 ng/ml
COCAINE	Negative	150 ng/ml
METHADONE	Negative	200 ng/ml
METHAMPHETAMINE	Negative	500 ng/ml
OPIATE	Negative	100 ng/ml
OXYCODONE	Negative	100 ng/ml
PCP	Negative	25 ng/ml
ТСА	Negative	300 ng/ml
THC	Negative	50 ng/ml
BUPRENORPHINE	Negative	10 ng/ml

Result comments:

This method provides screening results for medical purposes only. A more specific alternate chemical method must be used for a confirmed analytical result. Clinical considerations and professional judgment should be applied to any drug of abuse result, particularly when screening positive results are used.

Contact laboratory to order confirmatory testing.

*Positive results will not automatically reflex. The provider must order confirmatory test for positive result, if desired. Call the laboratory Customer Services at (858) 554-9552 to add the test.



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SEMEN ANALYSIS

SEMEN ANALYSIS	Reference Range
*Days of abstinence	2 - 7
*Volume	> 1.4 ml
*Appearance	2 - 3 turbidity, no unusual color
*Liquefaction	Liquefaction < or = 30 min
*1 hr progressive motility %	> 31%
*Motility	Motility > 4.7 mil/mL
*Motile sperm/ejaculate	> 7.1 million /ml
*pH	7.2-8.0
*Viscosity	Pours drop by drop
*Agglutination	NONE
WHO Normal Morphology % Normal	3.9 %
Abnormal head (ABHEAD)	No reference range established
Other forms (SMNOTH)	
Immature forms (IMFORM)	
Germ Cells	< 4.00 million / mL
Leukocytes	0 - 5 / HPF
Sperm Count	>14.9 million /ml
Sperm Count, Post Vasectomy	No sperm seen on wet mount. The presence of
	sperm may be below the limit of detection. A
	concentrating technique was not performed.

* Semen analysis Part I performing labs

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BODY FLUIDS

SYNOVIAL FLUID ANALYSIS	Reference Range
Color	Yellow, light yellow, straw, colorless
Appearance	Clear
Nucleated Cell Count	0-200/mcl
RBC Count	<15000/mcl
Glucose	None established
Protein	None established
Segmented Cells (% Neutrophils)	0-25 %
Lymphocytes %	None established
Mononuclear Cells %	None established
Crystals	No crystals