



Instructions for this form are available at: <a href="http://www.cdc.gov/nhsn/forms/instr/TOI-57.151-IRF.pdf">http://www.cdc.gov/nhsn/forms/instr/TOI-57.151-IRF.pdf</a>

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*required for saving	Tracking #:
*Facility ID:	*Survey Year:
Facility Characteristics (completed by Infection Preve	ntionist)
*Ownership (check one):	
☐ For profit ☐ Not for profit, including church	☐ Government ☐ Veterans Affairs
*Affiliation (check one): ☐ Independent ☐ Hospital system	☐ Multi-facility organization (specialty network)
*How would you describe your licensed inpatient rehabilita	ation facility? (check one)
☐ Free-standing	☐ Healthcare facility based
In the previous calendar year, indicate the following count *Total number of rehab beds:  *Average daily census:  *Number of patient days:  *Average length of stay:  *Indicate the number of admissions with the primary diagr (must sum to the total number of admissions listed below)  a. Traumatic spinal cord dysfunction:  b. Non-traumatic spinal cord dysfunction:  c. Stroke:  d. Brain dysfunction (non-traumatic or traumatic):  e. Other neurologic conditions (e.g. multiple sclerosis, f. Orthopedic conditions (incl. fracture, joint replacement) g. All other admissions:	nosis for each of the following rehabilitation categories
*Total number of admissions:  *Number of admissions on a ventilator:  *Number of pediatric (≤ 18 years old) admissions:	
Facility Microbiology Laboratory Practices (completed	d with input from Microbiology Laboratory Lead)
*1. Does your facility have its own on-site laboratory that partial yes   No   If No, where is your facility's antimicrobial susceptibility  Affiliated medical center   Commercial references	testing performed? (check one)
	illance system that would permit identification of any individual or institution is collected with ted, and will not otherwise be disclosed or released without the consent of the individual, or vice Act (42 USC 242b, 242k, and 242m(d)).

Public reporting burden of this collection of information is estimated to average 55 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 D-74, Atlanta, GA 30333, ATTN: PRA (0920-0666).

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Facility Microbiology Laborato	ory Practices (continued)			
*2. For the following organisms please indicate which methods are used for:				
(1) primary susceptibility testing and				
. ,	ental, or confirmatory testing (if perf	,		
If your laboratory does n laboratory.	ot perform susceptibility testing, ple	ease indicate the	e methods used at t	the outside
•	des listed below the table.			
Pathogen		) Secondary	Comment	ts
Staphylococcus aureus				<del></del>
Enterobacteriaceae				<del></del>
1 = Kirby-Bauer disk diffusion	5.1 = MicroScan walkaway rapid	10 = E test		
2 = Vitek (Legacy)	5.2 = MicroScan walkaway conventio		omycin agar screen (B	
2.1 = Vitek 2	5.3 = MicroScan auto or touchscan		(describe in Commen	its section)
3.1 = BD Phoenix	6 = Other micro-broth dilution method	I		
4 = Sensititre	7 = Agar dilution method			
	ted the revised cephalosporin and i iaceae recommended by CLSI as c		☐ Yes	□ No
*4. Has the laboratory implemen Enterobacteriaceae recomm	ted the revised carbapenem break ended by CLSI as of 2010?	points for	☐ Yes	□ No
*5. Does the laboratory perform	a special test for presence of carba	penemase?	☐ Yes ☐ No	
If Yes, please indicate what i	is done if carbapenemase production	on is detected: (	(check one)	
$\square$ Change susceptible	carbapenem results to resistant			
$\square$ Report carbapenem	MIC results without an interpretation	on		
☐ No changes are madinfection control purp	de in the interpretation of carbapend poses	ems, the test is	used for epidemiolo	ogical or
If Yes, which test is routinely	performed to detect carbapenema	se: (check all th	nat apply)	
□PCR	☐ MBL screen			
☐ Modified Hodge Test	☐ Carba NP			
□ E test	Other (specify):			
		<del></del>		Continued >>





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Facility Microbiology Labo	oratory Practices (c	continued)		
*6. Does the laboratory performagative bacilli?	orm colistin or polym	nyxin B susceptibility	testing for drug-resistant gran	n ☐ Yes ☐ No
If Yes, please indicate m	nethods: (check all th	nat apply)		
$\square$ Vitek (Legacy)	$\square$ MicroScan walk	away rapid	$\square$ Agar dilution method	
☐ Vitek 2	$\square$ MicroScan walk	away conventional	☐ E test	
☐ BD Phoenix	☐ MicroScan auto	or touchscan	Other (specify):	
☐ Sensititre	☐ Other micro-bro	th dilution method		
*7. Does your facility have it	s own laboratory tha	at performs antifungal	susceptibility testing for <i>Can</i> d	dida species?
☐ Yes ☐ No				
If No, where is your facil	ity's antifungal susc	eptibility testing perfo	rmed? (check one)	
$\Box$ Affiliated medical	center	[	$\Box$ Commercial referral laborat	tory
$\square$ Other local/region	al, non-affiliated refe	erence laboratory	$\square$ Not offered by my facility	
8. If antifungal susceptibility (check all that apply)	testing is performed	l at your facility or an	outside laboratory, what meth	nods are used?
$\square$ Broth macrodilution	☐ Broth microd	ilution $\square$ YeastOn	e colorimetric microdilution	☐ E test
$\square$ Vitek 2 card	$\square$ Disk diffusion	n 🗆 Other (sp	pecify):	
			rely without needing a specific ecies when cultured from norr	
Candida albicans: 🔲 Y	′es □ No			
If Yes, what antifung	al drugs are tested a	automatically/reflexiv	ely? (check all that apply)	
$\square$ Fluconazole	$\square$ Voriconazole	☐ Anidulafungin/C	aspofungin/Micafungin	
Candida glabrata: 🔲 v		automatically/reflexiv	ely? (check all that apply)	
$\square$ Fluconazole	$\square$ Voriconazole	☐ Anidulafungin/Ca	aspofungin/Micafungin	
	☐ Yes ☐ No al drugs are tested a ☐ Voriconazole		ely? (check all that apply) aspofungin/Micafungin	
Other Candida species:	☐ Yes ☐ No			
·		automatically/reflexiv	ely? (check all that apply)	
☐ Fluconazole	☐ Voriconazole		aspofungin/Micafungin	
			. 5	
$\square$ Automatic testing is n	ot performed for any	Candida species		Continued >>





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Facility Microbiology Laboratory Practices (continued)
*10. 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)
$\square$ Enzyme immunoassay (EIA) for toxin
$\square$ Cell cytotoxicity neutralization assay
☐ Nucleic acid amplification test (NAAT) (e.g., PCR, LAMP)
$\square$ NAAT plus EIA, if NAAT positive (2-step algorithm)
$\square$ Glutamate dehydrogenase (GDH) antigen plus EIA for toxin (2-step algorithm)
☐ GDH plus NAAT (2-step algorithm)
$\square$ 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): ("Other" should not be used to name specific laboratories, reference laboratories, or the brand names of C. difficile tests; most methods can be categorized accurately by selecting from the options provided. Please ask your laboratory or conduct a search for further guidance on selecting the correct option to report.)
*11. Does your facility produce an antibiogram (i.e., cumulative antimicrobial susceptibility report)?
☐ Yes ☐ No
If Yes, is the antibiogram produced at least annually?
☐ Yes ☐ No
If Yes, are data stratified by hospital location?
☐ Yes ☐ No
If No, please identify any obstacle(s) to producing an antibiogram. (Check all that apply)
$\square$ The laboratory data are difficult to access
$\square$ Limited or no information technology tool for data analysis
$\square$ Limited personnel time for data analysis
$\square$ Limited personnel skills for data analysis
$\square$ Limited interest in an antibiogram from staff who prescribe antibiotics
$\Box$ Our institution does not have enough isolates of any or most species (i.e., < 30 isolates per species) to produce an antibiogram
☐ Other (please specify):
Infection Control Practices
(completed with input from Hospital Epidemiologist and/or Quality Improvement Coordinator)
*12 Number or fraction of infection preventionists (IPs) in facility:
a. Total hours per week performing surveillance:
b. Total hours per week for infection control activities other than surveillance:
*13. Number or fraction of full-time employees (FTEs) for a designated hospital epidemiologist
(or equivalent role) affiliated with your facility:
Continued >>





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	ction Control Practices  upleted with input from Hospital Epidemiologist and/or Quality Improvement Coordinator)
*14.	Is it a policy in your facility that patients infected or colonized with MRSA are routinely placed in contact precautions while these patients are in your facility? (check one)
	$\square$ Yes, all infected or colonized patients
	$\square$ Yes, only all infected patients
	$\square$ Yes, only those with certain characteristics that make them high-risk for transmission (e.g., wounds, diarrhea, presence of an indwelling device)
	$\square$ Yes, only those admitted to high-risk settings (e.g., ICU)
	□ No
	$\square$ Not applicable: my facility never admits these patients
*15.	Is it a policy in your facility that patients infected or colonized with VRE are routinely placed in contact precautions while these patients are in your facility? (check one)
	$\square$ Yes, all infected or colonized patients
	$\square$ Yes, only all infected patients
	$\square$ Yes, only those with certain characteristics that make them high-risk for transmission (e.g., wounds, diarrhea, presence of an indwelling device)
	$\square$ Yes, only those admitted to high-risk settings (e.g., ICU)
	□ No
	$\square$ Not applicable: my facility never admits these patients
*16.	Is it a policy in your facility that patients infected or colonized with CRE (regardless of confirmatory testing for carbapenemase production) are routinely placed in contact precautions while these patients are in your facility? (check one)
	$\square$ Yes, all infected or colonized patients
	$\square$ Yes, only all infected patients
	$\square$ Yes, only those with certain characteristics that make them high-risk for transmission (e.g., wounds, diarrhea, presence of an indwelling device)
	$\square$ Yes, only those admitted to high-risk settings (e.g., ICU)
	□ No
	$\square$ Not applicable: my facility never admits these patients
	Is it a policy in your facility that patients infected or colonized with suspected or confirmed ESBL-producing or extended spectrum cephalosporin resistant Enterobacteriaceae are routinely placed in contact precautions while these patients are in your facility? (check one)
	$\square$ Yes, all infected or colonized patients
	$\square$ Yes, only all infected patients
	$\square$ Yes, only those with certain characteristics that make them high-risk for transmission (e.g., wounds, diarrhea, presence of an indwelling device)
	$\square$ Yes, only those admitted to high-risk settings (e.g., ICU)
	□ No
	☐ Not applicable: my facility never admits these patients  Continued >>





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Infection Control Practices (completed with input from Hospital Epidemiologist and/or Quality Improvement Coordinator)
*18. Does the facility routinely perform screening testing (culture or non-culture) for CRE?
☐ Yes ☐ No
If Yes, in which situations does the facility routinely perform screening testing for CRE? (check all that apply)
$\square$ Surveillance testing at admission for all patients
$\square$ Surveillance testing of epidemiologically-linked patients of newly identified CRE patients (e.g., roommates)
$\square$ Surveillance testing at admission of high-risk patients (e.g., admitted from LTAC or LTCF)
$\square$ Surveillance testing at admission of patients admitted to high-risk settings (e.g. ICU)
☐ Other (please specify):
*19. Does the facility routinely perform screening testing (culture or non-culture) for MRSA?
☐ Yes ☐ No
If yes, in which situations does the facility routinely perform screening testing for MRSA? (check all that apply)
☐ Surveillance testing at admission for all patients
<ul><li>Surveillance testing at admission of high-risk patients (e.g., admitted from LTAC or LTCF)</li></ul>
☐ Surveillance testing at admission of patients admitted to high-risk settings (e.g. ICU)
☐ Surveillance testing of pre-operative patients to prevent surgical site infections
☐ Other (please specify):
*20. Does the facility routinely use chlorhexidine bathing on any patient to prevent infection or transmission of MDROs at your facility? (Note: this does not include the use of such bathing in pre-operative patients to prevent SSIs)
☐ Yes ☐ No
*21. Does the facility routinely use a combination of topical chlorhexidine <u>AND</u> intranasal mupirocin (or equivalent agent) on any patients to prevent infection or transmission of MRSA at your facility? (Note: this does not include the use of these agents in pre-operative surgical patients or dialysis patients)
☐ Yes ☐ No
*22. Among patients with an MDRO admitted to your facility from another healthcare facility, please estimate how often your facility receives information from the transferring facility about the patient's MDRO status?
☐ All the time
$\square$ More than half of the time
$\square$ About half of the time
$\square$ Less than half of the time
$\square$ None of the time
$\square$ Not applicable: my facility does not receive transferred patients with a known MDRO Continued >>
Continued >>





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(completed with input from Physician and Pharmacist Stewardship Champions)
*23. Does your facility have a written statement of support from leadership that supports efforts to improve antibiotic use (antibiotic stewardship)?
☐ Yes ☐ No
*24. Is there a leader responsible for stewardship activities at your facility?
☐ Yes ☐ No
If Yes, what is the position of this leader: (check one)
$\square$ Physician $\square$ Co-led by both Pharmacist and Physician
☐ Pharmacist ☐ Other (please specify):
*25. Is there at least one pharmacist responsible for improving antibiotic use at your facility? $\Box$ Yes $\Box$ No
*26. Does your facility provide any salary support for dedicated time for antibiotic stewardship leadership activities? $\Box$ Yes $\Box$ No
*27. Does your facility have a policy that requires prescribers to document an indication for all antibiotics in the medical record or during order entry?
☐ Yes ☐ No
If Yes, has adherence to the policy to document an indication been monitored? $\square$ Yes $\square$ No
*28. Does your facility have facility-specific treatment recommendations, based on national guidelines and local susceptibility, to assist with antibiotic selection for common clinical conditions?
☐ Yes ☐ No
If Yes, has adherence to facility-specific treatment recommendations been monitored?
☐ Yes ☐ No
*29. Is there a formal procedure for all clinicians to review the appropriateness of all antibiotics at or after 48 hours from the initial orders (e.g. antibiotic time out)?
☐ Yes ☐ No
*30. Do any specified antibiotic agents need to be approved by a physician or pharmacist prior to dispensing at your facility?
☐ Yes ☐ No  Continued >>
Continued





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Antibiotic Stewardship Practices (continued)
*31. Does a physician or pharmacist review courses of therapy for specified antibiotic agents and communicate results with prescribers at your facility?
☐ Yes ☐ No
If Yes, what type of feedback is provided to prescribers? (check all that apply)
$\square$ Feedback on antimicrobial route and/or dosage
$\square$ Feedback on the selection of antimicrobial therapy and/or duration of therapy
$\square$ Other (please specify) :
*32. Does your facility monitor antibiotic use (consumption) at the unit, service, and/or facility wide?  ☐ Yes ☐ No  If Yes, by which metrics? (Check all that apply)  ☐ Days of Therapy (DOT) ☐ Purchasing Data
$\Box$ Defined Daily Dose (DDD) $\Box$ Other (please specify):
☐ Yes ☐ No
*33. Has your facility provided education to clinicians and other relevant staff on improving antibiotic use?
☐ Yes ☐ No