

Form Approved OMB No. 0920-0666 Exp. Date: xx/xx/20xx www.cdc.gov/nhsn

Central Line Insertion Practices Adherence Monitoring

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*required for saving		Frank III		
Facility ID:		Event #:		
*Patient ID:		Social Security #:		
Secondary ID:		Medicare #:		
Patient Name, Last:	First:	Middle:	_	
*Gender: ☐ F ☐ M ☐ Other		*Date of Birth: / (mm/dd/yyyy)		
Ethnicity (specify):		Race (specify):		
*Event Type: CLIP *Location:				
*Person recording insertion practice data	a: \square Inserter \square	☐ Observer		
	Name, Last:	First:		
*Occupation of inserter:				
☐ Fellow	\square Medical student	\Box Other student \Box Other medical staff		
\square Physician assistant	☐ Attending physic	cian \square Intern/resident \square Registered nurse		
\square Advanced practice nurse \square	\Box Other (specify):			
*Was inserter a member of PICC/IV Tea				
*Reason for insertion:				
☐ New indication for central lin	e (e.g., hemodyna	mic monitoring, fluid/medication administration, etc.)		
☐ Replace malfunctioning central line				
☐ Suspected central line-associ				
Other (specify):				
-		e central line exchanged over a guidewire? Y N		
*Inserter performed hand hygiene prior to central line insertion: \square Y \square N (if not observed directly, ask inserter)				
*Maximal sterile barriers used: Mask	\square Y \square N	Sterile gown ☐ Y ☐ N		
_	•	\square N Sterile gloves \square Y \square N Cap \square Y \square N		
*Skin preparation (check all that apply)	☐ Chlorhexidine	gluconate \square Povidone iodine \square Alcohol		
	☐ Other (specify):		
If skin prep choice was <u>not</u> chlorhe	exidine, was there	a contraindication to chlorhexidine? \square Y \square N \square U		
If there was a contraindication to c	hlorhexidine, indic	ate the type of contraindication:		
\square Patient is less than 2 months of age - chlorhexidine is to be used with caution in patients less than 2 months of age				
\square Patient has a documented/l	known allergy/reac	tion to CHG based products that would preclude its use		
\Box Facility restrictions or safety	concerns for CHC	G use in premature infants precludes its use		
*Was skin prep agent completely dry at		·)	
*Insertion site:	☐ Lower extremit	.y □ Scalp □ Subclavian □ Umbilical □ Upper extremit	ty	
Antimicrobial coated catheter used:] Y 🗆 N			
Assurance of Confidentiality: The voluntarily provided collected with a guarantee that it will be held in strict conficonsent of the individual, or the institution in accordance Public reporting burden of this collection of information is data sources, gathering and maintaining the data needed person is not required to respond to a collection of information.	information obtained in thi idence, will be used only fo with Sections 304, 306 and estimated to average 5 mi , and completing and revie ation unless it displays a c	s surveillance system that would permit identification of any individual or institution in the purposes stated, and will not otherwise be disclosed or released without the disclosed of the Public Health Service Act (42 USC 242b, 242k, and 242m(d)). Inutes per response, including the time for reviewing instructions, searching existing existing the collection of information. An agency may not conduct or sponsor, and a currently valid OMB control number. Send comments regarding this burden estimate ing this burden to CDC, Reports Clearance Officer, 1600 Clifton Rd., MS D-74,)	

Atlanta, GA 30333, ATTN: PRA (0920-0666). CDC 57.125 (Front) Rev 5, v8.5



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*Central line catheter type:			
\square Non-tunneled (other than dialysis)			
\square Tunneled (other than dialysis)	☐ Umbilical		
☐ Dialysis non-tunneled	Other (specify):		
☐ Dialysis tunneled	("Other" should not specify brand names or number of lumens; most lines can be categorized accurately by selecting from options provided.)		
*Did this insertion attempt result in a successful			
Custom Fields			
Label	Label		
Comments			
Comments			