

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 Tracking #: Facility ID: *Survey Year: **Facility Characteristics (completed by Infection Preventionist)** *Ownership (check one): □ Veterans Affairs ☐ For profit ☐ Not for profit, including church ☐ Government *Affiliation (check one): ☐ Hospital System ☐ Independent ☐ Multi-facility organization (specialty hospital network) ____ Free-standing ____ Within a hospital *Setting/classification: If classified as "Free-standing," does your LTAC hospital share physical housing with one or more of the following on-site facilities or units (check all that apply)? □ No ☐ Inpatient rehabilitation facility ☐ Skilled nursing facility (SNF)/nursing home □ Neuro-behavioral unit or facility □ Other (specify): _____ ☐ Residential facility (assisted living If classified as "Within a hospital," is your LTAC hospital located: In a building that does not provide acute care services (for example, psychiatric hospital?)

□ Yes □ No Near (but not within) an acute 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 following categories (categories should equal total): a. Intensive care unit (CIU) or critical care beds: b. High observation/special care/high acuity beds (not ICU): c. General LTAC beds: *Total number of LTAC beds (licensed capacity): Assurance of Confidentiality: The voluntarily provided information obtained in this surveillance system that would permit identification of any individual or 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)). CDC 57.150 (Front). Rev 10, v13.0

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2 = bioMérieux/Vitek

3 = BD Phoenix

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				www.c	dc.gov/nh
Facility Characteristics (continu	ued)				
*Number of single occupancy room *Number of double occupancy room *Number of triple occupancy rooms *Number of quadruple occupancy i	ms: s:				
*Total number of admissions with on not developing during LTAC stay):	_		•	present of a	noissimt
If helpful for your facility in identifying associated with these conditions for a. Ventilator dependence:b. Hemodialysis:	ound here: http://www.cdc.gov/nhsr				
Facility Microbiology Laborator	y Practices (completed with inpe	ut from Mic	crobiology Laborat	tory Lead)	
*1. Does your facility have its susceptibility testing?	own on-site laboratory that perform	s bacterial	antimicrobial	□ Yes	□ No
1a. If No, where is your fac	cility's antimicrobial susceptibility te	esting perfor	rmed: (check one)		
☐ Affiliated medical cente	er □ Commercial referral lab	oratory	☐ Other local/region reference laborate		liated
1b. If Yes, do you also ser	nd out any antimicrobial susceptibil	ity testing (d	check one)	□ Yes	□ No
methods are used for: (1) Primary susceptibility t	udomonas aeruginosa and/or Acino esting and ntal, or confirmatory testing (if perfo		<i>aumannii</i> complex,	indicate whic	;h
If your laboratory does not Use the testing codes listed	perform susceptibility testing, indicated below the table.	ate the me	thods used at the o	utside labora	tory.
(1) Primary	(2) Secondary	Comm	ents		
1 = Kirby-Bauer disk diffusion	4 = ThermoFiscer/Sensititre	7 = Gra	adient Dilution Strip	(for example	E test)

5 = Beckman Coulter/MicroScan

6 = Selux Diagnostics

8 = Sent out test, method not known

9 = Other (describe in Comments section)



Facility Microbiology Laboratory Practices (continued)

*3. Does either the primary or secondary/supplemental antimicrobial susceptibility testing (AST) include the following (check all that apply):

	Drug	Tested	Not Tested		
	Cefiderocol				
	Ceftazidime-Avibactam				
	Ceftolozane-Tazobactam				
	Eravacycline				
	Plazomicin				
	Imipenem-Relebactam				
	Meropenem-Vaborbactam				
	Aztreonam-Avibactam				
	Sulbactam-Durlobactam				
Ha: a. b. c. d. e. f. g. h.	Third Generation Cephalosporin a Enterobacterales in 2010 Carbapenem breakpoints for Enterocarbapenem breakpoints for Enterocarbapenem breakpoints for Pseu Fluroquinolone breakpoints for Enterocarbapenem breakpoints for Enterocarbapenem breakpoints for Pseu Fluroquinolone breakpoints for Enterocarbapenem breakpoints for Enteroca	erobacterales <u>in</u> 2010 bbacterales <u>in</u> 2012 udomonas aeruginosa <u>in</u> 2012 eudomonas aeruginosa <u>in</u> 2012 eterobacterales <u>in</u> 2019 interobacterales in 2023	eonam) breakpoints for	 Yes 	No No No No No No No No
j.	Piperacillin-tazobactam breakpoi	nts for <i>Enterobacterales</i> in 20	22	□ Yes	□ No
not	es the laboratory test bacterial isola include automated testing instrum. If Yes, indicate what is done if car. Change susceptible carbaper. Report carbapenem MIC resu. No changes are made in the infection control practices.	ent expert rules) bapenemase production is de lem results to resistant llts without an interpretation	tected: (check one)	□ Yes	



	5b. If \	es, which test is routinely perfor	med to detect carbapenem	ase: (check all that ap	oply)	
		☐ Nucleic Acid Amplification Test (PCR, Cepheid, etc.)	□ mCIM/CIM	☐ NG-Test Carba-5 lateral flow assay)	(or other	
		☐ Modified Hodge Test	□ Carba NP	□ Other		
	5c. If \	es, which of the following are ro	outinely tested for the prese	nce of carbapenemas	es: (check all that apply)	
		Enterobacterales spp.	□ Pseudomonas aerugin	osa 🗆 Aci	inetobacter baumannii	
*6.	resista	rour facility use commercial or la nce markers in bacterial bloodst e FilmArray, Luminex Verigene, e	ream infections? Examples	•		
		Yes				
		No [if checked, skip questions 7	⁷ and 8]			
	6a. If \	es, which test panel(s) does you	ur facility use? (check all tha	at apply)		
*7.	In a sc testing	Accelerate PhenoTest BC Cepheid Xpert MRSA/SA BC GenMark ePlex BCID-FP MALDI-TOF MS directly from p MALDI-TOF MS based antimic T2Biosystems T2Bacteria Other Commercial Test(s) (Lea Other Laboratory Developed To	☐ GenMark ePlex BCID ☐ Luminex Verigene BC ☐ cositive blood culture (e.g., strobial resistance detection ☐ T2Biosystems T2Candave Comment) ☐ est(s) (Leave Comment) ☐ ce marker and Staphylococc procedure(s) your facility co	GP GenMark GP Luminex SepsiTyper) dida T2Biosys cus aureus are detect anducts. (check one)	ed by rapid molecular	
	7a □ 7a	- Culture based phenotypic antimi	crobial susceptibility testing	is not performed. [If o	checked, skip question	
	CO	Culture based phenotypic antimi rresponding rapid molecular test phenotypic test result.		•	•	
	rap	Culture based phenotypic antimi oid molecular testing and/or inter	pretation is added.	•		
	blo	7a. If both rapid molecular and culture based phenotypic antimicrobial susceptibility testing are performed for a blood specimen to detect drug resistance in <i>Staphylococcus aureus</i> , and discordance is found between their results, how are results reported? (check one)				
		Further testing is not pursued.	Results are reported separa	itely.		
		Further testing is not pursued. an antimicrobial resistance ma		erridden by the rapid r	molecular test result when	
		Further testing is performed to further analysis.	identify the reason for the o	iscordance. Results a	are modified based on the	



*8.	In a scenario where the <i>bla_{CTX-M}</i> (CTX-M) resistance marker and <i>Escherichia coli</i> are detected by rapid molecular testing in a blood specimen, select the procedure(s) your facility conducts. (check one)						
	 ☐ Our laboratory does not perform bla_{CTX-M} (CTX questions 8a] 	-M) testing using rapid molecular methods. [If checked, skip					
	☐ Culture based phenotypic antimicrobial suscep 8a.]	tibility testing is not performed. [If checked, skip question					
	• • • • • • • • • • • • • • • • • • • •	tibility testing is performed. A text indicating results of the interpretation of the rapid molecular testing result is added to					
	• • • • • • • • • • • • • • • • • • • •	□ Culture based phenotypic antimicrobial susceptibility testing is performed. No text indicating corresponding rapid molecular testing and/or interpretation is added.					
	8a. If both rapid and culture based phenotypic antimic specimen to detect drug resistance in <i>Escherichia</i> are results reported? (check one)	crobial susceptibility testing are performed for a blood a coli and discordance is found between their results, how					
	☐ Further testing is not pursued. Results are reported separately.						
	 Further testing is not pursued. The phenotypic result is overridden by the rapid molecular test an antimicrobial resistance marker is detected. 						
	 Further testing is performed to identify the real further analysis. 	ason for the discordance. Results are modified based on the					
*9.	*9. Where is yeast identification performed for specimens	s collected at your facility? (check one)					
	□ On-site laboratory						
	☐ Affiliated medical center						
	☐ Commercial referral laboratory						
	☐ Other local/regional, non-affiliated reference labora	tory					
	☐ Yeast identification not available (specifically, yeas affiliate/commercial/other laboratory) [If checked, skip	,					
	Answer questions 10-14 for the laboratory that p*10.Which of the following methods are used for yeast ide						
	☐ MALDI-TOF MS System (Vitek MS) ☐ Mi	croScan					
	☐ MALDI-TOF MS System (Bruker Biotyper) ☐ No	on-automated Manual Kit (for example, API 20C, RapID, Tube, PNA-FISH, etc.)					
		NA sequencing					
	☐ BD Phoenix ☐ Of	her (specify):					



□ Yes	□ No	□ Unkr	nown	
*12. <i>Candid</i> that ap		e following body sites	are usually fully ide	ntified to the species level? (check all
□ Bloc	od		□ Respiratory	
□ Oth	er normally sterile body site	e (for example, CSF)	☐ Other (specify)	:
□ Urir	ne		☐ None are fully	identified to the species level
*13.Does t	he laboratory employ any F	PCR molecular tests to	identify Candida from	om blood specimens?
□ Yes	□ No	□ Unkr	nown	
13a. ap	If yes, which PCR molecuply)	ular tests are used to id	lentify <i>Candida</i> fron	n blood specimens? (check all that
	T2Candida Panel			
	BioFire BCID			
	GenMark ePlex BCID Other, specify:			
	Unknown	 		
13b.	If yes and you get a positi	ive result, does this lab	culture the blood t	o obtain an isolate?
	Yes, always			
	Yes, with clinical order			
	No			
	Unknown			
*14.Where	is antifungal susceptibility	testing (AFST) perform	ned for specimens	collected at your facility? (check one)
□ On-site	laboratory	☐ Other local/region	nal, non-affiliated re	eference laboratory
☐ Affiliate	d medical center			ST is not performed onsite or at any
□ Comme	rcial reference laboratory	affiliate/commercial/	other laboratory) [if	selected, skip questions 15 -19]
swer ques	stions 15-19 for the lab	oratory that <i>perforn</i>	ns AFST for you	<u>r facility</u> :
*15.What n apply)	nethods are used for antifu	ngal susceptibility testi	ng (AFST), exclud	ling Amphotericin B? (check all that
	th microdilution with tory developed plates	☐ YeastOne (Ther Sensititre™)	mo Scientific™	☐ Gradient diffusion (E test)
	k (bioMerieux)	- OII / :()		☐ Unknown



nods are used for ar	ntifungal susceptibility testir	ng (AFST) of Ampho	<i>tericin B</i> ? (check al	l that apply)	
☐ Broth microdilution with laboratory developed plates		mo Scientific™	☐ Gradient diffusion	on (E test)	
oioMerieux)	☐ Other (specify): _		☐ Unknown		
erformed for which o	of the following antifungal d	rugs? (check all that	apply)		
azole	□ Voriconazol	e	☐ Itraconazole		
onazole	☐ Micafungin		☐ Anidulafungin		
ungin	☐ Amphoteric	in B	☐ Flucytosine		
specify:	□ Unknown				
erformed on fungal	isolates in which of the follo	owing situations? (che	eck all that apply)		
Performed automatically Performed with a Not performed Unkno					
mally sterile body kample, CSF)					
•					
ecity):					
oratory developing a his laboratory?	ntibiograms or other report	s to track susceptibili	ty trends for <i>Candid</i>	a spp. isolates	
□ No	□ Unkn	own			
*20.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 Cell cytotoxicity neutralization assay Nucleic acid amplification test (NAAT) (for example, PCR, LAMP) 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 (<i>C. difficile</i> culture followed by detection of toxins) Other (specify):					
	mally sterile body cample, CSF) ry ecify): pratory developing a his laboratory? c primary testing me where your facility's nzyme immunoassa ell cytotoxicity neutracicleic acid amplification AAT plus EIA, if NAM lutamate dehydroge DH plus NAAT (2-st DH plus EIA for toxicoxigenic culture (C. exponents)	nicrodilution with developed plates Sensititre™) pioMerieux) □ Other (specify): _ performed for which of the following antifungal of azole □ Voriconazole □ Micafungin □ Amphoterici specify: □ Unknown performed on fungal isolates in which of the following automatically mally sterile body cample, CSF) Ty □ □ pecify): □ □ pratory developing antibiograms or other report his laboratory? □ No □ Unknown performed for C. difficile used meaning is performed? (che nazyme immunoassay (EIA) for toxin ell cytotoxicity neutralization assay uscleic acid amplification test (NAAT) (for example, AAT plus EIA, if NAAT positive (2-step algorithm) DH plus EIA for toxin, followed by NAAT for disposigenic culture (C. difficile culture followed by NAAT for disposigenic culture (C. difficile culture followed by NAAT for disposigenic culture (C. difficile culture followed by NAAT for disposigenic culture (C. difficile culture followed by NAAT for disposigenic culture (C. difficile culture followed by NAAT for disposigenic culture (C. difficile culture followed by NAAT for disposigenic culture (C. difficile culture followed by NAAT for disposigenic culture (C. difficile culture followed by NAAT for disposition culture (C. difficile culture followed by NAAT for disposition culture (C. difficile culture followed by NAAT for disposition culture (C. difficile culture followed by NAAT for disposition culture (C. difficile culture followed by NAAT for disposition culture (C. difficile culture followed by NAAT for disposition culture (C. difficile culture followed by NAAT for disposition culture (C. difficile culture followed by NAAT for disposition culture (C. difficile culture followed by NAAT for disposition culture (C. difficile culture followed by NAAT for disposition culture (C. difficile culture followed by NAAT for disposition culture (C. difficile culture followed by NAAT for disposition culture (C. difficile culture followed by NAAT for disposition culture (C. difficile culture followed by NAAT for disposition culture (C. dif	nicrodilution with developed plates Sensititre™) pioMerieux) □ Other (specify):	developed plates Sensititre™) Other (specify): Unknown	



*21.Which (check	of the following methods serve as the primary method used for bacterial identification at your facility?
	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
facility?	of the following methods serve as the secondary or backup method used for bacterial identification at your (for example, a secondary method if the primary method fails to give an identification, or if the primary I 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
nfection Con Coordinator)	trol Practices (completed with input from Hospital Epidemiologist and/or Quality Improvement
*23.Numbe	r or faction of infection preventionists (IPs) in facility:
	otal hours per week performing surveillance:
b. T	otal hours per week for infection control activities other than surveillance:
	r or fraction of full-time employees (FTEs) for a designated hospital epidemiologist (or equivalent role) d with your facility:
•	olicy in your facility that patients infected or colonized with MRSA are routinely placed in contact tions while these patients are in your facility? (check one)
□ Yes	□ No □ Not applicable: my facility never admits these patients
25a. (ch	If Yes, check the type of patients that are routinely placed in contact precautions while in your facility eck one):
	All infected and all colonized patients
	Only all infected patients
	Only infected or colonized patients with certain characteristics (check all that apply)



Infection Control Practices (continued)

		□ Patients admitted to high risk	ngs	
		☐ Patients at high risk for trans	on	
*26	•	olicy in your facility that patients infe ese patients are in your facility? (ch		re routinely placed in contact precautions
	□ Yes	□ No	□ Not applicable: my fa	cility never admits these patients
	26a. (ch	If Yes, check the type of patients theck one):	e routinely place in conta	act precautions while in your facility
		All infected and all colonized patier		
		Only all infected patients		
		Only infected or colonized patients	certain characteristics (c	heck all that apply)
		$\ \square$ Patients admitted to high risk	ngs	
		☐ Patients at high risk for trans	on	
*27		enemase production) are routinely p	,	regardless of confirmatory testing for while these patients are in your facility?
	□ Yes	□ No	□ Not applicable: my fa	cility never admits these patients
	27a. (ch	If Yes, check the type of patients theck one):	re routinely placed in con	tact precautions while in your facility
		All infected and all colonized patier		
		Only all infected patients		
		Only infected or colonized patients	certain characteristics (c	heck all that apply)
		$\ \square$ Patients admitted to high risk	ngs	
		☐ Patients at high risk for trans	on	
*28	extende		•	cted or confirmed ESBL-producing or ly placed in contact precautions while
	□ Yes	□ No	□ Not applicable: my fa	cility never admits these patients
	28a. (ch	If Yes, check the type of patients theck one):	e routinely placed in con	tact precautions while in your facility
		All infected and all colonized patier		
		Only all infected patients		
		Only infected or colonized patients	certain characteristics (c	heck all that apply)
		□ Patients admitted to high risk	ngs	
		☐ Patients at high risk for trans	on	



Infection Control Practices (continued)

	the facility routinely perform screening testing (culture or non-culture) for CRE? This includes screening for 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
а	oply)
	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
	□ 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. fa	If Yes, what method is routinely used by the lab conducting CRE testing of screening swabs from your icility? (check all that apply)
	Culture-based methods
	the facility routinely perform screening testing (culture or non-culture) for <i>Candida auris</i> ? This includes ning for patients at your facility performed by public health laboratories and commercial laboratories. □ Yes □ No If Yes, in which situations does the facility routinely perform screening testing for <i>Candida auris</i> ? (check
а	ll that apply)
	Surveillance testing at admission for all patients
	Surveillance testing of epidemiologically-linked patients of newly identified <i>Candida auris</i> 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
	□ Patients admitted to high-risk settings (for example, ICU)
	□ Other high-risk patients (specify):
С	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)



Infection Control Practices (continued)

30b. fro	If Yes, what method is routinely u m your facility?	sed by the lab co	nducting <i>Candida auris</i> tes	ting of screening swabs
	Culture-based methods	□ PCR	□ Other (specify): _	
*31.Does tl	he facility routinely perform screeni	ng testing (cultur	e or non-culture for MRSA	for any patients admitted? □ Yes □ No
31a. ap	If Yes, in which situations does th ply)	e facility routinely	perform screening testing	for MRSA? (check all that
	Surveillance testing at admission	for all patients		
	Surveillance testing at admission [LTAC] or long-term care facility [l	of high-risk patie		from long-term acute care
	Surveillance testing at admission	of patients admit	ted to high-risk settings (for	r example, ICU)
	Surveillance testing of pre-operat	ive patients to pre	event surgical site infection	s
	Other (specify):			
•	our facility have a policy to routinel ission of MDROs at your facility?	y use chlorhexidi	ne bathing for any adult pa	tients to prevent infection or
				□ Yes □ No
32a.	If Yes, indicate which patients: (se	elect all that apply	y)	
□ IC	U patients:	□ Patients outs	ide the ICU:	□ Pre-operatively for
0	All ICU patients	○ All patient	s outside the ICU	patients undergoing
0	Subset of ICU patients:	 Subset of 	patients outside the ICU:	surgery
	☐ Patients with central venous catheter or midline catheters	☐ Patients with central venous catheter or midline catheters		
	□ Other, specify:	□ Other, s	specify:	
antista	he facility have a policy to routinely phylococcal agent (mupirocin, iodo t healthcare-associated infections o	phor, or an alcoh	ol based intranasal agent)	for any adult patients to
Antibiotic St	ewardship Practices (completed	with input from	Physician and Pharmacis	st Stewardship Leaders)
☐ Providir interventic ☐ Allocati efforts. ☐ Having resources ☐ Present ☐ Ensurin board at le	r leadership has demonstrated coming stewardship program leader(s) cons. Ing resources (for example, IT suppose a senior executive that serves as a land support to accomplish its missing information on stewardship acting the stewardship program has an east annually. Unicating to staff about stewardship	dedicated time to port, training for so point of contact sion. ivities and outcor opportunity to dis	manage the program and of tewardship team) to support or "champion" to help ensu- mes to facility leadership ar scuss resource needs with	conduct daily stewardship ort antibiotic stewardship ore the program has ond/or board at least annually. facility leadership and/or



□ Providi	ng opportunities for	or hospital staff training and dev	velopment on antibiotic stewardship.
	ng a formal staten by the board).	nent of support for antibiotic ste	ewardship (for example, a written policy or statement
contributi	ng to stewardship		roups (for example, IT and hospital medicine) are
□ None c	of the above.		
*35.Our fa	cility has a leader	or co-leaders responsible for ar	ntibiotic stewardship program management and outcomes.
35a.	If Vee what is th	ne position of this leader? (chec	□ Yes □ No
	Physician	□ Co-led by both Pharn	•
	Pharmacist	•	RN, PA, NP, etc.; specify):
35b. lea	If Physician or C ader? (check all th		following describes your antibiotic stewardship physician
	Has antibiotic ste	ewardship responsibilities in the	eir contract or job description or performance review
	Is physically on-	site in your facility (either part-ti	ime or full-time)
	Completed an ID) fellowship	
	Completed a cer	tificate program on antibiotic st	tewardship
	Completed other	rtraining(s) (for example, confe	erences or online modules) on antibiotic stewardship
	None of the above	ve.	
•	o) leader): What pe		their contract or job description' is selected (for physician stewardship activities is specified in the physician (co)
	□ 1-10%	□ 11-25%	□ 26-50%
	□ 51-75%	□ 76-100%	□ Not specified
35d. lea	•	o-led is selected: In an averag ibiotic stewardship activities in	ge week , what percentage of time does the physician (co) your facility? (check one)
	□ 1-10%	□ 11-25%	□ 26-50%
	□ 51-75%	□ 76-100%	
35e. p h		Co-led is selected, which of the (check all that apply)	e following describes your antibiotic stewardship
	Has antibiotic ste	ewardship responsibilities in the	eir contract, job description or performance review
	Is physically on-	site in your facility (either part-ti	ime or full-time)
	Completed a PG	GY2 ID residency and/or ID fello	pwship
	Completed a cer	tificate program on antibiotic st	tewardship
	·	. •	erences or online modules) on antibiotic stewardship
	None of the above	• , , , , ,	,



	(co) leader): What perce	· ·	r contractor or job description stewardship activities is spec	` .
		□ 1-10%	□ 11-25%	□ 26-50%	
		□ 51-75%	□ 76-100%		
	35g. (co			erage week, what percentage es in your facility? (check one	e of time does the pharmacist
		□ 1-10%	□ 11-25%	□ 26-50%	
		□ 51-75%	□ 76-100%		
	35h. poi		ner is selected: Does your fa port for the non-physician I	acility have a designated physeader?	sician who can serve as a
					□ Yes □ No
		pharmacist is not the proving antibiotic use		e program, is there at least or	ne pharmacist responsible for
		J	•		□ Yes □ No
36.	Our fac	cility has the following	priority antibiotic stewards	hip interventions: (Check all t	
	36a. aud	If Prospective audit		Our antibiotic stewardship pro	of interventions, acceptance of
					□ Yes □ No
	□ Prea	uthorization for speci	fic antibiotic agents.		
	36b. (fo		s selected: Our antibiotic st g which agents are request		preauthorization interventions
					□ Yes □ No
	to assis	st with antibiotic selec		on national guidelines and loc onditions (for example, comm	cal pathogens susceptibilities, nunity-acquired pneumonia,
	36c.	If Facility-specific tre	eatment recommendations	is selected: For which commo	on clinical conditions?
		Community-acquire	d pneumonia		
		Urinary tract infection	on		
		Skin and soft tissue			
		None of the above			



to our	Facility-specific treatment recommendations is selected: Our stewardship program moni facility's treatment recommendations for antibiotic selection for common clinical conditionle, community-acquired pneumonia, urinary tract infection, skin and soft infections).		nerence
		□ Yes	□ No
36e. If	Yes: For which common clinical conditions?		
□ Co	ommunity-acquired pneumonia		
□ Ur	rinary tract infection		
	kin and soft tissue infection		
□ No	one of the above		
*37.Our facility that apply.	γ has a policy or formal procedure for other interventions to ensure optimal use of antibion.)	tics: (Cl	heck all
□ Early adm	ninistration of effective antibiotics to optimize the treatment of sepsis		
□ Treatment	t protocols for Staphylococcus aureus bloodstream infection		
□ Stopping ι	unnecessary antibiotic(s) in new cases of <i>Clostridioides difficile</i> infection (CDI)		
□ Review of	culture-proven invasive (for example, bloodstream) infections		
□ Review of	planned outpatient parenteral antibiotic therapy (OPAT)		
☐ The treating	ng team to review antibiotics 48-72 hours after initial order (specifically, antibiotic time-or	ut)	
□ Assess ar	nd clarify documented penicillin allergy		
•	shortest effective duration of antibiotics at discharge for common clinical conditions (for acquired pneumonia, urinary tract infection, skin and soft tissue infections) ne above	exampl	e,
selecto at disc	'Using the shortest effective duration of antibiotics at discharge for common clinical conded: Our stewardship program monitors adherence in using the shortest effective duration charge for common clinical conditions (for example, community-acquired pneumonia, uridons, skin and soft tissue infections), at least annually.	n of anti nary tra	biotics ct
*20 O fo cilit.		□ Yes	□ No
-	n has in place the following specific 'pharmacy-based' interventions: (Check all that apply n-driven changes from intravenous to oral antibiotics without a physician's order (for exar notocol)	-	ospital-
□ Alerts to p anaerobes)	providers about potentially duplicative antibiotic spectra (for example, multiple antibiotics	to treat	
□ Automatic	antibiotic stop orders in specific situations (for example, surgical prophylaxis)		
□ None of th	ne above		
*39.Our stewa	rdship 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' intervention tapply.)	ons: (Ch	eck all
Па	Nurses receive training on appropriate criteria for sending urine and/or respiratory culture	ic.	
	Nurses initiate discussions with the treating team on switching from intravenous to oral ar		
	Nurses initiate antibiotic time-out discussions with the treating team.	itibiotios	•
	Nurses track antibiotic duration of therapy.		
	None of the above.		
*40 04-	overdelein nach neuen neuen items (Oberell ell theter annh.)		
	ewardship program monitors: (Check all that apply.)		
	tic resistance patterns (either facility- or region-specific), at least annually		
	dioides difficile infections (or C. difficile LabID events), at least annually		
	tic use in days of therapy (DOT) per 1000 patient days or days present, at least quarterly		
	tic use in defined daily doses (DDD) per 1000 patient days, at least quarterly		
	tic expenditures (specifically, purchasing costs), at least quarterly		
	tic use in some other way, at least annually (specify):		
□ None o	of the above		
*41.Our ste	ewardship team provides the following antibiotic use reports to prescribers, at least annually	y: (Chec	k all that
□ Individ	ual, prescriber-level reports		
□ Unit- o	r service-specific reports		
□ None o	of the above		
	If 'Individual, prescriber-level reports' or 'Unit-or service-specific reports' is selected: Our segram uses these reports to target feedback to prescribers about how they can improve the escribing, at least annually.		
		□ Yes	□ No
*42.Our fac	cility distributes an antibiogram to prescribers, at least annually.		
		□ Yes	□ No
*43.Informa	ation on antibiotic use, antibiotic resistance, and stewardship efforts is reported to hospital sy.	staff, at l	east
		□ Yes	□ No
	of the following groups receive education on optimal prescribing, adverse reactions from a cic resistance (for example, Grand Rounds, in-service training, direct instruction) at least an apply.)		
□ Prescri	ibers		
□ Nursin	g staff		
□ Pharm	acists		
□ None o	of the above		



*45.Are pa	tients provided education on	important side eff	ects of prescribed	antibiotics?			
					□ Yes □ No		
45a.	If 'Yes' is selected: How is	education to patie	nts on side effects	shared? (Check all that a	apply.)		
	Discharge paperwork	·		·			
	Verbally by nurse						
	Verbally by pharmacist						
	Verbally by physician						
	None of the above						
Facility Wate	r Management Program (V	VMP) (Completed	d with input from	WMP team members)			
Legion	our facility have a water man ella and other opportunistic volderia, Stenotrophomonas, r	waterborne patho	gens (for example,	Pseudomonas, Acinetob			
					□ Yes □ No		
46a.	If Yes, who is represented	on your facility WI	MP team? (Check	all that apply):			
□ Hosp	ital Epidemiologist/Infection	Preventionist	□ Compliance/Sa	afety Officer			
□ Hosp	ital Administrator/Leadership)	□ Risk/Quality M	lanagement Staff			
□ Facili	ties Manager/Engineer		□ Infectious Disease Clinician				
□ Main	tenance Staff		□ Consultant				
□ Equip	oment/Chemical Acquisition/	Supplier	□ Laboratory Staff/Leadership				
□ Envir	onmental Services		□ Other (specify)):			
opport piping	our facility ever conducted an unistic waterborne pathogen infrastructure)? This may inc I water supply sources, treat	s for example cou lude a description	ld grow and spread of building water s	d in the facility water syst systems using text or bas	em (for example, sic diagrams that		
47a.	If Yes, when was the most	recent assessmer	nt conducted? (Ch	eck one)			
□W	ithin the most recent year	□ Between 1 an	d 3 years ago	☐ More than 3 years ag	o (>3		
(<1 <u>y</u>	/ear ago)	(<u>></u> 1 year and <u><</u> 3	years)	years)			
source examp 48a. □ W	our facility has ever conducte s, modes of transmission, pa le WICRA tool can be asses If Yes, when was the most ithin the most recent year	atient susceptibility sed at <u>https://www</u> recent assessmer □ Between 1 an	y, patient exposure y.cdc.gov/hai/pdfs/ nt conducted? (Cho d 3 years ago	e, and/or program prepare prevent/water-assessment eck one) More than 3 years ag	edness? An nt-tool-508.pdf. □ Yes □ No		
(<1)	/ear ago)	(<u>></u> 1 year and <u><</u> 3	years)	years)			



Facility Water Management Program (WMP) (continued)

*49.Does your facility regularly monitor the following parameters in the building water system(s)?							
Disinfectant (such as residual chlorine): □ Yes						□ No	
49a. If Yes, does your facility have a plan for corrective actions when disinfectant(s) are not within acceptable limits as determined by the water management program? □ Yes □ No					acceptable		
-		_			disinfectant	s)? (Check all that app	
	•	•	-	•			3,
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):							
Other (specify): Water temperature:						□ Yes	□ No
Other (specify): Water temperature: 49c. If Yes, does your fa	icility hav	/e a plan f	or correctiv	/e actions w	hen water t	□ Yes emperatures are not w	□ No ithin
Other (specify): Water temperature: 49c. If Yes, does your fa acceptable limits as det	cility hav	/e a plan f	or correctiv	/e actions w ement prog	rhen water t	□ Yes	□ No ithin □ No
Other (specify): Water temperature: 49c. If Yes, does your fa acceptable limits as det	cility hav	/e a plan f	or correctiv ter manag s your facil	/e actions w ement prog	rhen water t	□ Yes emperatures are not w □ Yes	□ No ithin □ No
Other (specify): Water temperature: 49c. If Yes, does your fa acceptable limits as det 49d. If Yes, where and h	cility have	ve a plan for the ware the depth of the ware the depth of the ware	or correctiv ter manag s your facil	e actions w ement prog ity monitor v	hen water t ram? vater tempe	□ Yes emperatures are not w □ Yes erature? (check all that	□ No ithin □ No apply)
Other (specify): Water temperature: 49c. If Yes, does your fa acceptable limits as det 49d. If Yes, where and h	cility have remined low freques Daily	ve a plan for the ware uently does weekly	or correctiv ter manag s your facil Monthly	e actions we ment progity monitor very Quarterly	hen water t ram? vater tempe Annually	☐ Yes emperatures are not w ☐ Yes erature? (check all that Other (specify):	□ No ithin □ No apply) N/A
Other (specify): Water temperature: 49c. If Yes, does your far acceptable limits as det 49d. If Yes, where and he Location Entry Points Cold Potable Water Storage	ecility have cermined bow freques Daily	ve a plan for the ware depth of the ware depth o	or corrective ter manages your faciled Monthly	/e actions w ement prog ity monitor v Quarterly	rhen water t ram? water tempe Annually	☐ Yes emperatures are not w ☐ Yes erature? (check all that ☐ Other (specify): ☐ ☐	□ No ithin □ No apply) N/A
Other (specify): Water temperature: 49c. If Yes, does your fa acceptable limits as det 49d. If Yes, where and h Location Entry Points Cold Potable Water Storage Tank(s) Hot Potable Water Storage Tank(s) Hot Water Supply	cility have cermined sow frequency Daily	ve a plan for the warently does	or correctivater manages your facil	ve actions we ement progity monitor very Quarterly	hen water tram? water tempe Annually	☐ Yes emperatures are not w ☐ Yes erature? (check all that Other (specify): ☐ ☐	□ No ithin □ No apply) N/A
Other (specify): Water temperature: 49c. If Yes, does your fa acceptable limits as def 49d. If Yes, where and h Location Entry Points Cold Potable Water Storage Tank(s) Hot Potable Water Storage Tank(s)	cility have cermined sow frequency Daily	ve a plan for the water that the wat	or correctivater manages your facil	ve actions we ement progity monitor very Quarterly	hen water tram? water tempe Annually	□ Yes emperatures are not w □ Yes erature? (check all that Other (specify): □ □ □	□ No ithin □ No apply) N/A □
Other (specify): Water temperature: 49c. If Yes, does your far acceptable limits as deft 49d. If Yes, where and he Location 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)	cility have cermined bow frequestions and the control of the contr	ve a plan for by the wateritly does weekly	or corrective ter manages your faciled Monthly	/e actions we ement progrity monitor we Quarterly	rhen water tram? water temperature Annually	□ Yes emperatures are not w □ Yes erature? (check all that Other (specify): □ □ □	□ No ithin □ No apply) N/A □ □
Other (specify): Water temperature: 49c. If Yes, does your far acceptable limits as def 49d. If Yes, where and he Location 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	cility have	ve a plan for by the water	or correctivater manages your facil	ve actions we ement progrity monitor we Quarterly	rhen water tram? vater tempe Annually	□ Yes emperatures are not w □ Yes erature? (check all that Other (specify): □ □ □ □ □	□ No ithin □ No apply) N/A □ □ □ □



Facility Water Management Prog	gram (W	MP) (con	tinued)				
Water pH:						□ Yes	□ No
49e. If Yes, does your fa	cility hav	e a plan fo	or correctiv	e actions w	hen water p	oH is not within accepta	able limits
as determined by the w		•				□ Yes	□ No
49f. If Yes, where and how f	requently	y does you	ur facility m	nonitor wate	r pH? (chec	k 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):							
Heterotrophic plate count (HPC) testing: 49g. 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? Yes No 49h. 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)							
_							
Tank(s)							
Tank(s) Hot Water Supply Hot Water Return Representative Locations Throughout Cold Potable Building Water System(s)							
Tank(s) Hot Water Supply Hot Water Return Representative Locations Throughout Cold Potable							



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						<u>ww</u>	w.cdc.gov/nhs
Facility Water Management Prog	gram (W	MP) (cont	tinued)				
Specific environmental Leg	ionella te	esting:				□ Yes	□ No
49i. If Yes, does your facility have a plan for corrective actions when environmental tests for Legionella are not							
within acceptable limits	as deter	mined by	the water i	managemen	it program?	□ Yes	□ No
49j. If Yes, where an how fr	equently	does you	r facility pe	rform <i>Legio</i>	<i>nella</i> testing	g? (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):						П	
	•	1	•				•
Specific environmental Pse		•				□ Yes	□ No
_	-	•				nmental tests for <i>Pseu</i>	domonas
are not within acceptab	ie iimits a	as determi	inea by the	water man	agement pr	_	- N.
401 If Van Juhana an haw fr		da a a	r fa ailite e na	wfawaa Daa	daaa.	☐ Yes	□ No
49l. If Yes, where an how fr	equently	does you	г тасшіу ре	riorm Pseud	ornonas te	sting? (check all that a	ірріу)
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):							



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www.cdc.gov/nhsn Facility Water Management Program (WMP) (continued) *50.Does your facility water management program address measures to prevent transmission of pathogens from wastewater premise plumbing to patients? ☐ Yes □ No □ N/A, my facility does not have a water management program Venous Thromboembolism (VTE) Practices *51. Our facility uses the following venous thromboembolism (VTE) prevention practices (select all that apply, and select at least one) □ Our facility has a VTE prevention policy. Our facility has a multidisciplinary team that addresses VTE prevention. 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. □ Yes □ No ☐ Our facility provides VTE prevention education for clinicians annually. Our facility provides VTE prevention education for patients during their stay at our facility. Our facility performs audits to determine whether patients are on risk-appropriate VTE prophylaxis and provides clinician feedback for quality improvement. Our facility tracks the incidence of VTE that develops during a patient's stay at our facility (VTE not present on admission). □ Our facility does not use any of the above VTE prevention practices. **Prevention Practices** *52. 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 Is checklist/bundle adherence shared routinely with the clinical team? □ Yes No □ Unknown 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
ш	
	At what minimum, regular frequency is adherence to the checklist/bundle monitored/measured? Check one.
	□ Weekly
	□ Monthly
	□ Quarterly
	□ Yearly
	□ PRN Î
	□ Other
	□ Not regularly monitored/measured
	Is checklist/bundle adherence shared routinely with the clinical team?
	□ Yes □ No □ Unknown
	HYST 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
	5y ····
	Is checklist/bundle adherence shared routinely with the clinical team?
	·
_	☐ Yes ☐ No ☐ Unknown
	None of the above



ye : 20:	ar̈́? *The	clifty (or any part of yer following prevention A/IDSA/APIC Practice	strategies are	e examples	from HAI pre	vention gu	ıidance document	ts (for example,
101		Yes		No		Unk	known	
If y	es, ched	ck all HAIs that apply						
	CLABS	Other (specify):	aily assessme line insertion s ne insertion e-containing d ontaining caps r antimicrobial	supplies to e ressings for covers for c	ensure efficier central lines central line po	nt access in patients		
	CAUTI	(check all that apply Documentation of d Bundling of indwellin access to supplies to Implementation of a automatic stop order an indwelling urinar Process for consider selected patients will Incorporation of approper of standardized Other (specify):	aily assessmeng urinary cathor aseptic indicates a nurse-driven by catheter eration of blade then appropriate indicational propriate institutional programments.	neter inserti welling urina indwelling u view of curi der manage te ations for ur	on supplies in ary catheter in urinary cathet rent indication ment alternate ine culturing	n conveniensertion er removans and ren ives to inc	ent location to ens al protocol or imple newal of order for d dwelling urethral c	ementation of continuation of atheterization in
	CDI La	bID Event (check all Use of an EPA-regior use of additional light disinfection) Establish process in cleaning Restriction of antibious and 4th general Implementation of launformed stool) or a difficile testing Implementation of lacare providers and Other (specify):	stered (EPA L disinfection of a collaboration otics with the h tion cephalosp aboratory prote a clinical decis aboratory alert infection contr	with environighest risk orins) ocol to ensuion support	t rooms with a nmental serv for CDI (for e are testing of system to he immediately r	no-touch to ices to rou xample, flo only appro lp reduce	echnologies (for e utinely assess ade uoroquinolones, c opriate specimens unnecessary Clos	example, UV equacy of room carbapenems, s (for example, stridioides



		Process for monitoring and validation of compliance of daily CHG bathing in applicable patient
		populations (for example, adult ICU patients)
		Process for multidisciplinary review of occurrences of hospital-onset MRSA bacteremia (for example,
		root cause analysis) to assess modifiable risk factors
		Establish process in collaboration with environmental services to routinely assess adequacy of room
		cleaning
		Implementation of a laboratory-based alert system that immediately notifies clinical care providers
		and infection control personnel of new MRSA-colonized and/or MRSA-infected patients
		Implementation of universal gowns and gloves upon entry into adult ICU patient rooms, regardless of
		MRSA status Other (aposity):
		Other (specify):
	COLO	SSI (check all that apply)
_		Use of combination of parenteral and oral antimicrobial prophylaxis with mechanical bowel prep,
		unless contraindicated, prior to elective colorectal surgery
		Monitor compliance with antimicrobial prophylaxis guidelines being appropriately provided
		Use of impervious plastic wound protectors for GI surgery
		Implementation of preoperative warming for at least 30 minutes prior to surgery to prevent
		intraoperative hypothermia
		Use of negative pressure dressings in patients who may benefit Use of antiseptic-impregnated sutures
		Other (specify):
		Other (aposity).
	HYST	SSI (check all that apply)
		Use antiseptic-containing preoperative vaginal preparatory agents for patients undergoing elective
		hysterectomy
		Monitor compliance with antimicrobial prophylaxis guidelines being appropriately provided
		Monitor compliance with antimicrobial prophylaxis guidelines being appropriately provided Implementation of preoperative warming for at least 30 minutes prior to surgery to prevent
		Monitor compliance with antimicrobial prophylaxis guidelines being appropriately provided Implementation of preoperative warming for at least 30 minutes prior to surgery to prevent intraoperative hypothermia
		Monitor compliance with antimicrobial prophylaxis guidelines being appropriately provided Implementation of preoperative warming for at least 30 minutes prior to surgery to prevent intraoperative hypothermia Use of negative pressure dressings in patients who may benefit
		Monitor compliance with antimicrobial prophylaxis guidelines being appropriately provided Implementation of preoperative warming for at least 30 minutes prior to surgery to prevent intraoperative hypothermia Use of negative pressure dressings in patients who may benefit Use of antiseptic-impregnated sutures
		Monitor compliance with antimicrobial prophylaxis guidelines being appropriately provided Implementation of preoperative warming for at least 30 minutes prior to surgery to prevent intraoperative hypothermia Use of negative pressure dressings in patients who may benefit
*54.Doe		Monitor compliance with antimicrobial prophylaxis guidelines being appropriately provided Implementation of preoperative warming for at least 30 minutes prior to surgery to prevent intraoperative hypothermia Use of negative pressure dressings in patients who may benefit Use of antiseptic-impregnated sutures
*54.Doe role	======================================	Monitor compliance with antimicrobial prophylaxis guidelines being appropriately provided Implementation of preoperative warming for at least 30 minutes prior to surgery to prevent intraoperative hypothermia Use of negative pressure dressings in patients who may benefit Use of antiseptic-impregnated sutures Other (specify):
	es your	Monitor compliance with antimicrobial prophylaxis guidelines being appropriately provided Implementation of preoperative warming for at least 30 minutes prior to surgery to prevent intraoperative hypothermia Use of negative pressure dressings in patients who may benefit Use of antiseptic-impregnated sutures Other (specify):
	es your	Monitor compliance with antimicrobial prophylaxis guidelines being appropriately provided Implementation of preoperative warming for at least 30 minutes prior to surgery to prevent intraoperative hypothermia Use of negative pressure dressings in patients who may benefit Use of antiseptic-impregnated sutures Other (specify): facility provide training and/or education on HAI prevention to healthcare personnel as it relates to their
	es your	Monitor compliance with antimicrobial prophylaxis guidelines being appropriately provided Implementation of preoperative warming for at least 30 minutes prior to surgery to prevent intraoperative hypothermia Use of negative pressure dressings in patients who may benefit Use of antiseptic-impregnated sutures Other (specify): facility provide training and/or education on HAI prevention to healthcare personnel as it relates to their
role	es your ?	Monitor compliance with antimicrobial prophylaxis guidelines being appropriately provided Implementation of preoperative warming for at least 30 minutes prior to surgery to prevent intraoperative hypothermia Use of negative pressure dressings in patients who may benefit Use of antiseptic-impregnated sutures Other (specify): facility provide training and/or education on HAI prevention to healthcare personnel as it relates to their Yes
	es your ?? If y	Monitor compliance with antimicrobial prophylaxis guidelines being appropriately provided Implementation of preoperative warming for at least 30 minutes prior to surgery to prevent intraoperative hypothermia Use of negative pressure dressings in patients who may benefit Use of antiseptic-impregnated sutures Other (specify): facility provide training and/or education on HAI prevention to healthcare personnel as it relates to their Yes
role	es your ?? If y	Monitor compliance with antimicrobial prophylaxis guidelines being appropriately provided Implementation of preoperative warming for at least 30 minutes prior to surgery to prevent intraoperative hypothermia Use of negative pressure dressings in patients who may benefit Use of antiseptic-impregnated sutures Other (specify): facility provide training and/or education on HAI prevention to healthcare personnel as it relates to their Yes No Unknown Ves, check all HAIs that apply.
role	es your e? If y	Monitor compliance with antimicrobial prophylaxis guidelines being appropriately provided Implementation of preoperative warming for at least 30 minutes prior to surgery to prevent intraoperative hypothermia Use of negative pressure dressings in patients who may benefit Use of antiseptic-impregnated sutures Other (specify): facility provide training and/or education on HAI prevention to healthcare personnel as it relates to their Yes
role	es your ? If y CLABS At wha	Monitor compliance with antimicrobial prophylaxis guidelines being appropriately provided Implementation of preoperative warming for at least 30 minutes prior to surgery to prevent intraoperative hypothermia Use of negative pressure dressings in patients who may benefit Use of antiseptic-impregnated sutures Other (specify): facility provide training and/or education on HAI prevention to healthcare personnel as it relates to their Yes
role	es your ?? If y CLABS At wha	Monitor compliance with antimicrobial prophylaxis guidelines being appropriately provided Implementation of preoperative warming for at least 30 minutes prior to surgery to prevent intraoperative hypothermia Use of negative pressure dressings in patients who may benefit Use of antiseptic-impregnated sutures Other (specify): facility provide training and/or education on HAI prevention to healthcare personnel as it relates to their Yes
role	es your ?? If y CLABS At wha	Monitor compliance with antimicrobial prophylaxis guidelines being appropriately provided Implementation of preoperative warming for at least 30 minutes prior to surgery to prevent intraoperative hypothermia Use of negative pressure dressings in patients who may benefit Use of antiseptic-impregnated sutures Other (specify): facility provide training and/or education on HAI prevention to healthcare personnel as it relates to their Yes



CAUTI At what frequency is training or education is provided? Check all that apply. Upon hire When new product or processes are implemented Quarterly Yearly PRN Other
CDI LabID Event At what frequency is training or education is provided? Check all that apply. Upon hire When new product or processes are implemented Quarterly Yearly PRN Other
MRSA Bacteremia LabID Event At what frequency is training or education is provided? Check all that apply. Upon hire When new product or processes are implemented Quarterly Yearly PRN Other
COLO SSI At what frequency is training or education is provided? Check all that apply. Upon hire When new product or processes are implemented Quarterly Yearly PRN Other
HYST SSI At what frequency is training or education is provided? Check all that apply. Upon hire When new product or processes are implemented Quarterly PRN PRN Other