

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

# 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): ☐ For profit □ Not for profit, including church □ Government □ Veterans Affairs \*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 ☐ Residential facility (assisted living ☐ Other (specify): \_\_\_\_\_ 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): \*Number of single occupancy rooms: \_\_\_\_\_ \*Number of double occupancy rooms: \_\_\_\_\_ \*Number of triple occupancy rooms: \_\_\_\_\_ 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)). Public reporting burden of this collection of information is estimated to average 89 minutes per response, including the time for reviewing instructions, searching existing data sources, gathering, and maintaining the data needed, and completing and reviewing the collection of information. An agency may not conduct or sponsor, and a person is not required to respond to a collection of information unless it displays a currently valid OMB control number. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to CDC, Reports Clearance Officer, 1600 Clifton Rd., MS



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*Number of quadruple occupancy	rooms:		www.cuc.gov/iiisii
*Total number of admissions with ont developing during LTAC stay):		•	admission (present of admission,
If helpful for your facility in identify associated with these conditions for a. Ventilator dependence: b. Hemodialysis:	ound here: http://www.cdc.go		
Facility Microbiology Laborato	ry Practices (completed wi	th input from Microbiol	ogy Laboratory Lead)
<ul><li>*1. Does your facility have its susceptibility testing?</li><li>1a. If No, where is your fa</li></ul>	own on-site laboratory that p		
☐ Affiliated medical center	er □ Commercial refer	•	er local/regional, non-affiliated nce laboratory
*2. For the following organism (1) Primary susceptibility (2) Secondary, suppleme	testing and ntal, or confirmatory testing ( perform susceptibility testing	re used for: (if performed). g, indicate the methods u	sed at the outside laboratory.
Pathogen (	1) Primary	(2) Secondary	Comments
Enterobacterales _	· · · · · · · · · · · · · · · · · · ·		
Pseudomonas aeruginosa _			
Acinetobacter baumanni complex			
1 = Kirby-Bauer disk diffusion	4 = Sensititre	7 = Agar diluti	on method
2 = Vitek (Legacy)	5.1 = MicroScan WalkAway	y 10 = Gradient	Dilution Strip (for example E test)
2.1 = Vitek 2	5.2 = MicroScan autoSCAN	N 13 = Other (de	escribe in Comments section)
3.1 = BD Phoenix	6 = Other broth microdilution	on method	
*3. Does either the primary or (check all that apply):	secondary/supplemental an	timicrobial susceptibility t	esting (AST) include the following

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Drug	Enterobacterales	Organism tested: Pseudomonas aeruginosa	Acinetobacter baumanni
Cefiderocol			
Ceftazidime-Avibactam			
Ceftolozane-Tazobactam			
Colistin			
Delafloxacin			
Eravacycline			
Imipenem-Relebactam			
Meropenem-Vaborbactam			

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# **Facility Microbiology Laboratory Practices (continued)**

*4.	Has	the	e laboratory implemented revised	breakpoints recommended by CLSI for	the following:
	a.		ird Generation Cephalosporin and <i>terobacterale</i> s <u>in</u> 2010	d monobactam (that is, aztreonam) brea	kpoints for ☐ Yes ☐ No
	b.	Ca	rbapenem breakpoints for <i>Entero</i>	<i>bacterales</i> <u>in</u> 2010	□ Yes □ No
	c.	Ert	apenem breakpoints for <i>Enteroba</i>	acterales <u>in</u> 2012	□ Yes □ No
	d.	Са	rbapenem breakpoints for <i>Pseud</i>	omonas aeruginosa <u>in</u> 2012	□ Yes □ No
	e.	Flu	roquinolone breakpoints for <i>Pseu</i>	idomonas aeruginosa <u>in</u> 2019	□ Yes □ No
	f.	Flu	roquinolone breakpoints for <i>Ente</i>	robacterales <u>in</u> 2019	□ Yes □ No
*5.	not	incl	lude automated testing instrumen	penemase production is detected: (chec	
			Report carbapenem MIC results	without an interpretation	
			infection control practices	erpretation of carbapenems, the rest is u	
	5b.	If Y	es, which test is routinely perforr	ned to detect carbapenemase: (check a	I that apply)
			□ NAAT (for example, PCR)	□ MLB Screen	□ mCIM/CIM
			☐ Modified Hodge Test	□ Carba NP	□ CARBA 5
			□ Rapid CARB Blue	□ Cepheid, BioFire, Verigene, Genmar	k, etc
			□ E test	□ Other (specify):	
*6.	Doe resi	□ es y	Enterobacterales spp. our facility use commercial or lab	ntinely tested for the presence of carbape □ Pseudomonas aeruginosa oratory developed tests for rapid molecu eam infections? Examples of commercia c.	☐ Acinetobacter baumannii
			Yes		
	6a.		No [if checked, skip questions 7 'es, which test panel(s) does you	and 8] r facility use? (check all that apply)	
			Cepheid Xpert MRSA/SA BC GenMark ePlex BCID-FP	☐ GenMark ePlex BCID-GP ☐ G	ioFire FilmArray BCID II enMark ePlex BCID-GN uminex Verigene BC-GN

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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)



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# **Facility Microbiology Laboratory Practices (continued)**

testing in a blood specimen, select the procedure(s) your facility conducts. (check one)
☐ Our laboratory does not perform <i>mecA</i> testing using rapid molecular methods. [If checked, skip question 7a.]
☐ Culture based phenotypic antimicrobial susceptibility testing is not performed. [If checked, skip question 7a.]
☐ Culture based phenotypic antimicrobial susceptibility testing is performed. A text indicating results of the corresponding rapid molecular testing and/or the interpretation of the rapid molecular testing result is added to the phenotypic test result.
☐ Culture based phenotypic antimicrobial susceptibility testing is performed. No text indicating corresponding rapid molecular testing and/or interpretation is added.
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 separately.
☐ 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.
*8. In a scenario where the <i>bla<sub>CTX-M</sub></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)
$\Box$ Our laboratory does not perform $bla_{CTX-M}$ (CTX-M) testing using rapid molecular methods. [If checked, skip questions 8a]
☐ Culture based phenotypic antimicrobial susceptibility testing is not performed. [If checked, skip question 8a.]
☐ Culture based phenotypic antimicrobial susceptibility testing is performed. A text indicating results of the corresponding rapid molecular testing and/or the interpretation of the rapid molecular testing result is added to the phenotypic test result.
☐ 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 antimicrobial susceptibility testing are performed for a blood specimen to detect drug resistance in <i>Escherichia coli</i> and discordance is found between their results, how are results reported? (check one)
☐ Further testing is not pursued. Results are reported separately.
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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)).

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\*7. In a scenario where the mecA resistance marker and Staphylococcus aureus are detected by rapid molecular



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		Further testing is not pursued. The phenotypic result is overridden by the rapid molecular an antimicrobial resistance marker is detected.	test result when		
		Further testing is performed to identify the reason for the discordance. Results are modified further analysis.	ed based on the		
*9.	Does y	our facility perform extended-spectrum beta-lactamase (ESBL) testing for E. coli, Klebsiella	ir facility perform extended-spectrum beta-lactamase (ESBL) testing for <i>E. coli, Klebsiella pneumoniae</i> ,		
	Klebsie	ella oxytoca, or Proteus mirabilis routinely or using a testing algorithm?	□ Yes □ No		
is collecte	ed with a gu	entiality: The voluntarily provided information obtained in this surveillance system that would permit identification of any interaction are that it will be held in strict confidence, will be used only for the purposes stated, and will not otherwise be disclosed dual, or the institution in accordance with Sections 304, 306 and 308(d) of the Public Health Service Act (42 USC 242b, 242l)	or released without the		



# **Facility Microbiology Laboratory Practices (continued)**

9a. If Yes, indicate what is done if ESBL is dete	ected: (check one)		
☐ Change susceptible Cefotaxime/Ceftria	exone/Cefepime results to resistant		
□ No changes are made in the interpretation of cephalosporins with a note of ESBL			
☐ Suppress cephalosporin susceptibility r			
*10. Where is yeast identification performed for spe	cimens collected at your facility? (check one)		
□ On-site laboratory			
☐ Affiliated medical center			
☐ Commercial referral laboratory			
☐ Other local/regional, non-affiliated reference	laboratory		
☐ Yeast identification not available (specifically affiliate/commercial/other laboratory) [If checked	v, yeast identification is not performed onsite or at any d, skip questions 11-15]		
Answer questions 11-15 for the laboratory *11. Which of the following methods are used for ye	that <u>performs yeast identification for your facility</u> : ast identification? (check all that apply)		
☐ MALDI-TOF MS System (Vitek MS)	☐ MicroScan		
☐ MALDI-TOF MS System (Bruker Biotyper)	☐ Non-automated Manual Kit (for example, API 20C, RapID, Germ Tube, PNA-FISH, etc.)		
☐ Vitek-2	□ DNA sequencing		
☐ BD Phoenix	□ Other (specify):		
*12.Does the laboratory routinely use chromogenic	agar for the identification or differentiation of Candida isolates?		
□ Yes □ No	□ Unknown		
*13. Candida isolated from which of the following bo that apply)	ody sites are usually fully identified to the species level? (check all		
☐ Blood	☐ Respiratory		
☐ Other normally sterile body site (for example	e, CSF)		
☐ Urine	$\hfill\square$ None are fully identified to the species level		
*14.Does the laboratory employ any molecular tests	s to identify Candida from blood specimens?		
□ Yes □ No	□ Unknown		
<ul> <li>14a. If yes, which molecular tests are used t</li> <li>□ T2Candida Panel</li> <li>□ BioFire BCID</li> <li>□ GenMark ePlex BCID</li> </ul>	to identify Candida from blood specimens? (check all that apply)		

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ETY NETW	VORK	Exp. Date: 12/31/2026 www.cdc.gov/nhsn
	Other, specify:	<u></u>
□ 14b.	Unknown If yes and you get a positive result, does this lab culture the blood to obtain an isolate?	
	Yes, always	
	Yes, with clinical order	
	No	
	Unknown	

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**Facility Microbiology Laboratory Practices (continued)** 

*15.Where is antifungal susceptible	lity testing (AFST) performe	ed for specimens co	llected at your facility	r? (check one)	
☐ On-site laboratory	☐ Other local/regiona	$\ \square$ Other local/regional, non-affiliated refe		erence laboratory	
☐ Affiliated medical center		☐ AFST not available (specifically, AFST affiliate/commercial/other laboratory) [if s			
☐ Commercial reference laborato	ry affiliate/commercial/o			ns 16 -19]	
Answer questions 16-19 for the I	aboratory that <u>perform</u>	s AFST for your t	<i>facility</i> :		
*16.What method is used for antifu apply)	ingal susceptibility testing (	AFST), <b>excluding A</b>	<b>Amphotericin B</b> ? (ch	neck all that	
<ul> <li>☐ Broth microdilution with laboratory developed plates</li> </ul>	□ YeastOne (Therm Sensititre™)	`		☐ Gradient diffusion (E test)	
☐ Vitek (bioMerieux)	☐ Other (specify): _		☐ Unknown		
*17.What method is used for antifu	ıngal susceptibility testing (	AFST) of <i>Amphote</i>	ricin B? (check all th	at apply)	
☐ Broth microdilution with laboratory developed plates	YeastOne (Therm Sensititre™)	no Scientific™	☐ Gradient diffusion (E test)		
☐ Vitek (bioMerieux)	☐ Other (specify): _	☐ Other (specify):		□ Unknown	
*18.AFST is performed for which o	of the following antifungal di	rugs? (check all that	apply)		
☐ Fluconazole	□ Voriconazol	Э	☐ Itraconazole		
☐ Posaconazole	☐ Micafungin		☐ Anidulafungin		
☐ Caspofungin	☐ Amphoterici	n B	☐ Flucytosine		
□ Other, specify:	□ Unknown				
*19.AFST is performed on fungal i	solates in which of the follo	wing situations? (ch	eck all that apply)		
, .	Performed automatically	Performed with a clinician's order	Not performed	Unknown	
Blood					
Other normally sterile body site (for example, CSF)					
Urine					
Respiratory					
Other (specify):					

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	laboratory developing antibiograms in this laboratory?	or other reports to track susceptibility trends for Candida spp. isolates	
□ Yes	□ No	□ Unknown	
	is the primary testing method for <i>C.</i> tory where your facility's testing is p	difficile used most often by your facility's laboratory or the outside erformed? (check one)	
	Enzyme immunoassay (EIA) for to	oxin	
	Cell cytotoxicity neutralization ass	ay	
	Nucleic acid amplification test (NA	AAT) (for example, PCR, LAMP)	
Facility Micro	obiology Laboratory Practices (co	ontinued)	
	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	n)	
	GDH plus EIA for toxin, followed b	by NAAT for discrepant results	
	Toxigenic culture (C. difficile cultu	re followed by detection of toxins)	
	\		
*22.Indicat	te the primary and definitive method	used to identify microbes from blood cultures collected in your facility.	
(check	cone)		
	MALDI-TOF MS System (Vitek MS	S)	
	MALDI-TOF MS System (Bruker E	Biotyper)	
	Automated Instrument (for examp	le, Vitek, MicroScan, Phoenix, OmniLog, Sherlock, etc.)	
	Non-automated Manual Kit (for ex	ample, API, Crystal, RapID, etc.)	
	□ Rapid Identification (for example, Verigene, BioFire FilmArray, PNA-FISH, Gene Xpert, etc.)		
	16S rRNA Sequencing		
	□ Other (specify):		
		s used for microbe identification from blood cultures collected in your	
•	•	s confirmed with the primary methods, a secondary method if the	
	y method fails to give an identification (all that apply)	on, or a method that is used in conjunction with the primary method).	
•			
	MALDI-TOF MS System (Vitek MS	•	
	MALDI-TOF MS System (Bruker E		
	·	le, Vitek, MicroScan, Phoenix, OmniLog, Sherlock, etc.)	
	Non-automated Manual Kit (for example	Verigene, BioFire FilmArray, PNA-FISH, Gene Xpert, etc.)	
	16S rRNA Sequencing	verigene, Diorne Filinanay, Fiva-Fion, Gene Apen, etc.)	
_			
	Other (specify):	<del></del>	
essurance of Confid	dontiality. The voluntarily provided information	obtained in this surveillance system that would permit identification of any individual or institution	

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□ None

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

er or faction of infection preventionists (IPs) in facility:  Total hours per week performing surveillance:  Total hours per week for infection control activities other than surveillance:  er or fraction of full-time employees (FTEs) for a designated hospital epidemiologist (or equivalent role)  ed with your facility:  poolicy in your facility that patients infected or colonized with MRSA are routinely placed in contact			
utions while these patients are in your facility? (check one)  ☐ No ☐ Not applicable: my facility never admits these patients			
□ Not applicable. Thy facility flever admits these patients			
ntrol Practices (continued)			
If Yes, check the type of patients that are routinely placed in contact precautions while in your facility			
(check one):			
☐ All infected and all colonized patients			
□ Only all infected patients			
□ Only infected or colonized patients with certain characteristics (check all that apply)			
□ Patients admitted to high risk settings			
□ Patients at high risk for transmission			
policy in your facility that patients infected or colonized with VRE are routinely placed in contact precautions hese patients are in your facility? (check one)			
□ No □ Not applicable: my facility never admits these patients			
If Yes, check the type of patients that are routinely place in contact precautions while in your facility neck one):			
All infected and all colonized patients			
Only all infected patients			
Only infected or colonized patients with certain characteristics (check all that apply)			
□ Patients admitted to high risk settings			
□ Patients at high risk for transmission			
- 1 duono de mgn not for transmission			

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carbap (check	,	ly placed in contact precautions while these patients are in your facility?	
□ Yes	□ No	☐ Not applicable: my facility never admits these patients	
28a. (ch	If Yes, check the type of patient neck one):	ts that are routinely placed in contact precautions while in your facility	
	All infected and all colonized pa	atients	
	Only all infected patients		
	Only infected or colonized patie	ents with certain characteristics (check all that apply)	
	$\ \square$ Patients admitted to high	risk settings	
	☐ Patients at high risk for tra	ansmission	
extend		infected or colonized with suspected or confirmed ESBL-producing or tant <i>Enterobacterales</i> are routinely placed in contact precautions while k one)	
□ Yes	□ No	☐ Not applicable: my facility never admits these patients	
29a. (ch	If Yes, check the type of patients that are routinely placed in contact precautions while in your facility heck one):		
	All infected and all colonized pa	atients	
	Only all infected patients		
	Only infected or colonized patie	ents with certain characteristics (check all that apply)	
	□ Patients admitted to high	risk settings	
	☐ Patients at high risk for tra	ansmission	
Infection Cor	ntrol Practices (continued)		
		ening testing (culture or non-culture) for CRE? This includes screening for ablic health laboratories and commercial laboratories.	
		□ Yes □ No	
30а. ар	If Yes, in which situations does ply)	the facility routinely perform screening testing for CRE? (check all that	
	Surveillance testing at admission	on for all patients	
	Surveillance testing of epidemic roommates)	ologically-linked patients of newly identified CRE patients (for example,	
	Surveillance testing at admission	on of high-risk patients (check all that apply)	
	☐ Patients admitted form long	g-term acute care (LTAC) or long-term care facility (LTCF)	
is collected with a gu	narantee that it will be held in strict confidence	on obtained in this surveillance system that would permit identification of any individual or institution te, will be used only for the purposes stated, and will not otherwise be disclosed or released without the ctions 304, 306 and 308(d) of the Public Health Service Act (42 USC 242b, 242k, and 242m(d)).	
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\*28.Is it a policy in your facility that patients infected or colonized with CRE (regardless of confirmatory testing for



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	☐ Patients with recent (for example, within 6 months) overnight nospital 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)
□ 30b. fac	Other (specify): If Yes, what method is routinely used by the lab conducting CRE testing of screening swabs from your ility? (check all that apply)
	Culture-based methods
	ne facility routinely perform screening testing (culture or non-culture) for <i>Candida auris</i> ? This includes ing for patients at your facility performed by public health laboratories and commercial laboratories.  □ Yes □ No
31a. all	If Yes, in which situations does the facility routinely perform screening testing for <i>Candida auris</i> ? (check 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):
	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):
31b. fro	If Yes, what method is routinely used by the lab conducting <i>Candida auris</i> testing of screening swabs m your facility?
	Culture-based methods
*32.Does tl	ne facility routinely perform screening testing (culture or non-culture for MRSA for any patients admitted?
	□ Yes □ No
	ntrol Practices (continued)
32a.	If Yes, in which situations does the facility routinely perform screening testing for MRSA? (check all that
	Oly)
	Surveillance testing at admission for all patients
is collected with a gu	entiality: The voluntarily provided information obtained in this surveillance system that would permit identification of any individual or institution tarantee 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 dual, 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)).
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[LTAC] or long-term care facility [	of nign-risk patients (for example, admitted to the control of the	ed from long-term acute care
	of patients admitted to high-risk settings	(for example ICII)
_	ive patients to prevent surgical site infect	` '
☐ Other (specify):	ivo pationio to provent cargical cito imeot	
*33.Does your facility have a policy to routine	 ly use chlorhexidine bathing for any adult	patients to prevent infection or
transmission of MDROs at your facility?		•
		□ Yes □ No
33a. If Yes, indicate which patients: (s	elect all that apply)	
□ ICU patients:	☐ Patients outside the ICU:	☐ Pre-operatively for
∘ All ICU patients	○ All ICU patients	patients undergoing
<ul> <li>Subset of ICU patients:</li> </ul>	○ Subset of ICU patients:	surgery
□ Patients with central venous	□ Patients with central venous	
catheter or midline catheters	catheter or midline catheters	
□ Other, specify:	□ Other, specify:	
· · · · · ·		
*34.Does the facility have a policy to routinely		
antistaphylococcal agent (mupirocin, iodo prevent healthcare-associated infections		
prevent nearincare-associated injections	or reduce transmission or resistant patho	gens≀ □ Yes □ No
A (11. 11. D) (1. 1. 1. 1.		
Antibiotic Stewardship Practices (completed	•	•
*35.Did the antibiotic stewardship leader(s) pa		s? (check one)
□ Yes, pharmacist lead □ Yes	, both pharmacist and physician leads	□ Yes, other lead
☐ Yes, physician lead ☐ No		
*36.Facility leadership has demonstrated con	•	• • • • • • • • • • • • • • • • • • • •
☐ Providing stewardship program leader(s)	dedicated time to manage the program ar	nd conduct daily stewardship
interventions.		
□ Allocating resources (for example, IT supplements)	port, training for stewardship team) to sup	port antibiotic stewardship
☐ Having a senior executive that serves as	a point of contact or "champion" to help er	neure the program has
resources and support to accomplish its mis-	·	isure the program has
☐ Presenting information on stewardship ac		and/or board at least annually.
☐ Ensuring the stewardship program has an	· ·	•
board at least annually.		
□ Communicating to staff about stewardship	activities, via email, newsletters, events,	or other avenues.
☐ Providing opportunities for hospital staff tr	aining and development on antibiotic stev	vardship.
	The train of the second	and the second second second
Assurance of Confidentiality: The voluntarily provided information is collected with a guarantee that it will be held in strict confidence,	*	· · · · · · · · · · · · · · · · · · ·
consent of the individual, or the institution in accordance with Section		
Public reporting burden of this collection of information is estimated	1 to average 89 minutes per response, including the time	for reviewing instructions, searching

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approved	by the board).		wardship (for example, a written policy or statement
	ng that staff from key ng to stewardship ac		oups (for example, IT and hospital medicine) are
	of the above.		
Antibiotic St	ewardship Practice	s (continued)	
*27 Our fo	oility boo a loader er	aa laadara raananaihla far ar	tibiotic stowardship program management and outcomes
37.Our la	cility has a leader or	co-leaders responsible for ar	ntibiotic stewardship program management and outcomes.  □ Yes □ No
37a.	If Yes, what is the	position of this leader? (chec	
	Physician	☐ Co-led by both Pharm	•
	Pharmacist	•	N, PA, NP, etc.; specify):
37b.		ed is selected, which of the f	ollowing describes your antibiotic stewardship <b>physician</b>
	Has antibiotic stew	ardship responsibilities in the	ir contract or job description or performance review
	Is physically on-site	e in your facility (either part-ti	me or full-time)
	Completed an ID fe	ellowship	
	Completed a certifi	cate program on antibiotic st	ewardship
	Completed other tr	aining(s) (for example, confe	rences or online modules) on antibiotic stewardship
	None of the above		
•	o) leader): What perd		their contract or job description' is selected (for physician stewardship activities is specified in the <b>physician</b> (co)
	□ 1-10%	□ 11-25%	□ 26-50%
	□ 51-75%	□ 76-100%	☐ Not specified
37d. lea	•	ed is selected: <b>In an averag</b> otic stewardship activities in	e week, what percentage of time does the physician (co) your facility? (check one)
	□ 1-10%	□ 11-25%	□ 26-50%
	□ 51-75%	□ 76-100%	
37e. <b>ph</b>	If Pharmacist or Conarmacist leader? (c		following describes your antibiotic stewardship
	Has antibiotic stew	ardship responsibilities in the	eir contract, job description or performance review
	Is physically on-site	e in your facility (either part-ti	me or full-time)
	Completed a PGY	2 ID residency and/or ID fello	wship
			veillance system that would permit identification of any individual or institution r the purposes stated, and will not otherwise be disclosed or released without the

A is 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)).



	Completed a certifica	te program on antibiotic s	tewardship	
	Completed other train	ning(s) (for example, confe	erences or online modules) o	n antibiotic stewardship
	None of the above			
(co	) leader): What percer	·	stewardship activities is spec	n' is selected (for pharmacist cified in the <b>pharmacist</b> (co)
	□ 1-10%	□ 11-25%	□ 26-50%	
	□ 51-75%	□ 76-100%		
37g.				e of time does the <b>pharmacist</b>
(co	•	•	es in your facility? (check one	e)
	□ 1-10% □ 54.75°/	□ 11-25% □ 72.4000/	□ 26-50%	
•	□ 51-75%	□ 76-100%		
	ewardship Practices	•		
37h.			acility have a designated phy	sician who can serve as a
рог	nt of contact and supp	ort for the non-physician I	eader?	- W - N
07: If -				☐ Yes ☐ No
	r pnarmacist is <b>not</b> the proving antibiotic use a		e program, is there at least o	ne pharmacist responsible for
11114	ordering antibiotic use a	t your facility!		□ Yes □ No
*38.Our fac	cility has the following p	oriority antibiotic stewards	hip interventions: (Check all	
□ Pros	pective audit and feed	back for specific antibiotic	agents	
38a. foll	•		For which categories of antim ot they are on formulary. (Ch	
	Cefepime, ceftazidim	e, or piperacillin/tazobacta	am	
	Vancomycin (intraver	nous)		
	Ertapenem, imipener	n/cilastatin, or meropenen	1	
	Ceftazidime/avibacta cilastatin/relebactam,		m, meropenem/vaborbactam	, imipenem-
	Fluoroquinolones			
	Daptomycin, linezolid	, or other newer anti-MRS	SA agents	
	Eravacycline or omac	lacycline		
	Lefamulin			
	Aminoglycosides			
	Colistin or polymyxin	В		
	Anidulafungin, caspo			
	entiality: The voluntarily prov	ided information obtained in this su	, ,	entification of any individual or institution

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	Isavuconazole, posaconazole, or voriconazole
	Amphotericin B and/or lipid-based amphotericin B
	None of the above
	If Prospective audit and feedback is selected: Our antibiotic stewardship program monitors prospective dit and feedback interventions (for example, by tracking antibiotic use, types of interventions, acceptance of commendations).
	Yes □ No
□ Prea	uthorization for specific antibiotic agents.
38c.	If Preauthorization is selected: For which categories of antimicrobials? Only answer for categories of
	imicrobials that are <i>on formulary</i> . (Check all that apply)
	Cefepime, ceftazidime, or piperacillin/tazobactam
	Vancomycin (intravenous)
	Ertapenem, imipenem/cilastatin, or meropenem
	Ceftazidime/avibactam, ceftolozane/tazobactam, meropenem/vaborbactam, imipenem-cilastatin/relebactam, or cefiderocol
	Fluoroquinolones
	Daptomycin, linezolid, or other newer anti-MRSA agents
	Eravacycline or omadacycline
	Lefamulin
ntibiotic Ste	ewardship Practices (continued)
	Aminoglycosides
	Colistin or polymyxin B
	Anidulafungin, caspofungin, or micafungin
	Isavuconazole, posaconazole, or voriconazole
	Amphotericin B and/or lipid-based amphotericin B
	None of the above
38d. (foi	If Preauthorization is selected: Our antibiotic stewardship program monitors preauthorization interventions rexample, by tracking which agents are requested for which conditions).
	□ Yes □ No
□ 38e.	Facility-specific treatment recommendations, based on national guidelines and local pathogens susceptibilities, to assist with antibiotic selections for common clinical conditions (for example, community-acquired pneumonia, urinary tract infection, skin and soft tissue infection).  If Facility-specific treatment recommendations is selected: For which common clinical conditions?
	Community-acquired pneumonia
	Urinary tract infection

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□ SI	kin and soft tissue infection
□ <b>N</b> ∈	one of the above
our fa	cility-specific treatment recommendations is selected: Our stewardship program monitors adherence to acility's treatment recommendations for antibiotic selection for common clinical conditions (for example, nunity-acquired pneumonia, urinary tract infection, skin and soft infections).
	□ Yes □ No
38g. If	Yes: For which common clinical conditions?
□ C	community-acquired pneumonia
□ U	rinary tract infection
□ SI	kin and soft tissue infection
□ <b>N</b>	one of the above
*39.Our facility that apply	y has a policy or formal procedure for other interventions to ensure optimal use of antibiotics: (Check all '.)
□ Early adm	ninistration of effective antibiotics to optimize the treatment of sepsis
□ Treatmen	nt protocols for <i>Staphylococcus aureus</i> bloodstream infection
□ Stopping	unnecessary antibiotic(s) in new cases of Clostridioides difficile infection (CDI)
□ Review of	f culture-proven invasive (for example, bloodstream) infections
□ Review of	f planned outpatient parenteral antibiotic therapy (OPAT)
□ The treati	ing team to review antibiotics 48-72 hours after initial order (specifically, antibiotic time-out)
□ Assess ar	nd clarify documented penicillin allergy
community-	shortest effective duration of antibiotics at discharge for common clinical conditions (for example, acquired pneumonia, urinary tract infection, skin and soft tissue infections)
□ None of th	пе авоче

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#### **Antibiotic Stewardship Practices (continued)**

H21-8, Atlanta, GA 30333, ATTN: PRA (0920-0666).

39a. 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.
*40.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)
☐ 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)
□ None of the above
*41.Our stewardship program has engaged bedside nurses in actions to optimize antibiotic use.
□ Yes □ No
41a. 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.
□ Nurses initiate discussions with the treating team on switching from intravenous to oral antibiotics.
□ Nurses initiate antibiotic time-out discussions with the treating team.
□ Nurses track antibiotic duration of therapy.
□ None of the above.
41b. If 'Nurses track antibiotic duration of therapy' is selected: Is that information available at the bedside (for example, on a whiteboard in the room)?
□ Yes □ No
*42.Our stewardship program monitors: (Check all that apply.)
☐ 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
☐ Antibiotic expenditures (specifically, purchasing costs), at least quarterly
□ Antibiotic use in some other way, at least annually (specify):
□ None of the above
*43.Our stewardship team provides the following antibiotic use reports to prescribers, at least annually: (Check all that apply.)
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		www.cuc	.gov/IIIIsii
□ Individ	ual, prescriber-level reports		
□ Unit- o	r service-specific reports		
□ None o	of the above		
43a.	If 'Individual, prescriber-level reports' or 'Unit-or service-specific reports' is selected: Our	stewardsl	nip
•	gram uses these reports to target feedback to prescribers about how they can improve the	ir antibio	tic
pre	escribing, at least annually.		
		□ Yes	□ No
Antibiotic Ste	ewardship Practices (continued)		
*44.Our fac	cility distributes an antibiogram to prescribers, at least annually.		
		□ Yes	□ No
*45.Informa annual	ation on antibiotic use, antibiotic resistance, and stewardship efforts is reported to hospitally.	staff, at le	east
		□ Yes	□ No
*46.Which	of the following groups receive education on optimal prescribing, adverse reactions from a	ntibiotics,	an
antibiot all that	tic resistance (for example, Grand Rounds, in-service training, direct instruction) at least ar apply.)	ınually? (	Check
□ Prescr	ibers		
□ Nursin	g staff		
□ Pharm	acists		
□ None o	of the above		
	tients provided education on important side effects of prescribed antibiotics?		
		□ Yes	□ No
47a.	If 'Yes' is selected: How is education to patients on side effects shared? (Check all that a	pply.)	
	Discharge paperwork	, ,	
	Verbally by nurse		
	Verbally by pharmacist		
	Verbally by physician		
	None of the above		
Optional Anti	ibiotic Stewardship Practices Questions		
Response to	the following questions are not required to complete the annual survey.		
Provide addit	tional information about your facility antibiotic stewardship activities and leadership	٠.	
48. Antibio	tic stewardship activities are integrated into quality improvement and/or patient safety initia	itives.	
		□ Yes	□ No
is collected with a gu	entiality: The voluntarily provided information obtained in this surveillance system that would permit identification of any interaction are that it will be held in strict confidence, will be used only for the purposes stated, and will not otherwise be disclosed dual, or the institution in accordance with Sections 304, 306 and 308(d) of the Public Health Service Act (42 USC 242b, 242d).	or released v	vithout the
existing data sources and a person is not re estimate or any other	den of this collection of information is estimated to average 89 minutes per response, including the time for reviewing instruct, gathering, and maintaining the data needed, and completing and reviewing the collection of information. An agency may required to respond to a collection of information unless it displays a currently valid OMB control number. Send comments reaspect of this collection of information, including suggestions for reducing this burden to CDC, Reports Clearance Officer, 30333, ATTN: PRA (0920-0666).	ot conduct o	r sponsor, burden



<ol> <li>Our facility accesses targeted rem specific support for our antibiotics</li> </ol>		pertise (for example, tele-stewardship to obt	ain facility-
opeome cappervier car arminette	storrar domp on orto).		□ Yes □ No
<ol><li>Our stewardship program works wall that apply)</li></ol>	vith the microbiology	laboratory to implement the following interv	
☐ Selective reporting of antimicrobia	al susceptibility testir	ng results	
☐ Placing comments in microbiology	y reports to improve	prescribing	
☐ None of the above	. , ,		
<ol><li>Which committees or leadership e all that apply)</li></ol>	entities provide overs	sight of your facility's antibiotic stewardship o	efforts? (Check
□ Pharmacy director	□ Executive leade	ership (for example, CEO, CMO)	
☐ Pharmacy & therapeutics	☐ Hospital board		
□ Patient safety	☐ Other (specify):		
□ Quality improvement	□ None		
Facility Water Management Program	(WMP) (Completed	with input from WMP team members)	
•	waterborne pathoge	(WMP) to prevent the growth and transmissions (for example, <i>Pseudomoas, Acinetobac</i> cobacteria, and fungi)?	
			□ Yes □ No
·	•	IP team? (Check all that apply):	
☐ Hospital Epidemiologist/Infection	n Preventionist	□ Compliance/Safety Officer	
☐ Hospital Administrator/Leadersh	ip	☐ Risk/Quality Management Staff	
□ Facilities Manager/Engineer		□ Infectious Disease Clinician	
□ Maintenance Staff		□ Consultant	
☐ Equipment/Chemical Acquistion	/Supplier	□ Laboratory Staff/Leadership	
□ Environmental Services		□ Other (specifiy):	
opportunistic waterborne pathoge piping infrastructure)? This may in	ns for example could aclude a description o	sessment to identify where <i>Legionella</i> and or d grow and spread in the facility water syste of building water systems using text or basic cessing steps, control measures, and end-u	m (for example, c diagrams that se points.
53a. If Yes, when was the mos	t recent assessment	t conducted? (Check one)	□ Yes □ No
is collected with a guarantee that it will be held in strict co	onfidence, will be used only	surveillance system that would permit identification of any in for the purposes stated, and will not otherwise be disclosed 308(d) of the Public Health Service Act (42 USC 242b, 242b	or released without the
existing data sources, gathering, and maintaining the data and a person is not required to respond to a collection of it	needed, and completing and information unless it displays	inutes per response, including the time for reviewing instruct d reviewing the collection of information. An agency may n s a currently valid OMB control number. Send comments re or reducing this burden to CDC, Reports Clearance Officer,	ot conduct or sponsor, egarding this burden



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☐ Within the most recent year	□ Between 1 and 3 years ago	☐ More than 3 years ago (>	•3
(<1 year ago)	(≥1 year and ≤3 years)	years)	
, , ,	— ,	,	
*54.Has your facility has ever conducted sources, modes of transmission, pa	tient susceptibility, patient exposure	e, and/or program preparedne	ess? An
example WICRA tool can be assess	sed at <a href="https://www.cdc.gov/hai/pdfs/">https://www.cdc.gov/hai/pdfs/</a>		
			Yes □ No
54a. If Yes, when was the most	recent assessment conducted? (Ch	eck one)	
☐ Within the most recent year (<1 year ago)	<ul><li>□ Between 1 and 3 years ago</li><li>(≥1 year and ≤3 years)</li></ul>	☐ More than 3 years ago (> years)	-3
*55.Does your facility regularly monitor	the following parameters in the build	ling water system(s)?	
Disinfectant (such as residual chlori	ine):	□ Yes	□ No
55a. If Yes, does your facility ha	ve a plan for corrective actions wher	n disinfectant(s) are not within	n acceptable
limits as determined by the water	er management program?	□ Yes	□ No
•	uently does your facility monitor disi	nfectant(s)? (Check all that a	pply)
Assurance of Confidentiality: The voluntarily provided info is collected with a guarantee that it will be held in strict con- consent of the individual, or the institution in accordance wi	fidence, will be used only for the purposes stated,	and will not otherwise be disclosed or i	released without the
Public reporting burden of this collection of information is existing data sources, gathering, and maintaining the data ne and a person is not required to respond to a collection of info	eeded, and completing and reviewing the collection	on of information. An agency may not c	conduct or sponsor,

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# Facility Water Management Program (WMP) (continued)

	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):
Daily								
Weekly								
Monthly								
Quarterly								
Annually								
Other (specify):								
acceptable	does you limits as	determined	by the wat	ter manag	ement pro	when water tempe gram? · water temperatur	□ Yes	□ No
	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):
Daily								
Weekly								
Monthly								
Quarterly								
Annually								
Other (specify):								
Water pH:							□ Yes	□ No

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55e. If Yes, does your facility have a plan for corrective actions when water pH i	s not within accepta	ble limits
as determined by the water management program?	□ Yes	□ No
55f. If Yes, where and how frequently does your facility monitor water pH? (check a	ll that apply)	
Assurance of Confidentiality: The voluntarily provided information obtained in this surveillance system that would permit id is collected with a guarantee that it will be held in strict confidence, will be used only for the purposes stated, and will not oth consent of the individual, or the institution in accordance with Sections 304, 306 and 308(d) of the Public Health Service Act	erwise be disclosed or relea	ased without the
Public reporting burden of this collection of information is estimated to average 89 minutes per response, including the time f existing data sources, gathering, and maintaining the data needed, and completing and reviewing the collection of information and a person is not required to respond to a collection of information unless it displays a currently valid OMB control number	. An agency may not cond	uct or sponsor,

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# Facility Water Management Program (WMP) (continued)

	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):
Daily								
Weekly								
Monthly								
Quarterly								
Annually								
Other (specify):								
acceptable	does you limits as where an	r facility har determined d how freque	ve a plan for the wat uently does	er manage your facili	ement pro ity perform	Representative	□ Yes neck all that apply) Representative	□ No ) Other
	Points	Potable Water Storage Tank(s)	Potable Water Storage Tank(s)	Water Supply	Water Return	Locations Throughout Cold Potable Building Water System(s)	Locations Throughout Hot Potable Building Water System(s)	(specify):
Daily								
Weekly								
Monthly								
Quarterly								
Annually								
Other (specify):								
Specific enviro	nmental <i>L</i>	.egionella te	esting:				□ Yes	□ No

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55i. If Yes, does	your facility have a plan fo	r corrective actions whe	en environmental test	s for <i>Legionella</i> a	re no	t
·	otable limits as determined e an how frequently does y	•	. •	□ Yes k all that apply)		No
•		,,	σ ,	,		
is collected with a guarantee that	e voluntarily provided information o it will be held in strict confidence, w astitution in accordance with Section	ill be used only for the purposes	s stated, and will not otherwise	e be disclosed or release	d with	
existing data sources, gathering, a and a person is not required to res	llection of information is estimated that maintaining the data needed, and spond to a collection of information use collection of information, including N: PRA (0920-0666).	completing and reviewing the concless it displays a currently valid	collection of information. An id OMB control number. Sen	agency may not conduct d comments regarding to	t or spo his burd	den



# Facility Water Management Program (WMP) (continued)

	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):
Daily								
Weekly								
Monthly								
Quarterly								
Annually								
Other (specify):								
				•	•		_ ,,	

Spec	cific environmental <i>Pseudomonas</i> testing:	□ Yes	□ No
55k.	If Yes, does your facility have a plan for corrective actions when envare not within acceptable limits as determined by the water management		lomonas
		□ Yes	□ No
55L I	If Yes, where an how frequently does your facility perform <i>Pseudomonas</i>	s testing? (check all that a	nnlv)

	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):
Daily								
Weekly								
Monthly								
Quarterly								
Annually								
Other (specify):								

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

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)).



Form Approved OMB No. 0920-0666 Exp. Date: 12/31/2026

Yes No N/A, my facility does not have a water management program
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)).
Public reporting burden of this collection of information is estimated to average 89 minutes per response, including the time for reviewing instructions, searching existing data sources, gathering, and maintaining the data needed, and completing and reviewing the collection of information. An agency may not conduct or sponsor, and a person is not required to respond to a collection of information unless it displays a currently valid OMB control number. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to CDC, Reports Clearance Officer, 1600 Clifton Rd., MS H21-8, Atlanta, GA 30333, ATTN: PRA (0920-0666).
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