

Patient NAME : Dummy	Report STATUS : Final Report
DOB/Age/Gender :	Barcode NO :
Patient ID / UHID :	Sample Type : Whole blood EDTA
Referred BY :	Report Date :
Sample Collected :	

Test Description	Value(s)	Unit(s)	Reference Range
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Absolute Eosinophil Count (AEC)

TLC <i>Electrical impedance and microscopy</i>	5	10 ³ /μl	4 - 10
Eosinophils <i>Flow-cytometry DHSS</i>	3.4	%	0 - 5
Eosinophils. <i>Calculated</i>	0.17	10 ³ /μl	0.02 - 0.5

Interpretation:
Increased eosinophil count is often associated with allergic reaction, parasitic infestation, mycosis, collagen vascular disorders, some diffuse skin diseases, brucellosis and in hematopoietic diseases like CML, AML, Hodgkin's disease, T cell lymphoma, eosinophilic leukemia.

*** End Of Report ***

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