

Lab Use Only

☐ **Acceptance Criteria Not Met**

- ☐ Inappropriate temperature
- ☐ Specimen too old
- ☐ Incomplete labeling/form
- ☐ Specimen inappropriate/damaged

Date: / / Initials:

Attach Printed Label Below

Patient Information	Last Name					
	First Name		MI			
	Maiden Name/Surname					
	Address/Attention:					
	Street Address:			Address 2:		City:
	State:	Zip Code:	County Code:	County Name:		Phone Number:
	Insurance ID Number: (if applicable)			Medicaid Number (if applicable):		
	Medical Record Number:			Date of Birth: ____/____/____		If Female, Pregnant? ☐ Yes ☐ No ☐ Unknown
	Sex: ☐ Male ☐ Transgender M2F ☐ Female ☐ Transgender F2M ☐ Unknown ☐ Transgender Unknown ☐ Ambiguous			Race (mark all that apply): ☐ White ☐ American Indian/ Alaska Native ☐ Black ☐ Native Hawaiian/ Pacific Isles ☐ Asian ☐ Unknown		Ethnicity: ☐ Hispanic or Latino Origin ☐ Non-Hispanic ☐ Unknown
Submitter	EIN: _____ - _____		Submitter Name:			
	Address:		Address 2:		City:	
	State:		Zip Code:		County Name:	
	Phone Number:		Email Address:		Fax Number:	
	Ordering Provider NPI:		Ordering Provider First and Last Name:			
Specimen (continued on page 2)	Specimen source(s):		Collection Date(s) and Time(s):		Collector's Initials:	Laboratory Number(s): <i>Do Not Write in this Space</i>
	☐ Acute Serum (within 7 days of onset)		____/____/____ :____ 24 Hr Time			
	☐ Convalescent Serum		____/____/____ :____ 24 Hr Time			
	☐ Whole Blood		____/____/____ :____ 24 Hr Time			
	☐ CSF		____/____/____ :____ 24 Hr Time			
	☐ Urine		____/____/____ :____ 24 Hr Time			
	☐ Amniotic Fluid		____/____/____ :____ 24 Hr Time			
	Onset Date: ____/____/____					Reason for Testing (ICD-10 Dx Code): _____
	Serologic Diagnostic Panels Available: (Check one or more boxes, as needed)					
	☐ Arboviral IgM ELISA Panel (Eastern Equine Encephalitis, La Crosse Encephalitis, and West Nile)					
☐ Rickettsia PCR Panel (<i>Rickettsia rickettsii</i> , <i>Rickettsia prowazekii</i> , <i>Rickettsia</i> species)						

Specimen (continued from page 1)	Exanthems: <i>(All suspect cases must be approved for testing by the Communicable Disease Branch (CDB) prior to submission of specimen to the State Lab. CDB can be reached at 919-733-3419. Testing will be sent to reference laboratory)</i> ☐ Measles ☐ Rubella ☐ Varicella Zoster, IgG ☐ Mumps, IgG										
	Single Agent Diagnostic Tests: (Check one or more boxes, as needed) ☐ Dengue IgM ELISA and PCR (if specimen collection is within 7 days of symptom onset) ☐ Chikungunya IgM ELISA and PCR (if specimen collection is within 7 days of symptom onset) ☐ Zika PCR **The Physician Attestation (below) must be signed prior to testing.** ☐ Other: _____ ☐ Prior approval/consultation received from: _____ ☐ Please forward specimen to CDC for testing. (Attach a completed CDC 50.34 DASH form).										
Other Patient Information	Patient Signs and Symptoms: <i>(Check all that apply)</i> <table style="width: 100%; border: none;"> <tr> <th style="text-align: left; width: 20%;">General</th> <th style="text-align: left; width: 20%;">Rash</th> <th style="text-align: left; width: 20%;">Respiratory</th> <th style="text-align: left; width: 20%;">CNS</th> <th style="text-align: left; width: 20%;">Cardiovascular</th> </tr> <tr> <td> ☐ Fever to ____°F ☐ Headache ☐ Fatigue ☐ Sore Throat ☐ Jaundice ☐ Conjunctivitis ☐ Arthralgia/Myalgia ☐ Nausea/Vomiting </td> <td> ☐ Macular ☐ Papular ☐ Vesicular ☐ Petechial ☐ Focal ☐ Hemorrhagic </td> <td> ☐ Cough ☐ Pneumonia ☐ Bronchitis ☐ Croup ☐ Pharyngitis </td> <td> ☐ Seizures ☐ Meningitis ☐ Encephalitis ☐ Nuchal rigidity ☐ Paralysis </td> <td> ☐ Chest Pain ☐ Pericarditis ☐ Myocarditis ☐ Pleurodynia </td> </tr> </table> <i>If pregnant, due date:</i> ____ / ____ / ____	General	Rash	Respiratory	CNS	Cardiovascular	☐ Fever to ____°F ☐ Headache ☐ Fatigue ☐ Sore Throat ☐ Jaundice ☐ Conjunctivitis ☐ Arthralgia/Myalgia ☐ Nausea/Vomiting	☐ Macular ☐ Papular ☐ Vesicular ☐ Petechial ☐ Focal ☐ Hemorrhagic	☐ Cough ☐ Pneumonia ☐ Bronchitis ☐ Croup ☐ Pharyngitis	☐ Seizures ☐ Meningitis ☐ Encephalitis ☐ Nuchal rigidity ☐ Paralysis	☐ Chest Pain ☐ Pericarditis ☐ Myocarditis ☐ Pleurodynia
	General	Rash	Respiratory	CNS	Cardiovascular						
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Recent Vaccination History: _____ _____ _____ Travel History: Area(s): _____ _____ Dates: _____											
Physician Attestation for Zika Testing	Zika virus assays are intended for use with specimens collected from individuals meeting CDC Zika virus clinical criteria (e.g., clinical signs and symptoms associated with Zika virus infection) and/or CDC Zika virus epidemiological criteria (e.g., history of residence in or travel to a geographic region with active Zika transmission at the time of travel, or other epidemiologic criteria for which Zika virus testing may be indicated as part of a public health investigation). NCSLPH provides testing to patients when the following criteria are met: • A pregnant woman who: ➢ Has ongoing possible Zika virus exposure ➢ Has had prenatal ultrasound findings consistent with congenital Zika infection • An individual with symptoms associated with Zika virus infection (rash, joint pain, fever, and/or conjunctivitis) who: ➢ Spent time in an area with risk for Zika virus transmission, or ➢ Had unprotected sex with a partner who spent time in an area with risk for Zika virus transmission										
	☐ I certify that the patient I am requesting Zika testing for meets the criteria outlined above.* Physician Name (Print) _____ Physician Signature _____ * For further guidance regarding eligibility for Zika testing, please visit the Zika Virus Testing page on the NCSLPH website at https://slph.dph.ncdhhs.gov/zika/										