

TEST CHANGE

Inflammatory Bowel Disease Differentiation Panel

3003748, IBD-PAN			
Specimen Requirements:			
Patient Preparation:	N/A		
Collect:	Serum Separator Tube (SST).		
Specimen Preparation:	Separate from cells ASAP or within 2 hours of collection. Transfer 1.5 mL serum to an ARUP <u>standard transport</u> <u>tube.Standard Transport Tube.</u> (Min: 0.6 mL)		
Transport Temperature:	Refrigerated.		
Unacceptable Conditions:	Contaminated, heat-inactivated, hemolyzed, or severely lipemic specimens.		
Remarks:	N/A		
Stability:	After separation from cells: Ambient: 48 hours; Refrigerated: 2 weeks; Frozen: <u>30 days1 year</u> (avoid repeated freeze/thaw cycles)		
Methodology:	Semi-Quantitative Indirect Fluorescent Antibody (IFA)/Semi- Quantitative Enzyme Immunoassay (EIA)		
Performed:	Sun-Sat		
Reported:	1-4 days		
Note:	This test may be a useful tool for distinguishing ulcerative colitis (UC) from Crohn disease (CD) in patients with suspected inflammatory bowel disease. ANCA IFA is simultaneously tested on ethanol- and formalin-fixed slides to allow differentiation of C- and P-ANCA patterns.		
CPT Codes:	86036; 86671 x2		
New York DOH Approval Status:	This test is New York DOH approved.		
Interpretive Data:			
Refer to report.			
Reference Interval:			



Test Number	Components	Reference Interval	
	S. cerevisiae Antibody, IgG		
		20.0 Units or less	Negative
		20.1-24.9 Units	Equivocal
		25.0 Units or greater	Positive
	S. cerevisiae Antibody, IgA	/	
		20.0 Units or less	Negative
		20.1-24.9 Units	Equivocal
		25.0 Units or greater	Positive
	ANCA IFA Titer	Less than 1:20	
	ANCA IFA Pattern	None Detected	