

Clostridioides difficile Infection (CDI) Surveillance

Hospital identified CDI case definition

A CDI episode is defined as a laboratory confirmed toxin-producing *C. difficile* infection from a diarrhoeal stool **or** evidence of pseudomembranous colitis on gross anatomic (includes endoscopic exams) or histopathologic exam on a symptomatic patient.

NOTE: Do not report positive results from screening specimens or asymptomatic patients

Inclusions

- > Cases from all patients attending an acute care facility while symptomatic (including inpatients, Hospital in the Home patients and patients attending Emergency departments, outpatient departments or haemodialysis units etc.).

Exclusions

- > Cases where a known previous positive specimen has been obtained within the previous 8 weeks (an isolate obtained from a patient more than 8 weeks since the last positive test is regarded as a new episode)
- > Patients less than 2 years old at date of attendance/admission.

CDI exposure classification

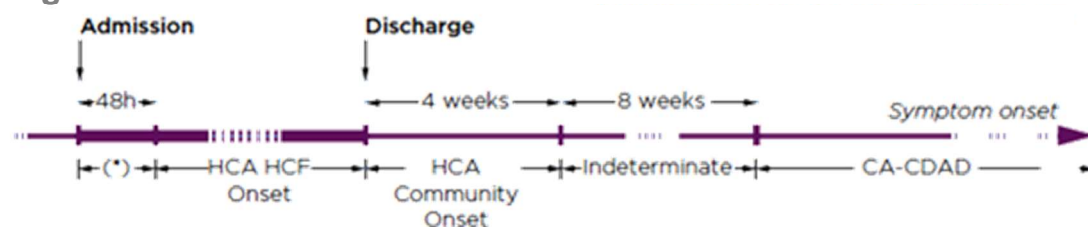
The Australian Commission for Safety and Quality in Health Care has developed an Implementation Guide for CDI surveillance that describes sub-categorisation of episodes according to their most likely place of acquisition into 5 separate categories. Available from: <https://www.safetyandquality.gov.au/our-work/healthcare-associated-infection/national-hai-surveillance-initiative>.

At the present time, South Australian contributors are requested to apply the sub-category A (HCA-HCF) exposure classification (see Figure 1 below).

Category A: Healthcare associated – Healthcare facility onset (HCA-HCF)

Date of CDI symptom onset more than 48 hours after admission to a health care facility and prior to discharge from the facility. If date of symptom onset is not available, the date/time of specimen collection is used as a proxy.

Figure 1



NOTE:

- #Identification/investigation of onset date is only required if the specimen collection date is >48hours after admission.
- Given state reporting currently only collects data on hospital identified and Category A (HCA-HCF) episodes as a sub-set, if a patient has a specimen taken during a current visit and it is identified as being associated with a previous admission, do not classify as a “post discharge” in the secondary acquisition, classify as NEW. Post-discharge information can be recorded as a comment in ICIMS.

Data Element Table

Field Name	Description	Details
UR or Postcode	Unique record identification number	<ul style="list-style-type: none"> This is the patient's medical record number (MRN) or postcode for Private hospitals that do not supply MRN Mandatory field, cannot be null
Gender	Sex of the patient	<ul style="list-style-type: none"> Mandatory field, cannot be null
Date of Birth	The patients full year of birth, including day and month	<ul style="list-style-type: none"> If date of birth is not known or cannot be provided, provision of a generic estimate is acceptable (the first day of the appropriate month or 01/01/ of the appropriate year Format date as dd/mm/yyyy Mandatory field, cannot be null
Date of Admission / Attendance	The date the patient attended the Emergency or Outpatient Dept or date the patient was admitted	<ul style="list-style-type: none"> Format date as dd/mm/yyyy Mandatory field, cannot be null
Attendance Type	Identifies if the type of hospital attendance/admission	<ul style="list-style-type: none"> Permissible values: Inpatient, Outpatient or Emergency Mandatory field, cannot be null
Ward	The ward where patient was located at time specimen was collected.	<ul style="list-style-type: none"> This is NOT the acquisition ward , only the ward at the time the specimen was taken Field should not be Null if "Attendance Type" = Inpatient
Specimen/Onset/ Diagnosis Date[#]	Identifies the date of either <ul style="list-style-type: none"> symptom onset specimen collection pseudomembranous colitis diagnosis 	<ul style="list-style-type: none"> Format date as dd/mm/yyyy Must be within the reporting month Mandatory field, cannot be null
LAB Name	Identifies the laboratory organisation that processed the specimen	<ul style="list-style-type: none"> Mandatory field, cannot be null
Specimen Number	Positive specimen's unique identification number	<ul style="list-style-type: none"> Identifier allocated by the laboratory to the pathology result Mandatory field, cannot be null
Category	Identifies record as healthcare associated where applicable	<ul style="list-style-type: none"> Permissible values: Cat A or N/A Field should not be null
Comment	Record any relevant additional information	Example: diagnosis of pseudomembranous colitis

QA NOTES:

- Outpatient and Emergency records – Attendance date should match specimen/symptom onset date*
- Category A records – ensure reported specimen/onset date is > 48hrs after admission.
- Specimen/onset date must be between admission and discharge dates

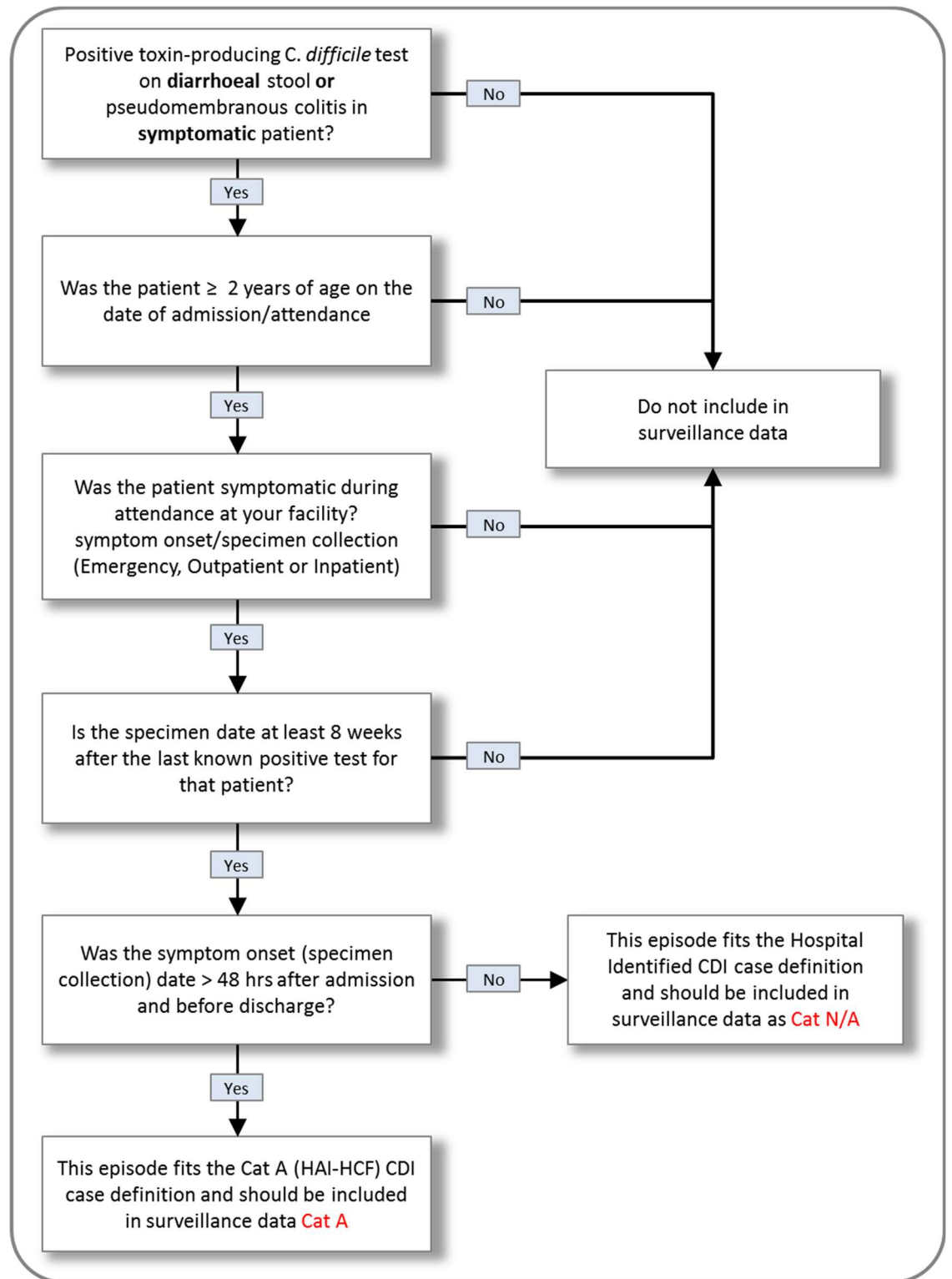
*Sources for identifying onset date may include (but are not limited to) medical notes, lab request date, specimen request form or the patient.

Careconnect.sa infection control users (ICIMS)

To ensure record is included in the Category A field select:

- Primary Acquisition = HCA
- Secondary Acquisition = New.

Flow Chart



For more information

Infection Control Service

Communicable Disease Control Branch

Telephone: 1300 232 272

www.sahealth.sa.gov.au/infectionprevention

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