Laboratory Diagnosis of Tropical Fever

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Tropical diseases are the diseases which are prevalent in the tropics but usually not seen in temperate climates. These may be vector borne or occur due to conditions of high humidity and temperature or sudden temperature changes which favour growth of particular microorganisms.

Diagnosis and management of tropical infectious diseases is important as they usually run a dramatic course if not treated. They may be caused by a variety of pathogens including bacteria, viruses, parasites and fungi; hence an accurate diagnosis leads to appropriate management. Diagnosis is also important from epidemiological purview to assess the burden of these diseases and monitor the effectiveness of national and international health programmes.

Recently the Indian society of Critical care medicine formulated certain guidelines and recommended a 'syndromic approach' to diagnosis and treatment of critical tropical infections. They have identified five major clinical syndromes: undifferentiated fever, fever with rash / thrombocytopenia, fever with acute respiratory distress syndrome (ARDS), fever with encephalopathy and fever with multi organ dysfunction syndrome.

Diagnosis of Tropical Diseases:

Diagnosis of a tropical infectious disease may require certain clues or hints which include a properly obtained history, epidemiological factors, recent travel and presenting clinical features 1. These infections have been categorised as arthropod-borne, rodent- associated, reservoir associated or human-human spread. An initial diagnosis may be based on the basis of interval between the exposure and the appearance of first symptom; which may be Short (\leq 10 days), Intermediate (7 – 28 days), Long (> 4 weeks) or Variable (weeks to years).

A full blood count and examination of blood smears is a nearly obligatory basic investigation. This may be accompanied by biochemical examinations like liver or kidney function tests or CSF examinations as the case may be. However, definitive diagnosis of any infectious disease relies on the microbiological investigations resulting in confirmation.

Laboratory Diagnosis in a Microbiology Laboratory:

Laboratory diagnosis may detect an organism directly by visualization under a microscope or by growing them in culture media. Culture of an organism and further identification by means of various tests proves the identity of the causative agent in an infectious disease, hence considered Gold standard. It also helps to test the organisms for susceptibility to antimicrobial agents under laboratory conditions. However, not all organisms can be cultured or identified routinely or results may not be available for days or weeks. For these agents, indirect methods of diagnosis are considered. These include serological or molecular methods. Serological tests include agglutination tests such as latex agglutination, enzyme immunoassays, Western blot,

precipitation tests, and complement fixation tests, and molecular tests may be nucleic and non-nucleic acid–based identification tests. In most Microbiology Laboratories, microscopic examination, culture facilities and some serological tests are available; other tests are done in special/research Microbiology Laboratories.

A list of bacterial, viral, parasitic and fungal diseases along with their causative agents, source of infection and relevant investigations are presented in Table 1.

Table 1: Tropical Infectious Diseases:

Tropical Dis- ease	Causative Agent	Source of Infection	Sample required	Investiga- tions	Labo- ratory Type	Sensitivity	Specificity (%)
Bacterial Tropica	al Diseases						
Legionellosis ²	Legionella pneumoph- ila	Cooling towers, Humidifiers,	Blood, Serum, Re- spiratory	Direct fluores- cent antibody staining	Special	25-70	90
		Respiratory therapy equipment, Potable/ hot water systems	Urine	Indirect fluorescent antibody test (IFA)	Special	78-91	>99
				Cultures of sputum, low- er respiratory tract secre- tions, tissue, blood	Special	10-80	100
				Urinary antigen	Routine	70-90	>99
				Polymerase chain reaction (PCR)	Special	30- 100	>90

Leptospirosis ³	Leptospira spp.	Urine, body fluids, or organs of infected animals, or by contaminated soil or water)	Blood, CSF, body fluids or tissues	Culture of body fluids or tissues like liver, muscle, kidney, skin, eyes. GOLD Stan- dard	Special	6-28	100
				Microscopic agglutination testing (MAT)	Special	86-96	>98
				IFA	Special	64	>95
				Lateral flow immunochro- matographic test	Routine	87	70
				PCR	Special	55	82
Melioidosis ⁴	Burkholderia pseudomallei	Inoculation, inhalation	Blood, Serum, urine, spu- tum, skin	Culture of specimen: GOLD Standard	Special	51-68	100
			lesions/ abscesses, throat/ rectal swabs	Gram stain, Immunoflu- orescence microscopy	Special	40-90	>90
				Indirect haemagglu- tination test, titres	Special	63-95	74-97
				IgM ELISA	Special	80	95
				PCR	Special		

Meningococcal	Neisseria meningitidis		Cerebro- spinal	CSF Gram stain	Routine	60-90	>97
Disease ⁵	lets/discharg- es from nose and throat of patients	fluid (CSF), Blood, Serum, Skin rash aspirates	Blood and CSF cultures: GOLD Stan- dard	Routine	70-85	100	
		and healthy carriers		Antigen detection in CSF/Serum by latex ag- glutination	Routine	50-93	>99
			-	Smears/ culture from petechiae	Routine	60-70	>90
				PCR	Special	91-94	>96
Q fever ⁶	fever ⁶ Coxiella Zoonosis. burnetii Cattle, sheep, goats or infected hu- mans through Inhalation, tick bites, un- pasteurized milk and milk products.	Whole blood, serum,, CSF, pleu- ral fluid, bone/ liver	Culture of affected tissue: GOLD Standard	Bio- safety level 3 (BSL 3) labora- tories	15-53	100	
		pasteurized milk and milk	biopsy/ excised heart valve, milk, placenta or foetal	Increased Phase I and II IgM, IgG titres by Mi- croimmuno- fluorescence	Special	58- 100	92-99
		i i	tissue	Increased Phase II IgM and IgG titres by ELISA	Special	80-84	>97
				Microaggluti- nation	Special	81	98
				Immunohis- tochemistry of tissue	Special	71	>90
				PCR	BSL 3	84	100

Tuberculosis ⁷	Mycobacteri- um tubercu- losis	Infected person	Sputum & other respiratory specimens,	Ziehl Neelsen Fluorescence Microscopy	Routine Special	20-80 30-90	>90 >90
			abscess, blood, bone mar-	Solid Culture- LJ Media	BSL 3	24	100
			row, body fluids, urine, gas-	Liquid Cul- ture: GOLD Standard	BSL 3	41	100
			tric lavage, faeces	Mantoux test	Clinical	65-94	50-95
				PCR	Special	43-98	90-99
				Serological tests	Not recom- mend- ed in India.	60-70	40-50

The presence of acid-fast-bacilli (AFB) on a sputum smear or other specimen often indicates TB disease. At least two sputum smears should be examined in a case of suspected pulmonary tuberculosis. A positive culture for M. tuberculosis confirms the diagnosis of TB. Culture examinations should be completed on all specimens, regardless of AFB smear results.

Typhoid and Paratyphoid fever ⁸	Salmonella enterica se- rotype Typhi,	Water or food contaminated by faeces of	Blood, bone mar- row, urine,	Culture of blood, bone marrow-	Routine	40- 90	100
	Paratyphi A, B or C	, ,	stool, Serum	Widal Test	Routine	88	70-80
				IgM Detection against S. Typhi	Routine	78-96	76-90
				Anti lipopoly- sachharide (LPS) haem- agglutination	Routine	60	98.2
				Antigen detection by ELISA or co-agglutina- tion	Routine		25-90
				PCR	Special	69-85	98- 100

Culture is the **GOLD Standard** of diagnosis. Sensitivity of blood culture varies according to the amount of blood cultured, number of specimens, antibiotic therapy, and timing of specimen collection. The sensitivity of culture is 85-90% for bone marrow, 40-50% for blood, around 60% for rose spots, 40-60% and <10% for stool and urine cultures, respectively.

Typhus: Scrub Typhus ⁹	Orientia tsut- sugamushi	Trombiculid mites	Blood, serum, biopsy or	Weil-Felix OXK aggluti- nation	Routine	89	89
			_	Scrub Ty- phus-Rapid Immunochro- matographic test	Routine	74-96	86-99
Louse-borne or Rickettsia prow louse (Pediculu bite wound	vazekii by excre	ement of body		Murine Typhus Immunoblot test	Routine	51-91	87- 100
Murine or Ende sia typhi, transm Flea (Xenopsyll	itted by Bite or e	excreta of Rat		Histopathological examination of tissue sections by Giemsa or Gimenez staining	Special	53-75	100
				IFA: GOLD Standard	Special	46- 100	78- 100
				Indirect Immunoperoxidase staining	Special	50- 100	80- 100
				Cell Culture	Special	29-59	100
				PCR	Special	<1 PFU/ ml	100

Viral Tropical D	iseases						
Avian influ- enza ¹⁰	Influenza A H5N1	Direct or close contact with infected	Throat /nasal swabs or	Viral Culture GOLD Stan- dard	BSL 3	100	100
		poultry	aspirates	Real-time reverse tran- scription-PCR	Special	100	100
				IFA test	Special	70- 100	80- 100
				Rapid Anti- gen Detection	Routine	70-75	90-95
Chikungunya ¹¹	ngunya ¹¹ CHIK virus of genus Alpha-virus, family transmitted	Serum, plasma or whole	IgM Antibody Capture (MAC) ELISA	Routine	84- 100	>99	
	Togaviridae	by bite of Aedes aegyptiand Aedes	blood	IFA	Special	75- 100	>99
		albopictus mosquitoes	-	Lateral flow immunochro- matography	Routine	10- 100	>95
				Culture GOLD Stan- dard	BSL 3	79- 100	100
				PCR	Special	0.001- 1 PFU /ml	100

Crimean-Con- go Haemor- rhagic Fever ¹²	Genus Nairo- virus, Family Bunyaviridae	Ticks and livestock animals, close contact with the blood, secretions, or-	Serum, Blood, Body flu- ids, Tissue Biopsy	IgM ELISA Antigen detection IFA	Routine Special	75-97 50- 100 75- 100	100 100 97- 100
		gans or other bodily fluids of infected persons		Pseu- do-plaque reduction neutralization	Special	98	100
				Reverse transcriptase polymerase chain reaction (RT-PCR)	Special	79-83	100
				Virus isolation by cell culture GOLD Stan- dard	BSL 3	Poor	100

Dengue ¹³	Genus Flavivirus of	Arboviral infection	Whole Blood,	RT-PCR	Special	80-90	89- 100
	the family Flaviviridae	by bite of Ae- T	Serum, Tissues	MAC ELISA	Routine	90	98
		des aegypti and Aedes		IgG ELISA	Routine	91	99
		albopictus mosquitoes		IgM Rapid test	Not Recom- mend- ed	21-99	77-98
				NS1 Antigen Detection	Routine	71- 100	98- 100
				Viral isolation	BSL 3	<50%	100
				GOLD Stan- dard			
				Plaque reduction and Neutraliza- tion test	Special	96	93-95
				Immunocyto- chemistry	Special	100	91
				Mosquito inoculation	Special	98- 100	100

During the initial five days, the virus can be detected in serum, plasma, circulating blood cells and other tissues and virus isolation in cell culture, detection of viral RNA by nucleic acid amplification tests (NAAT), or by detection of viral antigens (NS1) by ELISA can be done. At the end of the acute phase of infection, IgM antibodies appear in 50% of patients by days 3-5 after onset of illness, increasing to 80% by day 5 and 99% by day 10. A four-fold or greater increase in antibody levels measured by IgG ELISA or haemagglutination inhibition test in paired sera indicates an acute or recent flavivirus infection. During a secondary dengue infection IgG is detectable at high levels, even in the acute phase. Early convalescent stage IgM levels are significantly lower in secondary infections than in primary ones.

Haemorrhagic fever with re- nal syndrome ¹⁴	Genus Hantavirus of family Bunya- viridae	Aerosolized rodent excreta	Blood, tissue	IgM Rapid immunochro- matography test	Routine	80-97	90- 100
				IgM ELISA	Special	94	99
				IgM IFA	Special	96- 100	99
				Viral isolation by Cell culture	Special	80-95	100
				GOLD Stan- dard			
				RT-PCR	Special	94	100
Hepatitis A ¹	Genus Hepatovi-	Contaminated food or	Serum, Faeces	IgM anti-HAV ELISA	Routine	100	99
	rus Family: Picornavir- idae	water		RT-PCR GOLD Stan- dard	Special	-	100

Hepatitis B 1, 15	Genus Ortho- hepadnavi- rus, Family Hepadnavi-	Parenteral transmission, infected injec- tion needles,	Blood-Se- rum or Plasma	HBsAg Rapic Immunochro matographic test	1	94.5- 100	91- 100
	ridae	vertical and sexual trans- mission		HBsAg Latex Agglutination		66	98
				HBsAg ELISA	Routine	96-98	98- 100
				Anti-HBs ELISA	Routine	94-98	98- 100
				HBeAg ELISA	Routine	98-99	98- 100
				Anti-HBe ELISA	Routine	90-96	98- 100
				Anti-HBc ELISA	Routine	92-96	98- 100
				HBV RT- PCF GOLD Stan- dard	Special	90-95	100
		Laborat	ory Markei	s for Hepatiti	s B		•
Condition	HBsAg	HBeAg	HBV DNA	Anti HBs	Anti HBe	-	I Anti IBc
Acute Infection	+	+	+	-	-		+
Chronic Infection	+	+	+/-	-	-		+
Fulminant hepatitis	+/-	+	+	-	-		+
Vaccinated person	+#	-	-	-	-		-
Infection immunity	-	-	-	+	+/-		-
Healthy carrier	+	-	-	-	+		+

Hepatitis C ¹⁶	Flavivirus	transmission,	Blood	ELISA HCV Core Antigen	Routine	90-95	100
		infected nee- dles, vertical and sexual transmission		Recombinant immunoblot assay	Special	78	90
				ELISA Anti- HCV (IV generation)	Routine	99- 100	>99
				Saliva-based anti-HCV	Routine	87	99
			HCV RNA PCR GOLD Standard	Special	96	99- 100	
Hepatitis E ¹⁷	Genus Hepe- virus, Family		Blood, stool	ELISA HEV IgM	Routine	52-91	74- 100
	Hepeviridae	taminated water.		ELISA HEV IgG	Routine	60-91	96-98
				IgM HEV Immunoblot	Routine	95	100
				IgG HEV Immunoblot	Routine	97	85
				HEV PCR GOLD Stan- dard	Special	83- 100	100

Human Immunodeficiency virus (HIV) ¹⁸ Lentivirus, family Retroviridae	an HIV in- fected person through sex- ual or vertical transmission,	Whole blood, se- rum. Saliva and urine are not	HIV-1/2 Ab Rapid test Serum HIV- 1/2 ELISA	Routine Routine	99- 100 99- 100	98- 100 97- 100	
	mucocutaneous or parenteral exposure	taneous or parenteral	being used for testing in India.	HIV-1and HIV-2 Ab/ HIV-1 p24 antigen	Routine	100	>99
			HIV-1/2 Ab (Oral fluids)	Routine	54- 100	67- 100	
				HIV-1 Urine	Routine	92- 100	95- 100
			IFA an- ti-HIV-2	Special	93-99	98- 100	
			HIV Western Blot	Special	100	100	
			HIV DNA PCR GOLD Standard	Special	90-96	54- 100	

After Pre-test counselling, NACO guidelines for testing are followed. Three different kits with different antigen system and / or different principles of tests are required. If the first test is negative, the patient is considered non-reactive. If the test serum is reactive with two tests and non-reactive with the third, it is reported as "indeterminate" and patient is called back for repeat testing after 2-4 weeks. The test used as the screening test is one with the highest sensitivity and the supplementary second and third tests are with the highest specificity. If all the 3 tests are reactive, post-test counselling is done and then the patient is referred to ART centre for treatment. For confirmation and viral load determination, molecular tests are done.

Parasitic Tropical Diseases								
Amoebiasis ¹⁹ Entamoeba histolytica		Food or water contaminated with faeces containing infectious cysts	Stool, Abscess fluid, serum	Stool Micros- copy	Routine	10-60	10-50	
				Microscopy (abscess fluid)	Routine	<25	10-50	
				Culture with isoenzyme analysis	Special	<60	100	
				GOLD Stan- dard				
				HK-9 antigen detection (ELISA)	Routine	65- 100	>90	
				Abscess antigen detection (ELISA)	Routine	100	>90	
				Stool antigen detection (ELISA)	Routine	>95	>95	
				Serum antibody detection (ELISA)	Routine	70-90	85-90	
			PCR (stool)	Special	>70	>90		

Leishmania donovani Leishmania donovani Arthropod borne (Sandfly bite), Zoonotic in some countries	borne (Sand- fly bite), Zoo-	Blood, bone marrow,	Microscopy of leucocytocon-centrates	Routine	<80	>80	
	lymphoid tissues, Serum	Histological and impres- sion smear examination	Special	48-76	>80		
		Culture from Buffy coats GOLD Stan- dard	Special	<80	100		
		Antigen detection ELISA	Routine	98	96-99		
		IFA	Special	81	100		
			Western blots	Special	88	100	
				PCR	Special	88-95	100

Malaria ²¹ Plasmodium vivax, P. falciparum, P. malariae, P. ovale, P. knowlesi	Bite of infected mosquitoes, rarely by transfusion.	Blood, Serum	Microscopy Thin blood film	Routine	100 para- sites/ µl	100	
			Microscopy Thick blood film	Routine	10-20 para- sites/ µl	100	
				Fluorescent Microscopy	Special	81- 100	86- 100
				Quantitative buffy coat examination	Special	41-93	93
				Immunofluorescence (1:128)	Special	> 90	> 90
				P. falciparum Anti-HRP-2 antibody test	Routine	77-98	83-98
			Plasmodium pLDH or Aldolase Rapid test at 100-500/µL of blood	Routine	85- 100	98- 100	
			Culture	Special	-	100	
			PCR (1-100 parasites / μl of blood)	Special	95- 100	95- 100	

Microscopic examination of malarial parasites is considered **GOLD Standard** of diagnosis. Serological tests are approved only for emergencies and places where microscopy is not possible.

	gondii oocysts s in cat's fa vertical t mission,	Ingestion of oocysts shed	ts shed s faeces, affected tissues tissues	IgM ELISA	Routine	>93	90- 100
		in cat's faeces, vertical trans-		IgE ELISA	Routine	76	98
		mission, rare- ly infected		IgG ELISA	Routine	>99	>99
		blood/ organ		Western Blot	Special	99	100
	donation	donation		PCR for prenatal diagnosis	Special	90-92	>99
			PCR of pla- cental tissue	Special	42-71	98- 100	
			PCR (Blood, CSF) in cerebral toxo- plasmosis	Special	33-83	98- 100	
			IFN-γ release assay	Special	94	98	
PCR is considere	ed the GOLD St a	andard for Diagi	nosis.				
Fungal tropical	diseases						
Cryptococco- sis ²³	Cryptococcus neoformans Inhalation or inoculation of basidiospores	inoculation of	CSF, Blood, Serum, Urine, Sputum	Microscopy (India ink preparation)	Routine	50-90	>90
				Culture	Rou-	50-90	100
				GOLD Stan- dard	tine/ Special		
			Cryptococ- cal antigen detection	Routine	83-97	93- 100	
			Antibody detection by ELISA	Routine	80-85	100	
				PCR	Special	92- 100	100

Depending on the provisional clinical diagnosis of the abovementioned diseases, relevant investigations can be done according to the available facilities.

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