2022 CLABSI Medical Record Abstraction Tool

Refer to associated 2022 MRAT instructions

1. IDENTIFIERS AND ABSTRACTED DATA: Use Tables on page 1 to document information as needed to answer questions beginning on page 2.									
State Facility (NHSN) OrgID				(circle): ACH / LTACH / CancerH / Other/	Dat	Date of Audit/			
Patient ID				Patient DOB//	Rev	Reviewer Initials			
Review Start Time:				End Time:	Tim	e spent review	ing this record (minutes):		
FACILITY Adn	nission Date:			FACILITY Discharge Date://					
2. SCREENIN	NG QUESTIO	NS							
		-		or after facility day 3 or was the date of event	☐ Yes	☐ Yes -> Continue to 2-2			
(DOE) the da	y of transfer of	or discharge	, or the next day	?	☐ No -> (i.e., <u>ALL</u> positive blood specimens were drawn <u>before</u>				
						facility day 3) there was no HAI-CLABSI Event. STOP, record			
						outcome (a) No candidate VL CLABSI			
			ns taken during A	NY validation location (VL) stay, or on the day		> Continue to			
of or day afte	er VL discharg	se?			□ No	> STOP, record	d outcome (a) No candidate VL CLABSI		
2.2.14/	t : (C) :	f	2	AND in older desire a Market feet and a size of a		Cantinanata	2.4		
time?	trai line (CL) i	n place for a	>2 calendar days	AND in place during a VL stay for any period of	Yes -> Continue to 2-4				
				6.11	□ No -> STOP, record outcome (a) No candidate VL CLABSI				
		•		following criteria?	□ No -> Continue				
				E. coli, Enterohemorrhagic E. coli, Vibrio spp, spp. (These organisms are excluded pathogens	☐ Yes -> STOP, record outcome (a) No candidate VL CLABSI				
				e reported as the sole pathogen in a primary					
BSI.)	,,	,							
•	nyces, Histople	asma, Coccid	lioides, Paracoccia	lioides, Cryptococcus, Pneumocystis (These					
organis	sms are typical	lly causes of	community-assoc	ated infections and are rarely known to cause					
			and therefore are	· · · · · · · · · · · · · · · · · · ·					
			organisms identif	·					
_		nin a range o	f two days before	and day after a positive NCT with a recognized					
pathog									
Table 1a. Li	st Positive B	Validation	mens chronolo Optional: CL* on	gically:		1	T		
Positive BC*	Date BC	Location	this date or day		P or	Infection			
	Collection	BC?	before?	Organism genus/species	CC*	DOE*	RIT* End Date and RIT number		
1	, ,					, ,			
	//	ΥN	ΥN			//			
2	, ,	V N	V 81			, ,			
		ΥN	ΥN						
3	, ,					, .			
	//	ΥN	ΥN			-/-/-			
*BC=blood specimen, CL= Central Line, P=pathogen, CC=common commensal, DOE=Date of Event, RIT= Repeat Infection Timeframe. Add rows if needed.									



Table 1	b. Locations:				Table 1c. Central Lines:				
Facility	Physically	Discharge/	Location	Pt in	CL inserted or accessed	CL removed without replacement	Location housed with CL		
Location	Admit/	Transfer	Name (include	VL?					
Order	Transfer IN	OUT	ED)						
1		/		Y N					
2	//	//		YN					
3		//		YN					
4				YN					
5	//			YN					
6	//	//		YN					
7		//		YN	//				
8	8// Y N								
9				YN					
Add rows i	f needed			Ad	dd rows if needed				
3. LABORATORY CONFIRMED BLOOD STREAM INFECTION (LCBI) CRITERIA									
Evaluate all positive blood specimens in order as potential Laboratory Confirmed Bloodstream Infection (LCBI), using table columns in the MRAT Instructions;									
dete	determine if there was an LCBI, and which type (LCBI 1, LCBI 2, or LCBI 3) was met, if any.								

4. Did Infection Episode Qualify as LCBI Event? (begin loop)										
□ No	alternat	If LCBI definition was NOT met, record outcome (b) No LCBI, and reason (i.e., unmatched common commensal, asymptomatic matched common commensals, or alternative primary site infection with secondary BSI), and continue to next Infection Event. If no more positive blood specimens, STOP								
Yes	If Yes LCBI, document type of LCBI and Date of Event below. Note: there may be more than one LCBI during an episode of care.									
			Type of L	CBI (circle one)	:		Date of LCBI Event (date FIRST of required elements was met during the LCBI IWP):			
First LCBI	LCBI 1	MBI LCBI 1	LCBI 2	MBI LCBI 2	LCBI 3	MBI LCBI 3				
Second LCBI	H LCBI 1 MBI LCBI 1 LCBI 2 MBI LCBI 2 LCBI 3 MBI LCBI 3				LCBI 3	MBI LCBI 3				
Third LCBI	LCBI 1	MBI LCBI 1	LCBI 2	MBI LCBI 2	LCBI 3	MBI LCBI 3				
Add rows	Add rows if needed									



5. Wa	s LCBI Healthcare-Associated (HAI) or Present on Admission (POA)?							
Dia	LCBI occur during the time period of 2 days before facility admission to the day after facility admission (POA)?							
☐ Yes	If Yes, LCBI was POA; document outcome (c) POA LCBI type and evaluate next positive blood specimen outside of the event LCBI RIT.							
	If no more blood specimens, STOP							
□ No	If No, proceed to 6.							
6. HA	I-LCBI vs CLABSI?							
6a Was	this HAI-LCBI a CLABSI							
☐ Yes	If Yes, HAI-LCBI is CLABSI; proceed to 6b.							
□ No	If No, document outcome (d) HAI-LCBI not CLABSI and evaluate next positive blood specimen with date of event outside the LCBI RIT.							
	If no more blood specimens, STOP							
6b Was	there medical documentation of the patient suspected or observed self-injecting into their vascular access device within the infection window period?							
☐ Yes	If Yes, document outcome (d) HAI-LCBI not CLABSI and evaluate next positive blood specimen with date of event outside the LCBI RIT.							
	If no more blood specimens, STOP							
□ No	If No, HAI-LCBI is CLABSI; proceed to 6c.							
6c Was	there pus at the site of one of the following vascular access devices and a specimen collected from that site has at least one matching organism to an organism							
identifie	ed in blood							
☐ Yes	If Yes, then disassociate the LCBI from the central line – document outcome (d) HAI-LCBI not CLABSI and evaluate next positive blood specimen with date of							
	event outside of the LCBI RIT.							
□ No	If No, HAI-LCBI is CLABSI; proceed to 7.							

7. WAS	VALIDATION LOCATION (VL) the Location of Attribution (LOA)?						
7a. Was	patient in a VL on date of LCBI Event or day before Event?						
☐ Yes	If Yes, proceed to b.						
□ No	If No, document outcome (e) CLABSI not VL attributable and evaluate next positive blood specimen with date of event outside the previous LCBI RIT.						
	If no more blood specimens, STOP						
7b. Was	s patient transferred to VL from another bedded inpatient location, on date of LCBI Event or day before Event?						
☐ Yes	If Yes, location of attribution was the <u>transferring location</u> . Proceed to c.						
□ No	If No, location of attribution was location at time of infection; STOP record outcome (f) VL CLABSI						
7c. Was	7c. Was the transferring location a validation location (VL)?						
☐ Yes	If Yes, location of attribution (transferring location) WAS a validation location; STOP record outcome (f) VL CLABSI						
□ No	If No, location of attribution (transferring location) was NOT a validation location; record outcome (e) CLABSI not VL attributable and evaluate next positive						
	blood specimen with date of event outside the previous LCBI RIT.						
	If no more blood specimens, STOP						



8 Outcome Documentation								
Positive Blood specimen Number	Outcome (a	-	Detail for out (f) (See key b	-	b) through	Provid	de detail for Case Determination and reason (See key to below)	
1								
2								
3								
4								
5								
☐ Mater ☐ Alter ☐ Primary ☐ Date of ☐ Attach ☐ Circle co ☐ 1. ☐ 2.	t one): aminant (unn hing CCs with native primar source of BS alternative p NHSN checklis orrect NHSN E At least one the NHSN sit + repeat infe	natchen no syry sour I rimary st with assi Cho organi e-spec ction i	ed CC) emptoms ece of BSI (com evevent elements absorpter, Append ism from the le cific infection of	stracted ix B crite plood spe criterion	rion: ecimen match AND the bloo men is an ele	d specin	rganism identified from the site-specific infection that is used as an element to meet nen is collected during the secondary BSI attribution period (infection window period at is used to meet the NHSN site-specific infection criterion, and therefore is collected	
(c) POA LCBI Type of LCBI, S (d) HAI-LCBI r		.CBI1	MBI-LCBI1	LCBI2	MBI-LCBI2	LCBI3	MBI-LCBI3	
Type of LCBI, S (e) CLABSI not	Select one: L		MBI-LCBI1	LCBI2	MBI-LCBI2	LCBI3	MBI-LCBI3	
Type of LCBI, S (f) VL CLABSI;			MBI-LCBI1	LCBI2	MBI-LCBI2	LCBI3	MBI-LCBI3	
Type of LCBI, S Date of VL CLA Location of at	ABSI	LCBI1	MBI-LCBI1	LCBI2 —	MBI-LCBI2	LCBI3	MBI-LCBI3	
		e shou	ıld have an as	signed o	utcome a-f . T	There mo	ay be multiple LCBIs, or multiple CLABSIs during a single episode of care.	



Case Determination		
(A) Correctly Classified	(B) Over-reported HAI	(C) Underreported HAI
If CLABSI was misclassified (over- or underreported) by facility, what	was the reason?	
If CLABSI was misclassified (over- or underreported) by facility, what (I) General HAI definition misapplication (Ia) Incorrect location of attribution (Ib) Date of event incorrect (Ic) IWP set incorrectly (Id) RIT applied incorrectly (Ie) Did not identify elements present in IWP (If) POA/HAI applied incorrectly (Ih) Other (III) Additional Reasons (IIIa) Missed case finding/failure to review positive specimen/culture (IIIb) Clinical over-rule (IIIc) Used outdated criteria	(II) CLABSI criteria misapplied (IIa) Central Line not in > 2 days in an event (IIb) Missed CLABSI due to central line the date of event (IIc) Missed CLABSI due to location trabefore the date of event (IId) CLABSI incorrectly identified as s (IIe) Secondary BSI incorrectly identified (IIf) Other	e removed day of or day before ansfer/discharge day of or day econdary BSI
(IIId) No positive blood specimen in chart (IIIe) Other		

Don't forget to record the abstraction end time on page 1 Location of elements meeting criteria within Medical record: