U.S. WHO Influenza Collaborating Laboratories Testing Methods Assessment

	Lab Name: Lab ID Number:	
1.	What types of samples does your lab receive for influenza virus testing and what is the relative proportion of each? Please exclude proficiency testing samples. (check all that apply and give the approximate percent for each type checked) Original patient specimens% Viral isolates% Other% Specify:	
2.	What influenza testing methods does your lab offer? (check all that apply) Commercial rapid diagnostic Viral culture Immunofluorescent antibody testing RT-PCR Other Specify:	
3.	What best describes the origin of specimens your lab receives? Please rank order the following sources from 1 (source from which you obtain the most specimens) to 6 (source from which you receive the least specimens). Sentinel surveillance sites Local health departments Managed care Private physicians Hospitals Other Specify:	
4.	Approximately what proportion of the test results you report to CDC are obtained using PCR? None 1-24% 25-29% 50-74% 50-75-99% All	
5.	Is your laboratory currently testing all specimens for <i>both</i> influenza A and B viruses? ☐ Yes, all specimens are tested for both influenza A and B ☐ No, we are testing only for influenza A ☐ All specimens are tested for influenza A and only influenza A negative samples are tested for influenza B ☐ Other ☐ Specify: ☐ Specify:	
6.	 Are you fully subtyping all influenza A viruses identified by your laboratory? Yes, we are subytping all influenza A viruses to identify both 2009 influenza A (H1N1) and seasonal influenza A viruses. No, we identify only 2009 influenza A (H1N1) but don't further subtype those which are influenza A positive. No, we identify 2009 influenza A (H1N1) and test only a subset of influenza A positive and 2009 influenza A (H1N1) negative viruses for seasonal influenza subtypes No, we attempt subtyping on only a subset of influenza A viruses. No, we are not currently subtyping any influenza A viruses. Other 	

CDC 55.31A 9-95 This report is authorized by law (Public Health Service Act, 42 USC 241). Public reporting burden for this collection of information is estimated to average 10 minutes per response, including time for reviewing instructions, searching existing data sources, gathering and measuring the data needed, and completing and returning the collection of information. Send comments regarding the burden estimate or any other aspect of the collection of information, including suggestions for reducing the burden to PHS Reports Clearance Officer; ATTN: PRA, Hubert H. Humphrey Bldg., Rm 721-B; 200 Independence Ave., SW; Washington, DC 20201, and to the Office of Management and Budget; Paperwork Reduction Project (0920 0004); Washington, DC 20503.

Specify: ___