

NATIONAL HEALTHCARE SAFETY NETWORK Patient Safety Component—Annual Facility Survey for LTAC

Instructions for this form are available at: http://www.cdc.gov/nhsn/forms/instr/TOI-57.150-LTAC.pdf

*required for saving Facility ID:			Tracking #: *Survey Year:		
Facility Characteristics (co	mpleted by Infection Preve	ntionist)			
*Ownership (check one):	· · ·				
□ For profit *Affiliation (check one):	\Box Not for profit, including c	hurch	□ Government	🗆 Veteral	ns Affairs
□ Hospital System	Independent		Iulti-facility organization (sp	ecialty hosp	oital network)
*Setting/classification: If classified as "Free-standing, facilities or units (check all that	" does your LTAC hospital sh	are physi	Within a hospita cal housing with one or mor		owing on-site
□ No		🗆 Inpa	tient rehabilitation facility		
\Box Skilled nursing facility ((SNF)/nursing home	🗆 Neu	ro-behavioral unit or facility		
\Box Residential facility (ass	sisted living	□ Othe	er (specify):		
If classified as "Within a hospit	al," is your LTAC hospital loc	ated:			
In a building that does no	t provide acute care services	(for exan	nple, psychiatric hospital?)	□ Yes	□ No
Near (but not within) an a	cute care hospital?			□ Yes	□ No
In the previous calendar year,	indicate:				
*Number of patient days: _ *Number of admissions: _ *Average daily census: _ *Numbers of LTAC beds in the		ries shou	ld equal total):		
c. General LTAC beds:	or critical care beds: I care/high acuity beds (not IG C beds (licensed capacity):	CU): 			
*Number of single occupancy *Number of double occupancy *Number of triple occupancy ro *Number of quadruple occupan	rooms: poms:				
Assurance of Confidentiality: The voluntarily p guarantee that it will be held in strict confidence institution in accordance with Sections 304, 30 Public reporting burden of this collection of info gathering, and maintaining the data needed, ar a collection of information unless it displays a c	e, will be used only for the purposes stated, a 6 and 308(d) of the Public Health Service Ac mation is estimated to average 102 minutes nd completing and reviewing the collection of	and will not oth t (42 USC 242) ; per response ; information.	erwise be disclosed or released without th b, 242k, and 242m(d)). CDC 57.150 (Fror including the time for reviewing instruction an agency may not conduct or sponsor, ar	ne consent of the in nt). Rev 10, v13.0 ns, searching exis nd a person is not	ndividual, or the sting data sources, required to respond t

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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	one of the one of the following con): (Note: These categories are not r		Form Approved OMB No. 0920-0666 Exp. Date: 12/31/26 www.cdc.gov/nhsn ssion (present of admission,
	ying these conditions on admission found here: <u>http://www.cdc.gov/nhs</u> 		-
Facility Microbiology Laborate	ory Practices (completed with inp	out from Microbiology L	aboratory Lead)
susceptibility testing?	s own on-site laboratory that perforr acility's antimicrobial susceptibility t ter	esting performed: (check	
 *2. For Enterobacterales, Ps methods are used for: (1) Primary susceptibility (2) Secondary, supplem 	ental, or confirmatory testing (if per ot perform susceptibility testing, indi	netobacter baumannii cor formed).	☐ Yes ☐ No nplex, indicate which
	(1) Primary (2) S	Secondary	Comments
1 = Kirby-Bauer disk diffusion	4 = ThermoFiscer/Sensititre	7 = Agar dilution me	ethod
2 = bioMérieux/Vitek	5 = Beckman Coulter/MicroScan	10 = Gradient Diluti	on Strip (for example E test)
		13 = Other (describe	e in Comments section)
3 = BD Phoenix	6 = Selux Diagnostics		
*3. Does either the primary of (check all that apply):	or secondary/supplemental antimicr	obial susceptibility testing	(AST) include the following
Drug	Tested	Not Tested	
Cefiderocol			
Ceftazidime-Avibactam			
Ceftolozane-Tazobactam			
Eravacycline			
Plazomicin			
Imipenem-Relebactam			

			Form Approved OMB No. 0920-0666 Exp. Date: 12/31/26 www.cdc.gov/nhsn	
Meropenem-Vaborbactam				
Aztreonam-Avibactam				
Sulbactam-Durlobactam				
Facility Microbiology Laboratory	Practices (continued)			
*4. Has the laboratory implemented revised breakpoints recommended by CLSI for the following:				
a. Third Generation Cephalosporin and monobactam (that is, aztreonam) breakpoints for			🗆 Yes 🛛 No	

		Enterobacterales in 2010	
	b.	Carbapenem breakpoints for Enterobacterales in 2010	🗆 Yes 🛛 No
	c.	Ertapenem breakpoints for Enterobacterales in 2012	🗆 Yes 🛛 No
	d.	Carbapenem breakpoints for Pseudomonas aeruginosa in 2012	🗆 Yes 🛛 No
	e.	Fluroquinolone breakpoints for Pseudomonas aeruginosa in 2019	🗆 Yes 🛛 No
	f.	Fluroquinolone breakpoints for Enterobacterales in 2019	🗆 Yes 🛛 No
	g.	Aminoglycoside breakpoints for Enterobacterales in 2023	🗆 Yes 🛛 No
	h.	Aminoglycoside breakpoints for Pseudomonas aeruginosa in 2023	🗆 Yes 🛛 No
	i.	Piperacillin-tazobactam breakpoints for Pseudomonas aeruginosa in 2023	🗆 Yes 🛛 No
	j.	Piperacillin-tazobactam breakpoints for Enterobacterales in 2022	🗆 Yes 🛛 No
5.	not	es the laboratory test bacterial isolates for presence of a carbapenemase? (this does include automated testing instrument expert rules) If Yes, indicate what is done if carbapenemase production is detected: (check one)	🗆 Yes 🛛 No
		□ Change susceptible carbapenem results to resistant	
		□ Report carbapenem MIC results without an interpretation	
		$\hfill\square$ No changes are made in the interpretation of carbapenems, the rest is used for epidemio	logical or

infection control practices 5b. If Yes, which test is routinely performed to detect carbapenemase: (check all that apply)

Nucleic Acid Amplification Test (PCR, Cepheid, etc.)	□ mCIM/CIM	□ NG-Test Carba-5 (or other lateral flow assay)
☐ Modified Hodge Test	🗌 Carba NP	□ Other

5c. If Yes, which of the following are routinely tested for the presence of carbapenemases: (check all that apply)

		Enterobacterales spp.	🗆 Pseudomonas aeruginosa	🗌 Acinetobacter baumannii
6.	Does y	our facility use commercial or lat	poratory developed tests for rapid molecu	lar detection of antimicrobial
	resista	nce markers in bacterial bloodstr	eam infections? Examples of commercia	lly available systems include
	BioFire	FilmArray, Luminex Verigene, e	tc.	

□ Yes

- $\hfill\square$ No [if checked, skip questions 7 and 8]
- 6a. If Yes, which test panel(s) does your facility use? (check all that apply)
 - □ Accelerate PhenoTest BC □ BioFire FilmArray BCID □ BioFire FilmArray BCID II

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NATIONAL HEALTHCARE SAFETY NETWORK Cepheid Xpert MRSA/SA BC GenMark ePlex BCID-GP	GenMark ePlex BCID-GN
□ GenMark ePlex BCID-FP □ Luminex Verigene BC-GP	Luminex Verigene BC-GN
☐ MALDI-TOF MS directly from positive blood culture (e.g., Sepsi	-
MALDI-TOF MS based antimicrobial resistance detection	
T2Biosystems T2Bacteria T2Biosystems T2Candida	□ T2Biosystems T2Resistance
Other Commercial Test(s) (Leave Comment)	· · · · · · · · · · · · · · · · · · ·
Other Laboratory Developed Test(s) (Leave Comment)	
Facility Microbiology Laboratory Practices (continued)	
*7. In a scenario where the <i>mecA</i> resistance marker and <i>Staphylococcus au</i>	ureus are detected by rapid molecular
testing in a blood specimen, select the procedure(s) your facility conduct	
\Box Our laboratory does not perform <i>mecA</i> testing using rapid molecu	ılar methods. [If checked, skip question
7a.]	
Culture based phenotypic antimicrobial susceptibility testing is no	t performed. [If checked, skip question
7a.]	
□ Culture based phenotypic antimicrobial susceptibility testing is pe	0
corresponding rapid molecular testing and/or the interpretation of the the phenotypic test result.	e rapid molecular testing result is added to
\Box Culture based phenotypic antimicrobial susceptibility testing is pe	rformed. No text indicating corresponding
rapid molecular testing and/or interpretation is added.	normed. No text indicating corresponding
7a. If both rapid molecular and culture based phenotypic antimicrobial s	usceptibility testing are performed for a
blood specimen to detect drug resistance in Staphylococcus aureus	, and discordance is found between their
results, how are results reported? (check one)	
\Box Further testing is not pursued. Results are reported separately.	
Further testing is not pursued. The phenotypic result is overridde an antimicrobial resistance marker is detected.	en by the rapid molecular test result when
 Further testing is performed to identify the reason for the discord further analysis. 	dance. Results are modified based on the
*8. In a scenario where the <i>bla_{CTX-M}</i> (CTX-M) resistance marker and <i>Escheri</i> testing in a blood specimen, select the procedure(s) your facility conduct	
Our laboratory does not perform <i>bla_{CTX-M}</i> (CTX-M) testing using ra questions 8a]	pid molecular methods. [If checked, skip
Culture based phenotypic antimicrobial susceptibility testing is no 8a.]	t performed. [If checked, skip question
Culture based phenotypic antimicrobial susceptibility testing is pe corresponding rapid molecular testing and/or the interpretation of the the phenotypic test result.	-
Culture based phenotypic antimicrobial susceptibility testing is pe rapid molecular testing and/or interpretation is added.	rformed. No text indicating corresponding
 8a. If both rapid and culture based phenotypic antimicrobial susceptibilit specimen to detect drug resistance in <i>Escherichia coli</i> and discordar are results reported? (check one) 	
□ Further testing is not pursued. Results are reported separately.	



	ALIICAL	HCARE www.cdc.gov/nhsn
SAFE	TYNET	Further testing is not pursued. The phenotypic result is overridden by the rapid molecular test result when
		an antimicrobial resistance marker is detected.
		Further testing is performed to identify the reason for the discordance. Results are modified based on the
		further analysis.
	-	
Facilit	y Micro	biology Laboratory Practices (continued)
Facilit	y Micro	biology Laboratory Practices (continued)
+acilit *9.		biology Laboratory Practices (continued) is yeast identification performed for specimens collected at your facility? (check one)
	Where	

□ Affiliated medical center

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□ Commercial referral laboratory

□ Other local/regional, non-affiliated reference laboratory

□ Yeast identification not available (specifically, yeast identification is not performed onsite or at any	y
affiliate/commercial/other laboratory) [If checked, skip questions 11-15]	

Answer questions 11-15 for the laboratory that *performs yeast identification for your facility*:

*10.Which of the following methods are used for yeast identification? (check all that apply)

	□ MALDI-TOF MS System (Vitek MS)	□ MicroScan
	□ MALDI-TOF MS System (Bruker Biotyper)	\Box Non-automated Manual Kit (for example, API 20C, RapID, Germ Tube, PNA-FISH, etc.)
	□ Vitek-2	DNA sequencing
	□ BD Phoenix	Other (specify):
*11	.Does the laboratory routinely use chromogenic	agar for the identification or differentiation of Candida isolates?
	□ Yes □ No	🗌 Unknown

*12. Candida isolated from which of the following body sites are usually fully identified to the species le	vel?	(check all
that apply)		

Blood	□ Respiratory
\Box Other normally sterile body site (for example, CSF)	Other (specify):
	\Box None are fully identified to the species level

Does the laboratory	employ any PCR	molecular tests to iden	tifv <i>Candida</i> from blo	ood specimens?

🗆 Yes	s 🗆 No	🗆 Unknown
10-	If we a which DOD me leaveler to	te ave used to identify. Condided

If yes, which PCR molecular tests are used to identify *Candida* from blood specimens? (check all that 13a. apply)

- □ T2Candida Panel
- □ BioFire BCID
- □ GenMark ePlex BCID
- Other, specify: _____
- Unknown

13b. If yes and you get a positive result, does this lab culture the blood to obtain an isolate?

□ Yes, always

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□ No				
Unknown				
Facility Microbiology Laboratory Prac	tices (continued)			
*14.Where is antifungal susceptibility		d for specimens col	lected at your facility? (check or	ne)
□ On-site laboratory	□ Other local/regiona			- /
□ Affiliated medical center	-		is not performed onsite or at ar	ny
Commercial reference laboratory			elected, skip questions 16 -20]	2
Answer questions 16-20 for the laborer and the second seco	ngal susceptibility testing	g (AFST), excludin	g Amphotericin B? (check all t	hat
Broth microdilution with laboratory developed plates	☐ YeastOne (Therm Sensititre™)	o Scientific™	Gradient diffusion (E test)	
□ Vitek (bioMerieux)	\Box Other (specify): _		Unknown	
*16.What methods are used for antifu	ngal susceptibility testing	g (AFST) of Ampho	tericin B? (check all that apply)
Broth microdilution with laboratory developed plates	☐ YeastOne (Therm Sensititre™)	o Scientific™	\Box Gradient diffusion (E test)	
□ Vitek (bioMerieux)	\Box Other (specify): _		Unknown	
*17.AFST is performed for which of th	e following antifungal dr	ugs? (check all that	apply)	
☐ Fluconazole	□ Voriconazole)	□ Itraconazole	
Posaconazole	🗌 Micafungin		🗌 Anidulafungin	
🗌 Caspofungin	Amphotericir	ו B	□ Flucytosine	
Other, specify:	Unknown			
*18.AFST is performed on fungal isola	ates in which of the follow	ving situations? (ch	eck all that apply)	
	erformed automatically	Performed with a clinician's order	Not performed Unknow	n

Blood		
Other normally sterile body site (for example, CSF)		
Urine		
Respiratory		
Other (specify):		

*19.Is this laboratory developing antibiograms or other reports to track susceptibility trends for *Candida* spp. isolates tested in this laboratory?

🗆 Yes	🗆 No	🗌 Unknown



S*20.What is the primary testing method for C. difficile 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

- □ Cell cytotoxicity neutralization assay
- □ Nucleic acid amplification test (NAAT) (for example, PCR, LAMP)

Facility Microbiology Laboratory Practices (continued)

- □ NAAT plus EIA, if NAAT positive (2-step algorithm)
- Glutamate dehydrogenase (GDH) antigen plus EIA for toxin (2-step algorithm)
- □ GDH plus NAAT (2-step algorithm)
- GDH plus EIA for toxin, followed by NAAT for discrepant results
- □ Toxigenic culture (*C. difficile* culture followed by detection of toxins)
- Other (specify):

*21.Which of the following methods serve as the primary method used for bacterial identification at your facility? (check one)

- □ MALDI-TOF MS System (Vitek MS)
- □ MALDI-TOF MS System (Bruker Biotyper)
- □ Automated Instrument (for example, Vitek, MicroScan, Phoenix, etc.)
- □ Non-automated Manual Kit (for example, API 20C, biochemicals)
- □ Rapid Identification (for example, NAAT/PCR, Gene Xpert, etc.)
- □ 16S rRNA Sequencing
- Other (specify): _____
- □ None
- *22. Which of the following methods serve as the secondary or backup method used for bacterial identification at your facility? (for example, a secondary method if the primary method fails to give an identification, or if the primary method is unavailable). (check one)
 - □ MALDI-TOF MS System (Vitek MS)
 - □ MALDI-TOF MS System (Bruker Biotyper)
 - □ Automated Instrument (for example, Vitek, MicroScan, Phoenix, etc.)
 - □ Non-automated Manual Kit (for example, API 20C, biochemicals)
 - □ Rapid Identification (for example, NAAT/PCR, Gene Xpert, etc.)
 - □ 16S rRNA Sequencing
 - Other (specify): _____
 - □ None

Infection Control Practices (completed with input from Hospital Epidemiologist and/or Quality Improvement Coordinator)

*23.Number or faction of infection preventionists (IPs) in facility:

- a. Total hours per week performing surveillance:
- b. Total hours per week for infection control activities other than surveillance:
- *24.Number or fraction of full-time employees (FTEs) for a designated hospital epidemiologist (or equivalent role) affiliated with your facility:

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s#25.1s it a p	loncy in your facility that patients infe	ected or colonized with MRSA are routinely placed in conta	
precau □ Yes	tions while these patients are in your \Box No	r facility ? (check one) □ Not applicable: my facility never admits these pati	onte
			51115
Infection Cor	trol Practices (continued)		
05-			- f :1:4 -
25a. (ch	eck one):	nat are routinely placed in contact precautions while in your	facility
	All infected and all colonized patien	nts	
	Only all infected patients		
		with certain characteristics (check all that apply)	
	\Box Patients admitted to high risk	settings	
	\Box Patients at high risk for transr		
while th	nese patients are in your facility? (ch	ected or colonized with VRE are routinely placed in contact leck one)	precautions
□ Yes	□ No	\Box Not applicable: my facility never admits these patients	
26a. (ch	If Yes, check the type of patients th eck one):	nat are routinely place in contact precautions while in your t	acility
	All infected and all colonized patien	nts	
	Only all infected patients		
	Only infected or colonized patients	with certain characteristics (check all that apply)	
	\Box Patients admitted to high risk	settings	
	\Box Patients at high risk for transr	nission	
•	enemase production) are routinely p	ected or colonized with CRE (regardless of confirmatory tes laced in contact precautions while these patients are in yo	•
🗆 Yes	□ No	\Box Not applicable: my facility never admits these patie	ents
27a. (ch	If Yes, check the type of patients th eck one):	nat are routinely placed in contact precautions while in your	⁻ facility
	All infected and all colonized patien	nts	
	Only all infected patients		
	Only infected or colonized patients	with certain characteristics (check all that apply)	
	\Box Patients admitted to high risk	settings	
	\Box Patients at high risk for transr	nission	
extend		ected or colonized with suspected or confirmed ESBL-prod <i>Enterobacterales</i> are routinely placed in contact precautione)	-
🗆 Yes	□ No	\Box Not applicable: my facility never admits these patie	ents
28a. (ch	If Yes, check the type of patients th eck one):	nat are routinely placed in contact precautions while in your	facility
	All infected and all colonized patien	nts	
	Only all infected patients		
	rent) Dov 0 v12 0		Dage 9 of 21



SAFETY NETWORK Only infected or colonized patients with certain characteristics (check all that apply)

- \Box Patients admitted to high risk settings
- \Box Patients at high risk for transmission

Infection Control Practices (continued)

*29.Does the facility routinely perform screening testing (culture or non-culture) for CRE? This includes screening for patients at your facility performed by public health laboratories and commercial laboratories.

□ Yes □ No

- 29a. If Yes, in which situations does the facility routinely perform screening testing for CRE? (check all that apply)
 - □ Surveillance testing at admission for all patients
 - Surveillance testing of epidemiologically-linked patients of newly identified CRE patients (for example, roommates)
 - □ Surveillance testing at admission of high-risk patients (check all that apply)
 - □ Patients admitted form long-term acute care (LTAC) or long-term care facility (LTCF)
 - □ Patients with recent (for example, within 6 months) overnight hospital stay outside the United States
 - \Box Patients admitted to high-risk settings (for example, ICU)
 - □ Other high-risk patients (specify): _____
 - □ Surveillance testing of all patients in the facility or in a specific high-risk settings (for example, ICU) at prespecified intervals (for example, weekly point prevalence survey)
 - □ Other (specify): _
- 29b. If Yes, what method is routinely used by the lab conducting CRE testing of screening swabs from your facility? (check all that apply)
 - □ Culture-based methods □ PCR □ Other (specify): _____
- *30.Does the facility routinely perform screening testing (culture or non-culture) for *Candida auris*? This includes screening for patients at your facility performed by public health laboratories and commercial laboratories.

 \Box Yes \Box No

- 30a. If Yes, in which situations does the facility routinely perform screening testing for *Candida auris*? (check all that apply)
 - □ Surveillance testing at admission for all patients
 - □ Surveillance testing of epidemiologically-linked patients of newly identified *Candida auris* patients (for example, point prevalence surveys in response to a case, patients in the same room or unit as a case)
 - □ Surveillance testing at admission of high-risk patients (check all that apply)
 - □ Patients admitted form long-term acute care (LTAC) or long-term care facility (LTCF)
 - □ Patients with recent (for example, within 6 months) overnight hospital stay outside the United States
 - \Box Patients admitted to high-risk settings (for example, ICU)
 - Other high-risk patients (specify):

 - □ Surveillance testing of all patients in the facility or in a specific high-risk setting (for example, ICU) at prespecified intervals (for example, weekly point prevalence survey)
 - Other (specify): _____

NHSN NATIONAL HEALTHCARE SAFE 300 FTW 9R Yes, what meth from your facility?	nod is routinely u	used by the lab condu	cting <i>Candida auris</i> te	Form Approved OMB No. 0920-0666 Exp. Date: 12/31/26 www.cdc.gov/nhsr sting of screening swabs
Culture-based m	ethods		\Box Other (specify):	
*31.Does the facility routinely				
Infection Control Practices (co	ontinued)			
31a. If Yes, in which s apply)	ituations does th	ne facility routinely pe	rform screening testing	g for MRSA? (check all that
□ Surveillance testi	ng at admission	for all patients		
		of high-risk patients [LTCF], or dialysis pat		l from long-term acute care
□ Surveillance testi	ng at admission	of patients admitted	to high-risk settings (fo	or example, ICU)
\Box Surveillance testi	ng of pre-operat	tive patients to prever	nt surgical site infection	าร
□ Other (specify): _				
*32.Does your facility have a transmission of MDROs a		ly use chlorhexidine t	bathing for any adult pa	atients to prevent infection or
	at your facility?			🗆 Yes 🛛 No
32a. If Yes, indicate w	hich patients: (s	elect all that apply)		
\Box ICU patients:		□ Patients outside	the ICU:	\Box Pre-operatively for
 All ICU patients 		 All patients ou 	itside the ICU	patients undergoing
$^{ m O}$ Subset of ICU pa	tients:	 Subset of pati 	ents outside the	surgery
Patients with c		ICU:	the control veneus	
catheter or mid			th central venous midline catheters	
\Box Other, specify:			cify:	
*33.Does the facility have a p antistaphylococcal agent prevent healthcare-assoc	(mupirocin, iodo ciated infections	ophor, or an alcohol b or reduce transmissio	ased intranasal agent) on of resistant pathoge	for any adult patients to ens? Yes I No
Antibiotic Stewardship Practic	es (completed	with input from Phy	vsician and Pharmac	ist Stewardship Leaders)
 *34.Facility leadership has de □ Providing stewardship prointerventions. □ Allocating resources (for efforts. 	ogram leader(s) o example, IT supp	dedicated time to man port, training for stewa	nage the program and ardship team) to suppo	conduct daily stewardship ort antibiotic stewardship
Having a senior executive resources and support to ac	complish its mis	sion.		
-				nd/or board at least annually.
Ensuring the stewardship board at least annually.				
Communicating to staff al	-			
Providing opportunities fo Drewiding a formula statement	•	•		•
Providing a formal statem approved by the board).	ent of support fo	or antibiotic stewardsh	np (for example, a writ	ten policy or statement



STEEP Network that staff from key support departments and groups (for example, IT and hospital medicine) are contributing to stewardship activities.

 \Box None of the above.

	inty has a leader of a		ntibiotic stewardship program management and	outcomes
				s 🗆 No
35a.		osition of this leader? (chec	,	
	Physician	\Box Co-led by both Pharn	•	
	Pharmacist		RN, PA, NP, etc.; specify):	
35b. Iea	der? (check all that a		ollowing describes your antibiotic stewardship p	hysician
	Has antibiotic stewa	ardship responsibilities in the	eir contract or job description or performance rev	view
	Is physically on-site	in your facility (either part-ti	me or full-time)	
	Completed an ID fe	llowship		
	Completed a certific	ate program on antibiotic st	ewardship	
	Completed other tra	aining(s) (for example, confe	rences or online modules) on antibiotic stewards	ship
	None of the above.			
• •) leader): What perce		their contract or job description' is selected (for particular stewardship activities is specified in the physici	-
	□ 1-10%	□ 11-25%	□ 26-50%	
	□ 51-75%	□ 76-100%	\Box Not specified	
35d. lea	•	ed is selected: In an averag otic stewardship activities in	e week , what percentage of time does the phys your facility? (check one)	i cian (co
	□ 1-10%	□ 11-25%	□ 26-50%	
	□ 51-75%	□ 76-100%		
35e. pha	If Pharmacist or Co armacist leader? (ch		e following describes your antibiotic stewardship	
			eir contract, job description or performance revie	w
	Is physically on-site	in your facility (either part-ti	me or full-time)	
		ID residency and/or ID fello		
	•	ate program on antibiotic st	ewardship	
	Completed a certific	cate program on antibiotic st aining(s) (for example, confe	ewardship rences or online modules) on antibiotic stewards	ship
	Completed a certific Completed other tra			ship
□ □ 35f. If 'H (co)	Completed a certific Completed other tra None of the above Has antibiotic steward) leader): What perce	aining(s) (for example, confe		armacist

🚞 NH	SN		Form Approve OMB No. 0920-066 Exp. Date: 12/31/2	66 26
NATIONAL HEAL SAFETY NETV	^{THCARE} ^{VORK} □ 51-75%	□ 76-100%	www.cdc.gov/nhs	<u>3n</u>
35g.	If 'Pharmacist' or 'Co-led' b) leader spend on antibioti	-	week, what percentage of time does the pharmacis	st
(00	□ 1-10%	□ 11-25%		
	□ 51-75%	□ 76-100%		
Antibiotic St	ewardship Practices (con	inued)		
35h. po	If Pharmacist or Other is s int of contact and support fo		have a designated physician who can serve as a ??	
			🗆 Yes 🛛 No	
	a pharmacist is not the lead proving antibiotic use at you		gram, is there at least one pharmacist responsible for	•
			🗆 Yes 🛛 No	
*36.Our fac	cility has the following priori	y antibiotic stewardship in	terventions: (Check all that apply)	
	spective audit and feedback			
			ntibiotic stewardship program monitors prospective ng antibiotic use, types of interventions, acceptance o	of
	,		🗆 Yes 🛛 No	
🗆 Prea	authorization for specific ant	ibiotic agents.		
	•	8		
Antibiotic St	ewardship Practices (con	inued)]
36b.		ted: Our antibiotic steward	dship program monitors preauthorization interventions] s
36b.	If Preauthorization is sele	ted: Our antibiotic steward] s
36b.	If Preauthorization is select r example, by tracking whic	ted: Our antibiotic steward h agents are requested for	which conditions).] s
36b.	If Preauthorization is select r example, by tracking whic Facility-specific treatment susceptibilities, to assist v	ted: Our antibiotic steward h agents are requested for recommendations, based vith antibiotic selections for	which conditions).] S
36b. (fo	If Preauthorization is select r example, by tracking whice Facility-specific treatment susceptibilities, to assist w community-acquired pneu	ted: Our antibiotic steward h agents are requested for recommendations, based with antibiotic selections for monia, urinary tract infecti	which conditions).] s
36b. (fo	If Preauthorization is select r example, by tracking which Facility-specific treatment susceptibilities, to assist w community-acquired pneu- If Facility-specific treatment	ted: Our antibiotic steward h agents are requested for recommendations, based vith antibiotic selections for monia, urinary tract infecti nt recommendations is sel	which conditions).	s
36b. (fo	If Preauthorization is select r example, by tracking which Facility-specific treatment susceptibilities, to assist w community-acquired pneu If Facility-specific treatment Community-acquired pneu	ted: Our antibiotic steward h agents are requested for recommendations, based vith antibiotic selections for monia, urinary tract infecti nt recommendations is sel	which conditions).	s
36b. (fo	If Preauthorization is select r example, by tracking whice Facility-specific treatment susceptibilities, to assist w community-acquired pneu If Facility-specific treatment Community-acquired pneu Urinary tract infection	ted: Our antibiotic steward h agents are requested for recommendations, based with antibiotic selections for monia, urinary tract infecti nt recommendations is sel imonia	which conditions).	S
36b. (fo	If Preauthorization is select r example, by tracking which Facility-specific treatment susceptibilities, to assist w community-acquired pneu- If Facility-specific treatment Community-acquired pneu- Urinary tract infection Skin and soft tissue infect	ted: Our antibiotic steward h agents are requested for recommendations, based with antibiotic selections for monia, urinary tract infecti nt recommendations is sel imonia	which conditions).	s
36b. (fo 36c.	If Preauthorization is select r example, by tracking which Facility-specific treatment susceptibilities, to assist w community-acquired pneu- If Facility-specific treatment Community-acquired pneu- Urinary tract infection Skin and soft tissue infect None of the above	ted: Our antibiotic steward h agents are requested for recommendations, based with antibiotic selections for monia, urinary tract infecti nt recommendations is sel imonia	which conditions). □ Yes □ No on national guidelines and local pathogens common clinical conditions (for example, on, skin and soft tissue infection). ected: For which common clinical conditions?	
36b. (fo 36c. 36c. 36d. to	If Preauthorization is select r example, by tracking which Facility-specific treatment susceptibilities, to assist w community-acquired pneu- If Facility-specific treatment Community-acquired pneu- Urinary tract infection Skin and soft tissue infect None of the above If Facility-specific treatment our facility's treatment reco	eted: Our antibiotic steward h agents are requested for recommendations, based with antibiotic selections for monia, urinary tract infecti nt recommendations is sele imonia on ht recommendations is sele	which conditions). □ Yes □ No on national guidelines and local pathogens common clinical conditions (for example, on, skin and soft tissue infection). ected: For which common clinical conditions?	
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36b. (fo 36c.	If Preauthorization is select r example, by tracking which Facility-specific treatment susceptibilities, to assist w community-acquired pneu- If Facility-specific treatment Community-acquired pneu- Urinary tract infection Skin and soft tissue infect None of the above If Facility-specific treatment our facility's treatment recon- ample, community-acquired	eted: Our antibiotic steward h agents are requested for recommendations, based with antibiotic selections for monia, urinary tract infection the recommendations is selection monia	which conditions). □ Yes □ No on national guidelines and local pathogens common clinical conditions (for example, on, skin and soft tissue infection). ected: For which common clinical conditions?	
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36b. (fo 36c. 36c. 36c. 36d. to ex. 36e.	If Preauthorization is select r example, by tracking which Facility-specific treatment susceptibilities, to assist w community-acquired pneu- If Facility-specific treatment Community-acquired pneu- Urinary tract infection Skin and soft tissue infect None of the above If Facility-specific treatment our facility's treatment recon- ample, community-acquired If Yes: For which common Community-acquired pneu-	ted: Our antibiotic steward h agents are requested for recommendations, based with antibiotic selections for monia, urinary tract infecti nt recommendations is selection imonia on ht recommendations is selection mendations for antibiotic pneumonia, urinary tract clinical conditions?	 which conditions). ☐ Yes ☐ No on national guidelines and local pathogens common clinical conditions (for example, on, skin and soft tissue infection). ected: For which common clinical conditions? ected: Our stewardship program monitors adherence selection for common clinical conditions (for infection, skin and soft infections). 	



SAFETY NEW ORK None of the above

VATIONAL HEALTHCARE

- *37.Our facility has a policy or formal procedure for other interventions to ensure optimal use of antibiotics: (Check all that apply.)
 - \Box Early administration of effective antibiotics to optimize the treatment of sepsis
 - \Box Treatment protocols for Staphylococcus aureus bloodstream infection
 - □ Stopping unnecessary antibiotic(s) in new cases of *Clostridioides difficile* infection (CDI)
 - \Box Review of culture-proven invasive (for example, bloodstream) infections
 - □ Review of planned outpatient parenteral antibiotic therapy (OPAT)
 - □ The treating team to review antibiotics 48-72 hours after initial order (specifically, antibiotic time-out)
 - □ Assess and clarify documented penicillin allergy
 - Using the shortest effective duration of antibiotics at discharge for common clinical conditions (for example, community-acquired pneumonia, urinary tract infection, skin and soft tissue infections)
 - $\hfill\square$ None of the above

Antibiotic Stewardship Practices (continued)

37a. If 'Using the shortest effective duration of antibiotics at discharge for common clinical conditions' is selected: Our stewardship program monitors adherence in using the shortest effective duration of antibiotics at discharge for common clinical conditions (for example, community-acquired pneumonia, urinary tract infections, skin and soft tissue infections), at least annually.

□ Yes □ No

- *38.Our facility has in place the following specific 'pharmacy-based' interventions: (Check all that apply)
 - □ Pharmacy-driven changes from intravenous to oral antibiotics without a physician's order (for example, hospital-approved protocol)
 - \Box Alerts to providers about potentially duplicative antibiotic spectra (for example, multiple antibiotics to treat anaerobes)
 - □ Automatic antibiotic stop orders in specific situations (for example, surgical prophylaxis)
 - \Box None of the above
- *39.Our stewardship program has engaged bedside nurses in actions to optimize antibiotic use.

□ Yes □ No

- 39a. If Yes is selected: our facility has in place the following specific 'nursing-based' interventions: (Check all that apply.)
 - □ Nurses receive training on appropriate criteria for sending urine and/or respiratory cultures.
 - \Box Nurses initiate discussions with the treating team on switching from intravenous to oral antibiotics.
 - \Box Nurses initiate antibiotic time-out discussions with the treating team.
 - \Box Nurses track antibiotic duration of therapy.
 - \Box None of the above.

*40.Our stewardship program monitors: (Check all that apply.)

- \Box Antibiotic resistance patterns (either facility- or region-specific), at least annually
- Clostridioides difficile infections (or C. difficile LabID events), at least annually
- □ Antibiotic use in days of therapy (DOT) per 1000 patient days or days present, at least quarterly
- □ Antibiotic use in defined daily doses (DDD) per 1000 patient days, at least quarterly

m NH	SN	OMB No.	n Approved 0920-0666 e: 12/31/26
	HCARE /ORK tic expenditures (specifically, purchasing costs), at least quarterly	www.cd	lc.gov/nhsn
	tic use in some other way, at least annually (specify):		
	of the above		
*41.Our ste apply.)	ewardship team provides the following antibiotic use reports to prescribers, at least annual	y: (Chec	k all that
🗆 Individ	ual, prescriber-level reports		
🗆 Unit- o	r service-specific reports		
🗆 None d	of the above		
•	If 'Individual, prescriber-level reports' or 'Unit-or service-specific reports' is selected: Our ogram uses these reports to target feedback to prescribers about how they can improve the scribing, at least annually.		•
		□ Yes	🗆 No
Antibiotic Ste	ewardship Practices (continued)		
*42.Our fac	ility distributes an antibiogram to prescribers, at least annually.		
*43.Informa annual	ation on antibiotic use, antibiotic resistance, and stewardship efforts is reported to hospital ly.	□ Yes staff, at I	
	of the following groups receive education on optimal prescribing, adverse reactions from a tic resistance (for example, Grand Rounds, in-service training, direct instruction) at least a apply.)		s, an
🗆 Prescr	ibers		
🗆 Nursin	g staff		
🗆 Pharm	acists		
🗆 None d	of the above		
*45.Are pat	tients provided education on important side effects of prescribed antibiotics?	_	_
45 -		□ Yes	∐ No
45a.	If 'Yes' is selected: How is education to patients on side effects shared? (Check all that a Discharge paperwork	<i>,</i> рріу.)	
	Verbally by nurse		
	Verbally by pharmacist		
	Verbally by physician		
	None of the above		
Facility Wate	r Management Program (WMP) (Completed with input from WMP team members)		
*52.Does y Legion	our facility have a water management program (WMP) to prevent the growth and transmis ella and other opportunistic waterborne pathogens (for example, <i>Pseudomoas, Acinetoba</i> Ideria, Stenotrophomonas, nontuberculous mycobacteria, and fungi)?		
Danito		□ Yes	🗆 No

		Exp	Form Approved 3 No. 0920-0666 0. Date: 12/31/26 ww.cdc.gov/nhsn
SAFE 528 TW9R Yes, who is represented on your	facility WMP team? (Check	all that apply):	
\Box Hospital Epidemiologist/Infection Prevent	tionist 🛛 🗆 Compliance/S	afety Officer	
\Box Hospital Administrator/Leadership	\Box Risk/Quality N	lanagement Staff	
□ Facilities Manager/Engineer	\Box Infectious Dise	ease Clinician	
\Box Maintenance Staff	□ Consultant		
\Box Equipment/Chemical Acquistion/Supplier	\Box Laboratory Sta	aff/Leadership	
Environmental Services	\Box Other (specifiy	/):	
*53.Has your facility ever conducted an environ opportunistic waterborne pathogens for ex piping infrastructure)? This may include a map all water supply sources, treatment sy	ample could grow and sprea description of building water	d in the facility water system (f systems using text or basic dia ntrol measures, and end-use p	or example, Igrams that
53a. If Yes, when was the most recent a	assessment conducted? (Ch	eck one)	
	tween 1 and 3 years ago ear and ≤3 years)	☐ More than 3 years ago (>3 years)	3
-	assessment conducted? (Ch tween 1 and 3 years ago ear and ≤3 years)	□ \ eck one) □ More than 3 years ago (>3 years)	res 🗆 No
Disinfectant (such as residual chlorine): 55a. If Yes, does your facility have a pla	an for corrective actions whe	□ Yes n disinfectant(s) are not within	□ No
limits as determined by the water man	agement program?	□ Yes	□ No
55b. If Yes, where and how frequently o			μ <i>ι</i> λ)

NHSN NATIONAL HEALTHCARE

Facility Water Management Program (WMP) (continued)

Location	Daily	Weekly	Monthl	Quarterly	Annually	Other (specify):	N/A
			у			<u></u>	
Entry Points							
Cold Potable Water Storage Tank(s)							
Hot Potable Water Storage Tank(s)							
Hot Water Supply							
Hot Water Return							
Representative Locations Throughout Cold Potable Building Water System(s)							
Representative Locations Throughout Hot Potable Building Water System(s)							
Other (specify):							

Water temperature:

🗆 No

□ Yes

□ Yes

55c.If Yes, does your facility have a plan for corrective actions when water temperatures are not within
acceptable limits as determined by the water management program?Image: YesImage: No

55d. If Yes, where and how frequently does your facility monitor water temperature? (check all that apply)

Location	Daily	Weekl	Monthly	Quarterly	Annuall	Other (specify):	N/A
		у			у	<u> </u>	
Entry Points							
Cold Potable Water Storage Tank(s)							
Hot Potable Water Storage Tank(s)							
Hot Water Supply							
Hot Water Return							
Representative Locations Throughout Cold Potable Building Water System(s)							
Representative Locations Throughout Hot Potable Building Water System(s)							
Other (specify):							

Water pH:□ Yes□ No55e.If Yes, does your facility have a plan for corrective actions when water pH is not within acceptable limits

as determined by the water management program?

55f. If Yes, where and how frequently does your facility monitor water pH? (check all that apply)

□ No

NHSN

Facility Water Management Program (WMP) (continued)

Location	Daily	Weekly	Monthl	Quarterly	Annually	Other (specify):	N/A
			у				
Entry Points							
Cold Potable Water Storage Tank(s)							
Hot Potable Water Storage Tank(s)							
Hot Water Supply							
Hot Water Return							
Representative Locations Throughout Cold Potable Building Water System(s)							
Representative Locations Throughout Hot Potable Building Water System(s)							
Other (specify):							

Heterotrophic plate count (HPC) testing:

🗆 Yes 🛛 🗆 No

55g. If Yes, does your facility have a plan for corrective actions when heterotrophic plate counts are not within acceptable limits as determined by the water management program?

55h. If Yes, where and how frequently does your facility perform HPC testing? (check all that apply)

Location	Daily	Weekly	Monthly	Quarterly	Annually	Other (specify):	N/A
Entry Points							
Cold Potable Water Storage Tank(s)							
Hot Potable Water Storage Tank(s)							
Hot Water Supply							
Hot Water Return							
Representative Locations Throughout Cold Potable Building Water System(s)							
Representative Locations Throughout Hot Potable Building Water System(s)							
Other (specify):							

 Specific environmental Legionella testing:

 \[
 \] Yes

 \[
 \] Yes

 \[
 \] No

 55i. If Yes, does your facility have a plan for corrective actions when environmental tests for Legionella are not
 within acceptable limits as determined by the water management program?

 \[
 Yes

 \[
 No

55j. If Yes, where an how frequently does your facility perform Legionella testing? (check all that apply)

NHSN

Facility Water Management Program (WMP) (continued)

Location	Daily	Weekly	Monthl	Quarterly	Annually	Other (specify):	N/A
			у				
Entry Points							
Cold Potable Water Storage Tank(s)							
Hot Potable Water Storage Tank(s)							
Hot Water Supply							
Hot Water Return							
Representative Locations Throughout Cold Potable Building Water System(s)							
Representative Locations Throughout Hot Potable Building Water System(s)							
Other (specify):							

Specific environmental Pseudomonas testing:

🗆 Yes 🛛 🗆 No

55k. If Yes, does your facility have a plan for corrective actions when environmental tests for *Pseudomonas* are not within acceptable limits as determined by the water management program?

□ Yes □ No

55I. If Yes, where an how frequently does your facility perform *Pseudomonas* testing? (check all that apply)

Location	Daily	Weekly	Monthly	Quarterly	Annually	Other (specify):	N/A
Entry Points							
Cold Potable Water Storage Tank(s)							
Hot Potable Water Storage Tank(s)							
Hot Water Supply							
Hot Water Return							
Representative Locations Throughout Cold Potable Building Water System(s)							
Representative Locations Throughout Hot Potable Building Water System(s)							
Other (specify):							

*56.Does your facility water management program address measures to prevent transmission of pathogens from wastewater premise plumbing to patients?

□ Yes

🗆 No

 \Box N/A, my facility does not have a water management program

NHSN NATIONAL HEALTHCARE VTE OUESTION

Justification: provide data (baseline and annually) on VTE prevention practices in hospitals/facilities and help identify gaps between evidence-based guidelines for VTE prevention and implementation of those guidelines in practice. The baseline data would also be helpful in the evaluation of future VTE prevention initiatives.

1. Our facility uses the following venous thromboembolism (VTE) prevention practices (select all that apply, and select at least one)

- 0 Our facility has a VTE prevention policy.
- 0 Our facility has a multidisciplinary team that addresses VTE prevention.
- 0 Our facility has a facility-wide VTE prevention protocol that includes VTE and bleeding risk assessments linked to clinical decision support for appropriate VTE prophylaxis options.

Our facility has embedded the VTE prevention protocol in admission order sets.

0 <mark>Yes No</mark>

- 0 Our facility provides VTE prevention education for clinicians annually.
- 0 Our facility provides VTE prevention education for patients during their stay at our facility.
- 0 Our facility performs audits to determine whether patients are on risk-appropriate VTE prophylaxis and provides clinician feedback for quality improvement.
- 0 Our facility tracks the incidence of VTE that develops during a patient's stay at our facility (VTE not present on admission).
- 0 Our facility does not use any of the above VTE prevention practices.

Validity Testing Questions

Justification: For the purposes of the Consensus Based Entity measure endorsement process, validity testing demonstrates the measure score (in our case, the SIR) correctly reflects the quality of care provided, adequately identifying differences in quality. The goal of these questions is to correlate process measures (for example, implementation of HAI prevention strategies) with the outcome measures of the NHSN SIRs.

Hypothesis: Facilities that implement an increased number of evidence-based HAI prevention measures between 2024 and 2025 will have an improvement in their SIR between the two years.

Alternative Hypothesis: Facilities that implement high number of evidence-based HAI prevention measures will have lower SIR compared to facilities that implement a lower number of prevention measures.

- 1. Our facility utilizes a checklist or bundle for prevention of the following HAIs. (Check all that apply)
 - CLABSI

At what minimum, regular frequency is adherence to the checklist/bundle monitored/measured? Check one.

- Weekly
- Monthly
- Quarterly
- Yearly
- PRN
- Other
- Not regularly monitored/measured

No

Is checklist/bundle adherence shared routinely with the clinical team?

Yes •

Unknown

CAUTI

At what minimum, regular frequency is adherence to the checklist/bundle monitored/measured? Check one.

Weekly



- Monthly
- Quarterly
- Yearly

•

- PRN
- Other
- Not regularly monitored/measured

Is checklist/bundle adherence shared routinely with the clinical team?

Yes
 No
 Unknown

CDI LabID Event

At what minimum, regular frequency is adherence to the checklist/bundle monitored/measured? Check one.

- Weekly
- Monthly
- Quarterly
- Yearly
- PRN
- Other

• Not regularly monitored/measured

Is checklist/bundle adherence shared routinely with the clinical team?

Yes
 No
 Unknown

 MRSA Bacteremia LabID Event At what minimum, regular frequency is adherence to the checklist/bundle monitored/measured? Check one.

- Weekly
- Monthly
- Quarterly
- Yearly
- PRN
- Other
- Not regularly monitored/measured

Is checklist/bundle adherence shared routinely with the clinical team?

Yes • No

Unknown

COLO SSI

•

At what minimum, regular frequency is adherence to the checklist/bundle monitored/measured? Check one.

- Weekly
- Monthly
- Quarterly
- Yearly
- PRN
- Other
- Not regularly monitored/measured

• No

Is checklist/bundle adherence shared routinely with the clinical team?

• Yes

Unknown

HYST SSI



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At what minimum, regular frequency is adherence to the checklist/bundle monitored/measured? Check one.

- Weekly
- Monthly
- Quarterly
- Yearly