



**For CDI Data Validation,
Use CDI Forms A-C**



CDI Validation - Form A Summary of Laboratory Data

Instructions: To begin validation of CDI, ask your laboratory to produce a line list directly from the laboratory information system for a 3-month time period to include

- Toxin-Positive **C difficile tests** (PCR, assay, culture) from both inpatients and ED patients.

Reports should include the patient’s hospital location (unit) when the Cdiff test was performed.

Ask to have printed twice: sorted by date and then sorted by patient name or medical record number.

Months selected for validation:

- Jan Feb Mar Apr May Jun Jul Aug Sep Oct Nov Dec

Laboratory information system (LIS): _____ e.g. Meditech, Sunquest, Cerner
IMPORTANT: For validation, do not produce this positive blood culture line list from a surveillance software system (e.g. Medmined, TheraDoc, AICE, etc) or infection control integration software programs. Get directly from LIS.

STEP 1: Indicate # of Positive **C difficile tests** from 3 mo period _____ **[INCLUDE IN CDI REVIEW]**

STEP 2: Using the lab line list sorted by name, number each positive CDI on your lab line list.

STEP 3: Enter corresponding specimens and unit locations in table on CDI Validation **Form B.**

STEP 4: Follow instructions on CDI Validation Forms B and C to complete validation review.