

Form Approved OMB No. 0920-0666 Exp. Date: 12/31/2026 www.cdc.gov/nhsn

## **MDRO and CDI Monthly Denominator Form**

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*required for sav	ing	**conditi	**conditionally required based upon monitoring selection in Monthly Reporting Plan								
Facility ID #:		*Month:		*Year:		*Location (	Code:				
Line 1: Setting: **Total Facility	Inpatient	-			**Total Facility Admissions:						
Line 2: If your facility has a CMS-certified rehab unit (IRF) or CMS-certified psych unit (IPF), please subtract these counts from "Total Facility Patient Days" and "Total Admissions" (Line 1). If you do not have these units, enter the same values you entered on Line 1.											
Counts= [Total Facility – (IRF + IPF)]											
Patient Days: Admissions:											
Line 3: If your facility has a CMS-certified IRF, CMS-certified IPF, NICU, or Well Baby Unit, please subtract those counts from "Total Facility Patient Days" and "Total Admissions" (Line 1). If you do not have these units, enter the same values you entered on Line 1.											
Counts= [Total Facility – (IRF + IPF + NICU + Well Baby Unit)]  Patient Days: Admissions:											
**For this quarter, what is the primary testing method for <i>C. difficile</i> used most often by your facility's laboratory or the outside laboratory where your facility's testing is performed? (check one)											
☐ Enzyme immunoassay (EIA) for toxin ☐ GDH plus NAAT (2-step algorithm)											
☐ Cell cytotoxicity neutralization assay ☐ GDH plus EIA for toxin, followed by NAAT for discrepant results											
<ul> <li>□ Nucleic acid amplification test (NAAT) (e.g., PCR, LAMP)</li> <li>□ Toxigenic culture (<i>C. difficile</i> culture followed by detection of toxins)</li> </ul>											
□ NAAT plus EIA, if NAAT-positive (2-step algorithm) □ Other:											
☐ Glutamate dehydrogenase (GDH) antigen plus EIA for toxin (2-step algorithm)											
Note: "Other" should not be used to name specific laboratories, reference laboratories, generic testing methods (such as "PCR") or the brand names of <i>C. difficile</i> tests; most methods can be categorized accurately by selecting from the options provided. Please ask your laboratory or contact the NHSN helpdesk for further guidance.											
Organism Selection/Confirmation of No Events											
Specific Organism Type	MRSA	C. difficile	MSSA	CephR- Klebsiella	CRE- E. coli	CRE- Enterobacter	CRE- Klebsiella	MDR- Acinetobacter	VRE		
Infection Surveillance											
LabID Event (All specimens)											
LabID Event (Blood specimens only)											
	ntiality: The v	oluntarily prov	ided information	obtained in this su	ırveillance system	that would permit ident	ification of any individ	lual or institution is collect	ed with		

Assurance of Commentancy: The voluntarity provided information to brained in institution is collected with a guarantee that it will be held in strict confidence, will be used only for the purposes stated, and will not otherwise be disclosed or released without the consent of the individual, or the institution in accordance with Sections 304, 306 and 308(d) of the Public Health Service Act (42 USC 242b, 242k, and 242m(d)).

Public reporting burden of this collection of information is estimated to average 15 minutes per response, including the time for reviewing instructions, searching existing data sources, gathering, and maintaining the data needed, and completing and reviewing the collection of information. An agency may not conduct or sponsor, and a person is not required to respond to a collection of information unless it displays a currently valid OMB control number. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to CDC, Reports Clearance Officer, 1600 Clifton Rd., MS H21-8, Atlanta, GA 30333, ATTN: PRA (0920-0666).

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Process Measures: Hand Hygiene, Gown and Glove Use, and AST									
Hand Hygiene			Gown and Gloves						
**Performed:	**Indicated:		**Used: **Indicated:						
	-								
Active Surveillance Testing (AST)									
**Active Surveillance Testing performed									
**Timing of AST <sup>†</sup> (circle one)	Adm Both	Adm Both							
**AST Eligible Patients ‡ (circle one)	AII NHx	All NHx							
Admission AST	<u>l</u>								
**Performed									
**Eligible									
Discharge/Transfer AST	1								
**Performed									
**Eligible									
Outcome Measures: AST									
Prevalent Cases									
(Specific Organism Type)	MRSA VR		E						
**AST/Clinical Positive									
**Known Positive									
Incident Cases									
**AST/Clinical Positive									
Custom Fields									
Label									
Data									
† Adm – Admission testing B									
<b>‡ All</b> – All patients tested <b>NHx</b> – Only patients tested are those who have no documentation at the admitting facility in the previous months of MDRO-colonization or infection at the time of admission.									