Exhibit 351 Ambulatory Surgical Center INFECTION CONTROL SURVEYOR WORKSHEET (Rev. 68 Issued: 11-24-10, Effective: 11-24-10, Implementation: 11-24-10)

Name of State Agency or AO (please print at right)

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, unless the ASC is a low volume ASC with no procedures scheduled 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 CHARACTERIS	FICS
1. ASC Name (<i>please print</i>)	
2. Address, State and Zip Code <i>(please print)</i>	Address
	City State
3. 10-digit CMS Certification Number	
4. What year did the ASC open for operation?	y y y y
5. Please list date(s) of site	/ to / /
visit: m m d d	yyyy mmdd yyyy
6. What was the date of the most recent previous federal (CMS) survey:	m m d d y y y y
	FILL IN EACH BUBBLE USING A DARK PEN.
7. Does the ASC participate in Medicare status?	via accredited "deemed" O YES O NO

7a. If YES, by w CMS-recognized accreditation organization? (Check only ON)		O Ame AAAASI O Ame	editation A rican Asso ⁽⁷⁾ rican Ostee Joint Comi	ciate fo opathic	or Accr Assoc	ed. of	Ambı	ulator				
7b. If YES, accord ASC, what was t most recent accre	he date o		m	m	/d	d	/	у	у у	/ y		
8. What is the owner	ship of th	ne O	Physicia	n-owne	d							
facility?		0	Hospital	-owned								
		0	National	corpor	ation (i	includ	ing jo	int ve	ntures	with p	ohysic	ians)
		0	Other (p)	lease p	rint):							
9. What is the primar the ASC (i.e., what p majority of procedur (Fill in only ONE b	procedure es perfor	type refle	ects the	ASC Do 1	What a C? (Fill not inc stion 9.	l in al lude t	l that	apply	7)			
O Dental				0	Denta	ıl						
O Endoscopy				0	Endo	scopy						
O Ear/Nose/Throa	at			0	Ear/N	lose/T	hroat					
O OB/Gyn				0	OB/C	iyn						
O Ophthalmologi	с			0	Ophtl	nalmo	logic					
O Orthopedic				0	Ortho	pedic						
O Pain				0	Pain							
O Plastic/reconstr	uctive			0	Plasti		nstruc	ctive				
O Podiatry	г			_ 0	Podia	•						
O Other (please p	rint):			0	Other	· (plea	se prii	nt):				
11. Who does the AS procedures on? (<i>Fill in only</i> ONE b)	-	m O O O	Pediatric Adult pa Both ped	tients o	only		ents					
12. What is the avera	-											
procedures performe per month ?	d at the A	ASC							<i>p</i>	er moi	nth	
13. How many Operation	ating Roo	oms (inclu	ding	0	0	0	0	0	0	0	0	0
procedure rooms) do	es the AS	SC have?	-	1	2	3	4	5	6	7	8	9+
Number actively mail	intained:			0	0 2	O 3	0 4	0 5	0 6	0 7	0 8	0 9+
14. Please indicate h	ow the fo	ollowing s	ervices are	provid						-	-	·
			t Employ	-	Othe				ther, P	lease	print:	
Anesthesia		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				
INFECTION CONTROL	L PRO	GRAM	ſ					
15. Does the ASC have an	explic	it infec	tion control pr	ogram?		0 0	YES NO	
NOTE! If the ASC does no related to 42 CFR 416.51				control program	n, a conditio	on-le	evel deficiency	
16. Does the ASC's infection infection control guideline		ntrol pro	ogram follow r	nationally recog	gnized	0 0	YES NO	
NOTE! If the ASC does not follow nationally recognized infection control guidelines, a deficiency related to 42 CFR 416.51(b) must be cited. Depending on the scope of the lack of compliance with national guidelines, a condition-level citation may also be appropriate.								
16a. Is there document nationally-recognized i						0 0	YES NO	
nationally-recognized infection control guidelines has the ASC selected for its program? (<i>Fill in all that apply</i>)	O P	O O H O I O E Cerioper Guidelin	Hand hygiene (Disinfection an Environmental ative Standard les issued by a	solation Precau CDC/HICPAC d Sterilization i Infection Contr s and Recomme) in Healthcar rol in Health ended Practi cal <i>society</i> /	re Fancaro ices <i>org</i>	acilities e Facilities (AORN) <i>canization (List)</i>	
	0 0	thers <i>Please</i>	specify (pleas	e print and limi	t to the space	ce pi	rovided):	
		Γ	V + 1A	-				

NOTE! If the ASC cannot document that it considered and selected specific guidelines for use in its infection control program, a deficiency related to 42 CFR 416.51(b) **must** be cited. This is the case even if the ASC's infection control practices comply with generally accepted standards of practice/national guidelines. If the ASC neither selected any nationally recognized guidelines nor complies with generally accepted infection control standards of practice, then the ASC should be cited for a condition-level deficiency related to 42 CFR 416.51.

17. Does the ASC have a licensed health care professional qualified through	0	YES
training		
in infection control and designated to direct the ASC's infection control	0	NO
program?		

NOTE! If the ASC cannot document that it has designated a qualified professional with training (not necessarily certification) in infection control to direct its infection control program, a deficiency related to 42 CFR 416.51(b)(1) **must** be cited. Lack of a designated professional responsible for infection control should be considered for citation of a condition-level deficiency related to 42 CFR 416.51.

17a. If YES, Is this person as (<i>Fill in only ONE bubble</i>)	n:	0 0	ASC employee ASC contractor
L	n infection control (i.e., CIC) (Note: ire that the individual be certified in	0 0	YES NO
17c. If this person is NOT configuration infection control, what type control training has this pers	of infection		
the infection control program, but it	in the ASC hours per	end in l spend	ls sufficient time on-
18. Does the ASC have a system to been related to procedures performe	actively identify infections that may have d at the ASC?	0	YES NO
18a. If YES, how does the ASC obtain this information? (<i>Fill in ALL that apply</i>)	 O The ASC sends e-mails to patients a O The ASC follows-up with their patiafter discharge O The ASC relies on the physician period obtain this information at a follow-report it to the ASC O Other (please <i>print</i>): 	ents' pr	rimary care providers

18b. Is there supporting documentation confirming this tracking activity?

O YESO NO

NOTE! If the ASC does not have an and 42 CFR 416.51(b)(3) must be c	identification system, a deficiency related to 42 CFR 416.44(a)(3) ited.
18c. Does the ASC have a polic notifiable disease reporting requ	y/procedure in place to comply with State O YES irements? O NO
	reporting system, a deficiency must be cited related to 42 CFR the means for reporting; generally this would be done by the State
19. Do staff members receive infect	ion control training? O NO
19a. If YES, how do they receive infection control training? (<i>Fill in all that apply</i>)	 O In-service O Computer-based training O Other (<i>please</i> print):
19b. Which staff members receive infection control training? (Fill in all that apply)	 Medical staff Nursing staff Other staff providing direct patient care Staff responsible for on-site sterilization/high-level disinfection Cleaning staff Other (please print):
19c. Is training:	O the same for all categories of staffO different for different categories of staff
19d. Indicate frequency of staff infection control training (Fill in all that apply)	 O Upon hire O Annually O Periodically / as needed O Other (please print):
19e. Is there documentation con categories of staff listed above?	firming that training is provided to all OYES ONO

NOTE! If training is not provided to appropriate staff upon hire/granting of privileges, with some refresher training thereafter, a deficiency **must** by cited in relation to 42 CFR 416.51(b) and (b)(3). If training is completely absent, then consideration should be given to condition-level citation in relation to 42 CFR 416.51, particularly when the ASC's practices fail to comply with infection control standards of practice.

20. How many procedures were observed during the site visit?	O	0	O	0	O
	1	2	3	4	Other
If other, please <i>print</i> the number:			procedures		

PART 2 – INFECTION CONTROL & RELATED PRACTICES

INSTRUCTIONS:

- Please **completely fill in ONE bubble** for each "Was Practice Performed?" and "Manner of Confirmation" question, unless otherwise noted.
- Please use a dark pen to fully fill in each bubble.
- 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).
- If N/A is response, please explain why there is no associated observation, or why the question is not applicable, *in the COMMENTS box at the end of each section*.

I. Hand Hygiene

Observations are to focus on staff directly involved in patient are (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. Interviews are used primarily to provide additional evidence for what the surveyor has observed, but may in some cases substitute for direct observation to support a citation of deficient practice.

Practices to be Assessed	Practico		Manner of Confirmation	
A. All patient care areas have: Note: 42 CFR 416.51(a) should be cited only if the answer to both a ar	nd b	is "No."		
a. Soap and water available	0 0	Yes No	0 0 0	Observation Interview Both
b. Alcohol-based hand rubs available	000	Yes No	000	Observation Interview Both
I. If alcohol-based hand rub is available 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)		Yes No		
B. Staff perform hand hygiene:				
a. After removing gloves	0 0 0	Yes No N/A	0000	Observation Interview Both
b. After direct patient contact	0 0 0	Yes No N/A	0 0 0	Observation Interview Both
c. Before performing invasive procedures (e.g., placing an IV)	0 0	Yes No	0 0	Observation Interview

	0	N/A	0	Both
Practices to be Assessed		as actice rformed?		nner of nfirmation
d. After contact with blood, body fluids, or contaminated surfaces (even if gloves are worn)	000	Yes No N/A	0 0 0	Observation Interview Both
C. Regarding gloves, staff:				
a. Wear gloves for procedures that might involve contact with blood or body fluids	0 0 0	Yes No N/A	0 0 0	Observation Interview Both
b. Wear gloves when handling potentially contaminated patient equipment	0 0 0	Yes No N/A	0 0 0	Observation Interview Both
c. Remove gloves before moving to the next tasks and/or patien	0 t 0 0	Yes No N/A	0 0 0	Observation Interview Both
D. Additional breaches in hand hygiene, not captured by the questions above, were identified (If YES, please specify further in comments)	0 0 0	Yes No N/A	0 0 0	Observation Interview Both
Comments: (please print and limit comments to the space provided)				
II. Injection Practices (injectable medications, saline, other infus Observations are to be made of staff who prepare and administer injections (e.g., anesthesiologists, certified registered nurse anesthe	med	ications a	-	perform
Practices to be Assessed		as actice rformed?		nner of nfirmation
A. Needles are used for only one patient	000	Yes No N/A	000	Observation Interview Both
B. Syringes are used for only one patient	0 0 0	Yes No N/A	0 0 0	Observation Interview Both

Practices to be Assessed		s ctice formed?		nner of Ifirmation
C. Medication vials are always entered with a new needle	0	Yes No	0	Observation Interview
	0	N/A	0	Both
	0	Yes	0	Observation
, , , , , , , , , , , , , , , , , , ,	0	No	0	Interview
	0	N/A	0	Both
	0	Yes	0	Observation
initials of the person drawing, medication name, strength and expiration	0	No	0	Interview
date or time	0	N/A	0	Both
Note: A "No" answer should result in citation as a deficient practice in r Administration of Drugs	relat	ion to 42	CFR	. 416.48(a),
F. a. Single dose (single-use) medication vials are used for only	0	Yes	0	Observation
one patient (A "No" response must be cited in relation to 42	0	No	0	Interview
CFR 416.48(a).)	0	N/A	0	Both
h Manufactured genefilled avaia and an used for only and	0	Yes	0	Observation
b. Manufactured prefilled syringes are used for only one notiont	0	No	0	Interview
patient	0	N/A	0	Both
	0	Yes	0	Observation
c. Bags of IV solutions are used for only one patient	0	No	0	Interview
	0	N/A	0	Both
	0	Yes	0	Observation
d. Medication administration tubing and connectors are used	0	No	0	Interview
for only one patient	0	N/A	0	Both

G. *Please print* all injectable medications/infusates that are in a vial/container used for **more than one patient:**

Name of Medication	Average number of patients per vial/container

Practices to be Assessed	Was Practice Performed?	Manner of Confirmation		
H. Multi-dose injectable medications are used for only one patient	O Yes O No O N/A	O ObservationO InterviewO Both		

(Note: a "No" answer here is not necessarily a breach in infection control and does not result in a citation. However, a "No" response to the related questions I - K should be cited).

(Fill in N/A if no multi-dose medications/infusates are used).

If YES, please skip to "L"

If NO, please answer "I-K":

I. The rubber septum on a multi-dose vial used for more than one patient is disinfected with alcohol prior to each entry				0 0 0	Observation Interview Both
J. Multi-dose medication when they are first ope opening or according to comes first	0 0 0	Yes No N/A	0 0 0	Observation Interview Both	
K. Multi-dose medications, used for more than one patient , are not stored or accessed in the immediate areas where direct patient contact occurs				0 0 0	Observation Interview Both
L. All sharps are disposed of in a puncture-resistant sharps container			Yes No N/A	0 0 0	Observation Interview Both
M. Sharps containers are replaced when the fill line is reached			Yes No N/A	000	Observation Interview Both
N. Additional breaches in injection practices, not captured by the questions above were identified (If YES, please specify further in comments)			Yes No N/A	0 0 0	Observation Interview Both
Comments: (please print and limit comments to the space provided)					

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 who perform equipment reprocessing (e.g., surgical techs), unless these activities are performed under contract or arrangement off-site from the ASC.

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 Performed?		Manner of Confirmation		
A. a. If single-use devices are reprocessed, they are devices that a approved by the FDA for reprocessing					000000000000000000000000000000000000000	Yes No N/A	0 0 0	Observation Interview Both	
b. If single-use devices are reprocessed, they are reprocessed b an FDA-approved reprocessor.					000	Yes No N/A	000	Observation Interview Both	
			STERILIZATI	ON					
A. Critical equipment is sterilized					0 0 0	Yes No N/A	0 0 0	Observation Interview Both	
B. Are sterilization procedures performed on-site? (If NO, skip to "F")					000	Yes No N/A	000	Observation Interview Both	
(A "No" answer does not result in a citation, since ASCs are permitted to provide for sterilization off-site, under a contractual arrangement.)					-		-		
	eyor to confirm there is a c gement for off-site steriliza			ation of an					
	a. If YES to B , please indicate method of sterilization:	0 0 0	Steam autoclave Peracetic acid Other (<i>please</i>						
C. Items are pre-cleaned according to manufacturer's instructions or evidence-based guidelines prior to sterilization				actions or	0 0	Yes No	0 0	Observation Interview	

	0	N/A	0	Both	
Practices to be Assessed		Was Practice Performed?		Manner of Confirmation	
D.	0	Yes	0	Observation	
a. Medical devices and instruments are visually inspected for	0	No	0	Interview	
residual soil and re-cleaned as needed before packaging and sterilization	0	N/A	0	Both	
	0	Yes	0	Observation	
b. A chemical indicator is placed in each load	Ο	No	0	Interview	
	0	N/A	0	Both	
	0	Yes	0	Observation	
c. A biologic indicator is performed at least weekly and with all	0	No	0	Interview	
implantable loads	0	N/A	0	Both	
	0	Yes	0	Observation	
d. Each load is monitored with mechanical indicators (e.g. time,	0	No	0	Interview	
temperature, pressure)	0	N/A	0	Both	
	0	Yes	0	Observation	
e. Documentation for each piece of sterilization equipment is	0	No	0	Interview	
maintained and up to date and includes results from each load	0	N/A	0	Both	
	0	Yes	0	Observation	
E. Items are appropriately contained and handled during the	0	No	0	Interview	
sterilization process to assure that sterility is not compromised prior to use	0	N/A	0	Both	
	0	Yes	0	Observation	
F. After sterilization, medical devices and instruments are stored in a	0	No	0	Interview	
designated clean area so that sterility is not compromised	0	N/A	0	Both	
	0	Yes	0	Observation	
G. Sterile packages are inspected for integrity and compromised	0	No	0	Interview	
packages are reprocessed	0	N/A	0	Both	
	0	Yes	0	Observation	
H. Additional breaches in sterilization practices not captured by the	0	No	0	Interview	
questions above were identified (If YES, please specify further in comments)	0	N/A	0	Both	

Comments: (please print and limit comments to the space provided)

HIGH-LEVEL DISINFECTION

Practices to be Assessed	Was Practice Performed?	Manner of Confirmation		
A. Semi-critical equipment is high-level disinfected or sterilized	O Yes O No O N/A	O ObservationO InterviewO Both		
B. Is high-level disinfection performed on site?(If NO, Skip to "F")	O Yes O No O N/A	O ObservationO InterviewO Both		

(A "No" answer does not result in a citation, since ASCs are permitted to provide for high-level disinfection off-site, under a contractual arrangement.)

(Surveyor to confirm there is a contract or other documentation of an arrangement for off-site sterilization by viewing it)

		0	Manual				
	a. <i>If answer to B was</i> YES, please indicate method of high-level	0	Automated				
	disinfection:	0	Other (<i>please</i>				
		prin	<i>ut):</i>				
C L		C 1		0	Yes	0	Observation
	ms are pre-cleaned according to manu			0	No	0	Interview
evide	nce-based guidelines prior to high-leve	el disi	infection	0	N/A	0	Both
D.	a. Medical devices and instruments	are vi	sually inspected for	0	Yes	0	Observation
	residual soil and re-cleaned as needed before high-level disinfection			0	No	0	Interview
				0	N/A	0	Both
			, · · , · · · · · ·		Yes	0	Observation
	b. High-level disinfection equipmen manufacturer instructions	t is m	laintained according t	00	No	0	Interview
	manufacturer instructions			0	N/A	0	Both
	c. Chemicals used for high-level disinfection are:						
				0	Yes	0	Observation
I. Prepared according to man		ufact	urer instructions	0	No	0	Interview
				0	N/A	0	Both
	II. Tested for appropriate con	ncentr	ration according to	0	Yes	0	Observation

manufacturer's instruc	0 0	No N/A	0 0	Interview Both	
III. Replaced accordin	ng to manufacturer's instructions	000	Yes No N/A	0 0 0	Observation Interview Both
Practices to be Assessed	Pra	Practice		Manner of Confirmation	
IV. Documented to ha according to manufact	0 0 0	Yes No N/A	0 0 0	Observation Interview Both	
d. Instruments requiring high-	-level disinfection are:				
	appropriate length of time as turer's instructions or evidence-	0 0 0	Yes No N/A	0 0 0	Observation Interview Both
II. Disinfected at the appropriate temperature as specified by manufacturer's instructions on evidence-based guidelines				000	Observation Interview Both
E. Items that undergo high-level disinfection are allowed to dry before use				0 0 0	Observation Interview Both
F. Following high-level disinfection, clean area in a manner to prevent cor	-	0 0 0	Yes No N/A	0 0 0	Observation Interview Both
G. Additional breaches in high-level disinfection practices, not captured by the questions above were identified (If YES, please specify further in comments)				0 0 0	Observation Interview Both
Comments: (please print and limit comments to the space provided)					

IV. Environmental Infection Control

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

Practices to be Assessed			Manner of Confirmation	
A. Operating rooms are cleaned and disinfected after each surgical or invasive procedure with an EPA-registered disinfectant	0 0 0	Yes No N/A	0 0 0	Observation Interview Both
B. Operating rooms are terminally cleaned daily	000	Yes No N/A	000	Observation Interview Both
C. High-touch surfaces in patient care areas are cleaned and disinfected with an EPA-registered disinfectant	0 0 0	Yes No N/A	0 0 0	Observation Interview Both
D. The ASC has a procedure in place to decontaminate gross spills of blood	000	Yes No N/A	0000	Observation Interview Both
E. Additional breaches in environmental cleaning not captured by the questions above were identified (If YES, please specify further in comments)	0000	Yes No N/A	0 0 0	Observation Interview Both

Comments: (please print and limit comments to the space provided)

V. Point of Care Devices (e.g., blood glucose meter) Observations are to be made of staff who perform fingerstick testing (e.g., nurses)

If N/A is *filled in*, please clarify why *in the comments box below* why it was not applicable or not observed.

Practices to be Assessed	Was Practice Performed? Manner of Confirmation
--------------------------	--

1. Does the ASC have a blood glucose meter? If NO, STOP HERE.	0 0 0	Yes No N/A	000	Observation Interview Both
A. A new single-use, auto-disabling lancing device is used for each patient	0 0 0	Yes No N/A	0 0 0	Observation Interview Both
B. The glucose meter is not used on more than one patient unless the manufacturer's instructions indicate this is permissible	000	Yes No N/A	0 0 0	Observation Interview Both
C. The glucose meter is cleaned and disinfected after every use.	0 0 0	Yes No N/A	0 0 0	Observation Interview Both
D. Additional breaches in appropriate use of point of care devices (like glucose meters) not captured by the questions above were identified (If YES, please specify further in comments)			0 0 0	Observation Interview Both
Comments: (please print and limit comments to the space provided)				