CMS Manual System	Department of Health & Human Services (DHHS)
Pub. 100-07 State Operations Provider Certification	Centers for Medicare & Medicaid Services (CMS)
Transmittal 142	Date: July 17, 2015

SUBJECT: Revisions to State Operations Manual (SOM) Chapter 9 Exhibits

I. SUMMARY OF CHANGES: Revisions to Chapter 9, Exhibit 351. Revisions are being made due to updates and clarification to the Ambulatory Surgical Center (ASC) Infection Control Surveyor Worksheet questions.

NEW/REVISED MATERIAL - EFFECTIVE DATE: July 17, 2015 IMPLEMENTATION DATE: July 17, 2015

The revision date and transmittal number apply to the red italicized material only. Any other material was previously published and remains unchanged. However, if this revision contains a table of contents, you will receive the new/revised information only, and not the entire table of contents.

II. CHANGES IN MANUAL INSTRUCTIONS: (N/A if manual not updated.)
(R = REVISED, N = NEW, D = DELETED) – (Only One Per Row.)

R/N/D	CHAPTER/SECTION/SUBSECTION/TITLE
R	9/Exhibit 351 Ambulatory Surgical Center (ASC) Infection Control Surveyor Worksheet

III. FUNDING: No additional funding will be provided by CMS; contractor activities are to be carried out within their FY 20xx operating budgets.

Funding for implementation activities will be provided to contractors through the regular budget process.

IV. ATTACHMENTS:

	Business Requirements
X	Manual Instruction
	Confidential Requirements
	One-Time Notification
	Recurring Update Notification

Exhibit 351

Ambulatory Surgical Center (ASC) INFECTION CONTROL SURVEYOR WORKSHEET

(Rev: 142, Issued: 07-17-15, Effective: 07-17-15, Implementation: 07-17-15)

Name of State Agency or AO (please specify)

Instructions: The following is a list of items that must be assessed during the on-site survey, in order to determine compliance with the infection control Condition for Coverage. Items are to be assessed primarily by surveyor observation, with interviews used to provide additional confirming evidence of observations. In some cases information gained from interviews may provide sufficient evidence to support a deficiency citation.

The interviews and observations should be performed with the most appropriate staff person(s) for the items of interest (*e.g.*, the staff person responsible for sterilization should answer the sterilization questions). A minimum of one surgical procedure must be observed during the site visit. The surveyor(s) must identify at least one patient and follow that case from registration to discharge to observe pertinent practices. For facilities that perform brief procedures, e.g., colonoscopies, it is preferable to follow at least two cases. When performing interviews and observations, any single instance of a breach in infection control would constitute a breach for that practice.

Citation instructions are provided throughout this instrument, indicating the applicable regulatory provision to be cited on the Form CMS-2567 when deficient practices are observed.

PART 1 - ASC CHARACTERISTI	cs	
1. ASC Name		
2. Address, State and Zip Code	Address	
3. 10-digit CMS Certification Numb	City State Zip	
4. What year did the ASC open operation?	for	
5. Please list date(s) of site visit: m m	d d y y y Y m m d d y y	у у
6. What was the date of the most recent previous federal (CMS) surv	vey: m m d d y y y y	
7. Does the ASC participate in Med	dicare via accredited "deemed" status? O YES O NO	
7a. If YES, by which CMS-recognized accreditation organization(s)?	Accreditation Association for Ambulatory Health Care (AAAHC) American Associate for Accred. of Ambulatory Surgery Facilities (AAAAS American Osteopathic Association (AOA) The Joint Commission (TJC)	SF)

	7b. If YES, according to the ASC, what was the date of the most recent accreditation survey?		m	m] / [d	d	/	у	у у	у	
	What is the ownership of the facility? LECT only ONE bubble)	О H О N	hysician- ospital-c ational c ther (ple	wneo orpo	d ration (iı	ncluding	g joint v	enture:	s with p	hysicia	ns)
the ma	What is the primary procedure perfor ASC (i.e., what procedure type reflectionity of procedures performed at the lect only ONE bubble)	ts the		ASC.	What ad? (Select not inclustion 9.	all that	tapply)	·		t the
000000000	Dental Endoscopy Ear/Nose/Throat OB/Gyn Ophthalmologic Orthopedic Pain Plastic/reconstructive Podiatry Other (please specify):			0000000000	OB/Gyr Ophtha Orthop Pain Plastic/ Podiatr	se/Thro n nImologi edic reconst	ructive				
pro	cedures on?	O A	ediatric p dult pation	ents c	only	t patien	ts				
pro	What is the average number of cedures performed at the ASC per nth ?									peri	month
	How many Operating Rooms (includincedure rooms) does the ASC have?	ng	0	O 2	O 3	O 4	O 5	O 6	O 7	0	O 9+
Nu	mber actively maintained:		0	O 2	O 3	O 4	O 5	O 6	O 7	O 8	O 9+

14. Please indicate how the follow	ing services a	re provided: (fill in all that ap	ply)
	Contract	Employee	Other	If Other, Please print:
Anesthesia/Analgesia	0	0	0	
Environmental Cleaning	0	0	0	
Linen	0	0	0	
Nursing	0	0	0	
Pharmacy	0	0	0	
Sterilization/Reprocessing	0	0	0	
Waste Management	0	0	0	
15. Does the ASC have an explicit i				O YES O NO
NOTE! If the ASC does not have an CFR 416.51 must be cited.	explicit infec	tion control pr	ogram, a condit	on-level deficiency related to 42
16. Does the ASC's infection contriguidelines?	ol program fo	llow nationally	recognized infe	ction control O YES O NO
NOTE! If the ASC does not follow r CFR 416.51(b) must be cited. Depe condition-level citation may also b	ending on the	scope of the la	_	· · · · · · · · · · · · · · · · · · ·
16a. Is there documentation the recognized infection control grown NOTE! If the ASC cannot document for use in its infection control programmer in the case even if the generally accepted standards of prany nationally recognized guideling control standards of practice, there deficiency related to 42 CFR 416.5	t that it considerant, a deficiency of the ASC's infect ractice/nation es nor complication the ASC should be a sho	ts program? dered and selectory related to ion control praid guidelines. It is with general	cted specific gui 42 CFR 416.51(lectices comply well f the ASC neither lly accepted info	O YES O NO delines o) must rith or selected ection

16b. If YES to (a), which	0	CDC/HICI	PAC Guidelines:
nationally-recognized		•	Guideline for Isolation Precautions (CDC/HICPAC)
infection control guidelines has the ASC selected for its		O F	land hygiene (CDC/HICPAC)
program?			Disinfection and Sterilization in Healthcare Facilities (CDC/HICPAC)
(Select all that apply)			nvironmental Infection Control in Healthcare Facilities
			CDC/HICPAC)
	0	Periopera	ative Standards and Recommended Practices (AORN)
	0	Guideline	es issued by a specialty surgical society / organization (List)
		Please s	pecify (please limit to the space provided):
	0	Others	
		Please s	pecify (please limit to the space provided):
47.0 11.4601 11			
		-	ofessional qualified through training O YES ASC's infection control program? O NO
			- O NO
			esignated a qualified professional with training (not necessarily ection control program, a deficiency related to 42 CFR
The state of the s			professional responsible for infection control should be
considered for citation of a cond	<mark>ition-</mark>	<mark>level defic</mark>	ciency related to 42 CFR 416.51.
17a. If YES, Is this person an:			O ASC employee
(Select only ONE bubble)			O ASC contractor
17b. Is this person certified in			
(Note: §416.50(b)(1) does infection control.)	<u>not</u> r	equire tha	t the individual be certified in ONO
17c. If this person is NOT cert	ified	in	Ţ
infection control, what typ			
control training has this pe	erson	received?	
17d. On average, how many h		•	
does this person spend in directing the infection con			hours per week
	•		nt of time the person must spend in the ASC directing the
			t the designated individual spends sufficient time on-site
directing the program, taking int	<mark>o cor</mark>	sideration	the size of the ASC and the volume of its surgical activity.)

18. Does the ASC have a system to active related to procedures performed at the NOTE! If the ASC does not have a document to 42 CFR 416.51(b)(3) must be	ASC? nente	e <mark>d identification system, a deficien</mark>		YES NO
				Profession
18a. If YES, how does the ASC obtain this information?	0	The ASC sends e-mails to patie		_
(Select ALL that apply)	0	The ASC follows-up with their after discharge	patients'	primary care providers
	0	The ASC relies on the physicial obtain this information at a for report it to the ASC	•	•
	0	Other (please specify):		
18b. Is there supporting documenta	ation	confirming this tracking activity?	0	YES
			0	NO
NOTE! If the ASC does not have support	ing d	ocumentation, a deficiency related	<mark>d <i>to</i> 42 CF</mark>	R 416.51(b)(3) must be
<mark>cited.</mark>				
18c. Does the ASC have a policy/pronotifiable disease reporting require			0	YES NO
NOTE! If the ASC does not have a report CMS does not specify the means for rep	_	•		
19. Do staff members receive infection	contr	ol training?	0	YES
If training is completely absent, then co				NO
level citation in relation to 42 CFR 416.5 to comply with infection control standa		The state of the s	s tall	
19a. If YES, how do they receive	0	In-service		
infection control training?	0	Computer-based training		
(Select all that apply)	0	Other (please specify):		
	0	Medical staff		
19b. Which staff members receive	0	Nursing staff		
infection control training? (Select all that apply)	0	Other staff providing direct patie	nt care	
(0.000000000000000000000000000000000000	0	Staff responsible for on-site steril	lization/h	gh-level disinfection
	0	Cleaning staff		
	0	Other (please specify):		

19c. Is training:	0	the same for all ca	· ·		
19d. Indicate frequency of staff infection control training (Select all that apply)	0 0 0	Upon hire Annually Periodically / as n Other (please spe	Г		
19e. Is there documentation conficategories of staff listed above? NOTE! If training is not provided to aptraining thereafter, a deficiency must	<mark>propri</mark>	ate staff upon hire,	granting of		e refresher
20. How many procedures were observed during the site visit?	O 1	O 2	O 3	O 4	O Other
If other, please specify the nu	mber:			procedures	

PART 2 - INFECTION CONTROL & RELATED PRACTICES

INSTRUCTIONS:

- Please select ONE bubble for each "Was Practice Performed?" question, unless otherwise noted.
- If N/A or unable to observe is selected as the response, please explain why there is no associated observation, or why the question is not applicable, in the surveyor notes box. Surveyors should attempt to assess the practice by interview or document review if unable to observe the actual practice during survey.
- During the survey, observations or concerns may prompt the surveyor to request and review specific policies and procedures. Surveyors are expected to use their judgment and review only those documents necessary to investigate their concern(s) or to validate their observations.

I. Hand Hygiene

Observations are to focus on staff directly involved in patient care (e.g., physicians, nurses, CRNAs, etc.). Hand hygiene should be observed not only during the case being followed, but also while making other observations in the ASC throughout the survey.

Unless otherwise indicated, a "No" response to any question below **must** be cited as a deficient practice in relation to 42 CFR 416.51(a).

Practices to be Assessed		s Practice formed?	Surveyor Notes:		
A. All patient care areas have <i>readily accessible, in</i>	. All patient care areas have readily accessible, in appropriate locations:				
a. Soap and water	0	Yes			
	0	No			
b. Alcohol-based hand rubs	0	Yes			
	0	No			
I. If alcohol-based hand rub is available	0	Yes			
in patient care areas, it is installed as required. (There are LSC requirements at 42 CFR 416.44(b)(5) for installation of alcohol-based hand rubs)	0	No			
B. Staff perform hand hygiene:					
a. After removing gloves	0	Yes No			
b. Before direct patient contact	0	Yes No			
c. After direct patient contact	0	Yes No			
d. Before performing invasive procedures (e.g. placing an IV)	0 0	Yes No <i>Unable to observe</i>			

A. Needles are used for only one patient.	0 0	Yes No <i>Unable to observe</i>
Practices to be Assessed	Per	s Practice Surveyor Notes formed?
	nsmi that to ta	ssion that warrant engagement of public health a survey has identified evidence of one or more of the king appropriate enforcement action to ensure the
(e.g., anesthesiologists, certified registered nurs Unless otherwise indicated, a "No" response to a relation to 42 CFR 416.51(a).	and ase and	edministering medications and performing injections esthetists, nurses). uestion below must be cited as a deficient practice in arveyor notes box why it was not observed and attempt
D. Personnel providing direct patient care do not wear artificial fingernails and/ or extenders when having direct contact with patients.	0	Yes No
c. Remove gloves before moving to the next tasks and/or patient	0	Yes No Unable to observe
b. Wear gloves when handling potentially contaminated patient equipment	0 0 0	Yes No <i>Unable to observe</i>
C. Regarding gloves, staff: a. Wear gloves for procedures that might involve contact with blood or body fluids	000	Yes No <i>Unable to observe</i>
 e. After contact with blood, body fluids, or contaminated surfaces (even if gloves are worn) 	0 0 0	Yes No <i>Unable to observe</i>

Practices to be Assessed		Practice ormed?	Surveyor Notes
B. Syringes are used for only one patient (this	0	Yes	
includes manufactured prefilled syringes).	0	No	
	0	Unable to observe	
C. The rubber septum on a medication, whether	0	Yes	
unopened or previously accessed, vial is disinfected	0	No	
with alcohol prior to piercing.		Unable to observe	
D. Medication vials are always entered with a new	0	Yes	
needle.		No	
		Unable to observe	
E. Medication vials are always entered with a new	0	Yes	
syringe ,	Ö	No	
-,6		Unable to observe	
F. Medications that are pre-drawn are labeled	0	Yes	
with the date and time of draw, initials of the	0	No	
person drawing, medication name, strength and	0	Unable to observe	
beyond-use date and time	Ū	Chable to observe	
NOTEe: A "No" answer should result in citation as	<mark>a defi</mark>	<mark>cient practice in rela</mark>	ation to 42 CFR 416.48(a),
Administration of Drugs			
G. a. Single dose (single-use) medication vials	0	Yes	
are used for only one patient	0	No	
	0	Unable to observe	
b. Bags of IV solutions are used for only one	0	Yes	
patient (and not as a source of flush solution	0	No	
for multiple patients).	0	Unable to observe	
c. Medication administration tubing and	0	Yes	
connectors are used for only one patient	0	No	
, ,			

Practices to be Assessed		s Practice formed?	Surveyor Notes
H. The ASC has voluntarily adopted a policy that medications labeled for multi-dose use for multiple patients are nevertheless only used for one patient. (Fill in N/A if no multi-dose medications/infusates	O O O s are u	Yes No N/A sed).	
(NOTE: a "No" answer to question H. does not incresult in a citation. <i>However</i> , a "No" response to cited). If YES, please skip to "K" If NO, you <i>must also</i> assess the practices <i>at ques</i>	<mark>either</mark>	or both of the	the contract of the contract o
I. Multi-dose vials are dated when they are first opened and discarded within 28 days unless the manufacturer specifies a different (shorter or longer) date for that opened vial. Note: This is different from the expiration date for the vial. The multi-dose vial can be dated with either the date opened or the <i>beyond-use date</i> as per ASC policies and procedures, so long as it is clear what the date represents and the same policy is used consistently throughout the ASC.	0 0 0	Yes No <i>Unable to obs</i>	erve
J. Multi-dose medication vials used for more than one patient are stored appropriately and do not enter the immediate patient care area (e.g., operating room, anesthesia carts). NOTE: If multi-dose vials enter the immediate patient care area, they must be dedicated for single patient use and discarded immediately after use.	0 0 0	Yes No <i>Unable to obse</i>	erve
K. All sharps are disposed of in a puncture- resistant sharps container	0	Yes No	
L. Sharps containers are replaced when the fill line is reached	0 0	Yes No	

III. Single Use Devices, Sterilization, and High Level Disinfection

Pre-cleaning must always be performed prior to sterilization and high-level disinfection

Sterilization must be performed for critical equipment (i.e., instruments and equipment that enter normally sterile tissue or the vascular system, such as surgical instruments)

High-level disinfection must be performed for semi-critical equipment (i.e., items that come into contact with non-intact skin or mucous membranes such as reusable flexible endoscopes, laryngoscope blades)

Observations are to be made of staff *performing* equipment reprocessing (e.g., surgical techs), unless these activities are performed under contract or arrangement off-site from the ASC.

Unless otherwise indicated, a "No" response to any question below **must** be cited as a deficient practice in relation to 42 CFR 416.51(a).

SINGLE-USE DEVICES

(Choose N/A if single-use devices are never reprocessed and used again) (Surveyor to confirm there is a contract or other documentation of an arrangement with a reprocessing facility by viewing it)

Practices to be Assessed			Was Practice Surveyor Notes Performed?			
Α.	a. If single-use devices are rep		0	Yes		
	devices that are approved by t	the FDA for	0	No		
	reprocessing		0	N/A		
	b. If single-use devices are rep	rocessed, they are	0	Yes		
	reprocessed by an FDA-approv	ved reprocessor.	0	No		
			0	N/A		
		STERIL	IZATIOI	N		
A. (Critical equipment is sterilized		0	Yes		
_			0	No		
В.	Are sterilization procedures perf	formed on-site?	0	Yes		
(If I	NO, skip to "F")		0	No		
<mark>per</mark>	"No" answer does not result in a mitted to provide for sterilization tractual arrangement.)		<mark>are</mark>			
doc	rveyor to confirm there is a cont cumentation of an arrangement viewing it)		on			
/	a. If YES to B, please indicate	O Steam autocla	ve			
	method of sterilization:	O Peracetic acid				
		O Other (please				
		specify):				

C. Items are pre-cleaned according to manufacturer's instructions or, if the manufacturer does not provide instructions, evidence-based guidelines prior to sterilization		Was Practice Surveyor Notes Performed?		
		000	Yes No <i>Unable to observe</i>	?
inspe	edical devices and instruments are visually cted for residual soil and re-cleaned as ed before packaging and sterilization	000	Yes No <i>Unable to observe</i>	;
correc	hemical indicator (process indicator) is placed ctly, as described in manufacturer's ctions for use, in the instrument packs in load.	0 0 0	Yes No Unable to observe	,
each s impla	iological indicator is <i>used</i> at least weekly <i>for</i> sterilizer and with every load containing ntable items, as evidenced by ASC mentation (i.e., log).	000	Yes No <i>Unable to observe</i>	?
	h load is monitored with mechanical tors (e.g. time, temperature, pressure)	000	Yes No <i>Unable to observe</i>	•
equip	cumentation for each piece of sterilization ment is maintained and up to date and les results from each load	0	Yes No	
	opropriately contained and handled during on process to assure that sterility is not prior to use	0 0 0	Yes No <i>Unable to observe</i>	•
	zation, medical devices and instruments are signated clean area so that sterility is not	0	Yes No	
G. Sterile pack compromised are reprocesse		0	Yes No	

Practices to be Assessed	Was Practice Performed?	Surveyor Notes
H. Is immediate-use steam sterilization (IUSS) performed on-site? If NO, skip to "High Level Disinfection Section"	O Yes O No	
If YES, you must also assess the practices at questions "I - K": (A "No" answer does not result in a citation)		
 Work practices ensure proper cleaning and decontamination, inspection, and arrangement of the instruments into the recommended sterilizing trays or other containment devices before sterilization. Once clean, the item is placed within a container intended for immediate use. The sterilizer cycle and parameters used are selected according to the manufacturers' instructions for use for the device, container, and sterilizer. The sterilizer function is monitored with monitors (e.g., mechanical, chemical and biologic) that are approved for the cycle being used. The processed item must be transferred immediately, using aseptic technique, from the sterilizer to the actual point of use, the sterile field in an ongoing surgical procedure. 	O Yes O No O Unable to observe O N/A	
Note: "Immediate use" is defined as the shortest possible time between a sterilized item's removal from the sterilizer and its aseptic transfer to the sterile field. A sterilized item intended for immediate use is not stored for future use, nor held from one case to another. IUSS is not equivalent to "short cycle" sterilization performed in accordance with manufacturers' IFUs. IUSS must not be a routine or frequent practice in the ASC.		

Practices to be Assessed	Was Practice Performed?	Surveyor Notes
 J. Immediate-use steam sterilization is NOT performed on the following devices: Implants. Post-procedure decontamination of instruments used on patients who may have Creutzfeldt-Jakob disease or similar disorders. Devices that have not been validated with the specific cycle employed. Single-use devices that are sold sterile. 	O Yes O No	
K. Is IUSS performed on a routine basis?	O Yes	
(A "Yes" answer must be cited as a deficient practice in relation to 42 CFR 416.51(a).	O No	
HIGH-LEVEL DI	SINFECTION	
Practices to be Assessed	Was Practice Performed?	Surveyor Notes
A. Semi-critical equipment is high-level disinfected or sterilized	O Yes O No O N/A	
B. Is high-level disinfection performed on site? (If NO, Skip to "F")	O Yes O No O N/A	
(A "No" answer does not result in a citation, since ASCs ar site, under a contractual arrangement.)	e permitted to prov	vide for high-level disinfection off-
(Surveyor to confirm there is a contract or other documer viewing it)	ntation of an arrang	ement for off-site sterilization by
a. If answer to B was YES , please O Manu	al	
indicate method of high-level O Auton disinfection:	nated	
O Other (please specify):		
C. Items are pre-cleaned according to manufacturer's instructions or, if the manufacturer does not provide instructions, evidence-based guidelines prior to high-level disinfection	O Yes O No O <i>Unable to obse</i>	erve

Practices to be Assessed		Was Practice Surveyor Notes Performed?		Surveyor Notes
D.	a. Medical devices and instruments are visually	0	Yes	
	inspected for residual soil and re-cleaned as needed	0	No	
	before high-level disinfection	0	Unable to observe	
	b. High-level disinfection equipment is maintained	0	Yes	
	according to manufacturer instructions	0	No	
		0	Unable to observe	
	c. Chemicals used for high-level disinfection are:			
	I. Prepared according to manufacturer	0	Yes	
	instructions	0	No	
		0	Unable to observe	
	II. Tooked for a representation according			
	II. Tested for appropriate concentration according to manufacturer's instructions	_	Yes	
	to manufacturer's mistructions	0	No Unable to observe	
		0	Unable to observe	
	III. Replaced according to manufacturer's	0	Yes	
	instructions	0	No	
		0	Unable to observe	
	IV. Documented to have been prepared and	0	Yes	
	replaced according to manufacturer's instructions	0	No	
	d. Instruments requiring high-level disinfection are:			
	I. Disinfected for the appropriate length of time	0	Yes	
	as specified by manufacturer's instructions or, if	0	No	
	the manufacturer does not provide instructions,	0	Unable to observe	
	evidence-based guidelines			
	II. Disinfected at the appropriate temperature as	0	Yes	
	specified by manufacturer's instructions or, if the		No	
	manufacturer does not provide instructions,	Ö	Unable to observe	,
	evidence-based guidelines	Ü	Onable to observe	
E. It	ems that undergo high-level disinfection are allowed	0	Yes	
	ry before use	Ō	No	
		0	Unable to observe	
F. F.	ollowing high-level disinfection, items are placed in a	0	Yes	
	gnated clean area in a manner to prevent contamination	•	No	
5.551		•		

IV. Environmental Infection Control

Observations are to be made of staff performing environmental cleaning (e.g., surgical technicians, cleaning staff, etc.)

If unable to observe is selected, please clarify in the surveyor notes box why it was not observed and attempt to assess by means of interview or documentation review.

Unless otherwise indicated, a "No" response to any question below **must** be cited as a deficient practice in relation to 42 CFR 416.51(a).

relation to 42 CFR 416.51(a).	
Practices to be Assessed	Was Practice Surveyor Notes Performed?
A. Operating rooms are cleaned and disinfected after each surgical or invasive procedure with an EPA-registered disinfectant	O Yes O No O Unable to observe
B. Operating rooms are terminally cleaned daily	O Yes O No O Unable to observe
C. Environmental surfaces in patient care areas are cleaned and disinfected, using an EPA-registered disinfectant on a regular basis (e.g., daily), when spills occur and when surfaces are visibly contaminated.	O Yes O No O Unable to observe
D. The ASC has a procedure in place to decontaminate gross spills of blood.	O Yes O No
V. Point of Care Devices (e.g., blood glucose meter) Observations are to be made of staff performing fingerstic If unable to observe or N/A is selected, please clarify in the sapplicable and attempt to assess by means of interview or during the same of the	urveyor notes box why it was not observed or ocumentation review.
Practices to be Assessed	Was Practice Performed? Surveyor Notes
1. Does the ASC <i>use</i> a point-of-care <i>testing</i> device, such as a blood glucose meter? If NO, STOP HERE.	O Yes O No

Practices to be Assessed		s Practice formed?	Surveyor Notes
A. Hand hygiene is performed before and after performing a	0	Yes	
finger stick procedure to obtain a sample of blood and using the point-of-care testing device.	0	No	
B. Gloves are worn by health care personnel when	0	Yes	
performing a finger stick procedure to obtain a sample of blood, and are removed after the procedure (followed by hand hygiene).	0	No	
C. Finger stick devices are not used for more than one	0	Yes	
patient.	0	No	
	0	Unable to observe	
NOTE: This includes both the lancet and the lancet holding device.			
D. If used for more than one patient, the point-of-care	0	Yes	
testing device (e.g., blood glucose meter, INR monitor) is	0	No	
cleaned and disinfected after every use according to the manufacturer's instructions.	0	N/A	
NOTE: if the manufacturer does not provide instructions for cleaning and disinfection, then the device should not be used for >1 patient.	d		