

Ref. By : DR S. S. PADHI

Address :

54 Years / Male Reg No. : 18486

Reg. Date : 29/07/2022 07:38AM

Collected At : MedZone Center

INVESTIGATION REPORT

CLINICAL BIOCHEMISTRY

<u>TEST</u>		<u>RESULT</u>	<u>UNIT</u>	BIOLOGICAL	REF RANGE	TEST METHOD
Glucose - Fasting						
Sample Type		: PLASMA - N	aF			
Blood Glucose-Fasting GOD/POD)	(Methodology :	: 110	mg/dl	70 - 110		
<u>Glucose - PLBS</u>						
Sample Type		: PLASMA - N	aF			

Glucose - PLBS : 125 mg/dL 70 - 140



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<u>TEST</u>	<u>RESULT</u>	<u>UNIT</u>	BIOLOGICAL REF RANGE	TEST METHOD		
Glycosylated Hemoglobin (GHb/HBA1c)						
Sample Type	: WB - EDTA					
Glycosylated Hemoglobin (GHb/HBA1c)	: 6.5	%	4.8 - 6.0 : Non Diabetic 6.0 - 7.0 : Good Control 7.0 - 8.0 : Weak Control More than 8 : Poor Control	Biorad D10 HPLC		

Glycosylated hemoglobin (*hemoglobin A1c, HbA1c, A1C, or Hb1c*; sometimes also *HbA1c*) is a form of hemoglobin used primarily to identify the average plasma glucose concentration over prolonged periods of time. It is formed in a non-enzymatic pathway by hemoglobin's normal exposure to high plasma levels of glucose. Glycation of hemoglobin has been associated with cardiovascular disease, nephropathy and retinopathy in diabetes mellitus. Monitoring the HbA1c in type-1 diabetic patients may improve treatment. HbA1c is a weighted average of blood glucose levels during the preceding 120 days, which is the average life span of red blood cells. A large change in mean blood glucose can increase HbA1c levels within 1-2 weeks. Sudden changes in HbA1c may occur because recent changes in blood glucose levels in the 30 days immediately preceding blood sampling contribute 50% to the HbA1c level, whereas glucose levels in the preceding 90-120 day period contribute only 10%. Thus, it does not take 120 days to detect a clinically meaningful change in HbA1c following a significant change in mean plasma glucose level.

METHOD: Ion Exchange Chromatography High performance liquid chromatography(HPLC)

INSTRUMENT: D -10 Bio-Rad Laboratories;FRANCE



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TEST	<u>RESULT</u>	<u>UNIT</u>	BIOLOGICAL REF RANGE	TEST METHOD
Lipid Profile				
Sample Type	: SERUM			
Cholesterol Total	: 111.9	mg/dl	Desirable : < 200 Moderate Risk : 200 - 239 High Risk :> 240	CHOD-PAP
Cholesterol HDL	: 42.66	mg/dl	Desirable : > 37 Moderate Risk : 25 - 37 High Risk : < 12 - 18	Direct Clearance
Cholesterol LDL	: 43.46	mg/dl	Desirable : < 130 Moderate Risk : 130 - 159 High Risk :> 160	Direct Clearance
Cholesterol VLDL	: 25.78	mg/dl	6 - 40	
Triglycerides	: 128.9	mg/dl	< 160 : Normal 160 - 400 : Slightly Elevated 400 - 600 : Elevated > 600 : Highly Elevated	GPO
T.Chol / HDL Chol Ratio	: 2.62		2.9 - 5.1	
LDL / HDL Ratio	: 1.02		1.7 - 3.5	

NOTE : Lipid Profile RANGES AS PER NCEP-ATP III are:

Serum cholesterol (Total) : Desirable : < 200 mg/dl, Borderline : 200 - 239 mg/dl, Elevated : >/= 240 mg/dl Serum high density lipoprotein cholesterol(HDL) : Desirable : > 60 mg/dl, Borderline : 40- 60 mg/dll, Elevated : 40 mg/dl Total cholesterol : HDL cholesterol ratio : Low risk : 3.3-4.4, Average risk : 4.4-7.1, Moderate risk : 7.1-11.0, High risk : >11.0 Serum low density lipoprotein (LDL) cholesterol : Desirable : 100 mg/dl, Borderline : 100- 159 mg/dll, Elevated : >/= 160 mg/dl Triglycerides : Desirable : 150 mg/dl, Borderline : 150- 199 mg/dll, High : 200 - 499 mg/dl, Very High : >/= 500 mg/dl HDL measurement done by Direct HDL clearance method (CDC approved). As per the Friedwald Equation, VLDL & LDL values are not applicable for triglyceride values above 400 mg/dl.



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CLINICAL BIOCHEMISTRY

<u>TEST</u>	<u>RESULT</u>	<u>UNIT</u>	BIOLOGICAL REF RANGE	TEST METHOD
Thyroid Profile				
Sample Type	: SERUM			
Tri lodothyronine (T3)	: 1.09	ng/mL	0.6-2.7 : 1 - 10 Years 0.6-1.81 : Adults Pregnancy 0.9 - 3.0 : Ist Trimester 0.9 - 3.6 : 2nd & 3rdTr	ECL
Total Thyoxine (T4)	: 7.31	µg/dL	7.8 - 16.5 : 1 - 12 Months 4.6 - 11.6 : Adults 9.1 - 14.0 : Pregnancy (15 - 40 Weeks)	ECL
Thyroid Stimulating Hormone (TSH)	: 0.387	μIU/mL	0.52 - 16.0 : 1 - 30 Days 0.46 - 8.10 : 1 Mn - 5 Yrs 0.37 - 4.8 : Adults Cord blood : 2.3 - 13.2	ECL

Three common ways in which there may be inadequate amounts of the thyroid hormone for normal metabolism. **1.** Primary hypothyroidism, in which there is a raised TSH and a low T4 and low T3. This is due to failure of the thyroid gland, possibly due to autoantibody disease, possibly due to toxic stress or possibly due to iodine deficiency. **2.** The second, the most common cause of thyroid failure, occurs at the pituitary level. In this condition there is inadequate thyroid stimulating hormone (TSH) produced from the pituitary and so one tends to see low or normal TSH, low T4s and variable T3s. This condition is most common in many patients with chronic fatigue syndrome, where there is a general suppression of the hypothalamic-pituitary-adrenal axis. **3.** The third type of under-functioning is due to poor conversion of T4 to T3. This requires enzymes and co-factors, in particular selenium, zinc and iron. In this condition there are normal or possibly slightly raised levels of TSH, normal levels of T4 but low levels of T3. This requires micronutrients and also T3 to correct.

Therefore, in any patient suspecting of thyroid problem routinely TSH, a Free T4 and a Free T3 are also advisable. Any patients who are taking T3 as part of their thyroid supplement need to have their T3 levels monitored as well as T4. T3 is much more quickly metabolized than T4 and blood tests should be done between 4-6 hours after their morning dose.

METHOD: One-step sandwich and competitive FEIA INSTRUMENT: TOSHO AIA-360 JAPAN



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RENAL FUNCTION TEST				
Sample Type	: SERUM			
Blood urea	: 27.3	mg/dl	10-40	Urease UV
Serum Creatinine	: 0.78	mg/dl	0.6-1.4	Alkaline Picrate
Blood Urea Nitrogen	: 12.75	mg/dl	7-21	
Serum Sodium	: 138	mmol/L	136-145	ISE
Serum Potassium	: 5.08	mmol/L	3.5-5.1	ISE
chloride	: 104.6	Meq/L	96-106	

Sample Registered On 29/07 Sample Received On 29/07 Report Released On 29/07 Sample Barcode :

: 29/07/2022 07:38AM : 29/07/2022 07:40AM : 29/07/2022 03:09PM

--- End Of Report ---

Checked By:gopal

Dr. VANDANA CHANDANI



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INVESTIGATION REPORT

HAEMATOLOGY

TEST	RESULT	<u>UNIT</u>	BIOLOGICAL REF RANGE	TEST METHOD
CBP (Complete Blood Picture)				
Sample Type	: WB - EDTA			
Haemoglobin	: 12.3	gm%	12.0 - 18.0	
Total Erythrocyte Count	: 4.90	M/cmm	4.0 - 6.2	Cell Counter
Hemotocrit (PCV)	: 39.8	Vol %	35.0 - 50.0	
Mean Corpuscular Volume	: 81.2	fL	80 - 100	
Mean Corpuscular Hemoglobin	: 25.1	PG	26 - 34	
МСНС	: 30.9	g/L	31 - 35	
RDW	: 14.2	%	11.5 - 14.5	
Total Leucocyte Count.	: 6750	/cumm	4000 - 11000	
DIFFERENTIAL COUNT :				
Neutrophils	: 45	%	40 - 75	
Lymphocytes.	: 44	%	20 - 40	Cell Counter
Monocytes.	: 06	%	2 - 10	Cell Counter
Eosinophils	: 05	%	1 - 6	Cell Counter
Basophils	: 0	%	0 - 1	Cell Counter
Platelet Count	: 276000	/cmm	150000 - 450000	

ESR (Erythrocyte Sedimentation Rate)

Sample Type	: PLASMA -Na	Citrate		
ESR (Erythrocyte Sedimentation Rate)	: 08	mm/hr	0 - 15 :1st Hour	Sedimentation me

		End Of Report	
Sample Registered On	: 29/07/2022 07:38AM		
Sample Received On	: 29/07/2022 07:40AM		Dhain.
Report Released On	: 29/07/2022 01:09PM		Dr. VANDANA CHANDANI
Sample Barcode :		Checked By:RAVI	