National Healthcare Safety Network (NHSN) External Validation Guidance and Toolkit 2015, for Validation of

- 2015 Central Line-Associated Bloodstream Infection (CLABSI) in ICUs
- 2015 Catheter-Associated Urinary Tract Infection (CAUTI) in ICUs
- Surgical Site Infection (SSI) following 2015 Abdominal Hysterectomy (HYST) Procedure
- Surgical Site Infection (SSI) following 2015 Colon (COLO) Procedure
- 2015 Methicillin-Resistant Staphylococcus aureus (MRSA) Bacteremia LabID Event
- 2015 Clostridium difficile Infection (CDI) LabID Event

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About the 2015 NHSN External Validation Guidance and Toolkit

The 2015 NHSN External Validation Guidance and Toolkit provides guidance for NHSN data validation. Like 2014, CDC provides guidance and tools for validation of six healthcare-associated infection (HAI) metrics: ICU CLABSI, ICU Catheter-Associated Urinary Tract Infection (CAUTI), selected Surgical Site Infections (following colon (COLO) and abdominal hysterectomy (HYST) procedures), Methicillin-Resistant *Staphylococcus aureus* (MRSA) Bacteremia LabID Event and *Clostridium difficile* infection (CDI) LabID Event for 2015 HAI validation. The guidance and tools for CLABSI and CAUTI were designed to work in settings including and beyond acute care hospitals; validation of CLABSI is appropriate for long-term acute care hospitals (LTACs, termed long-term care hospitals by the Centers for Medicare and Medicaid Services, CMS), and validation of CAUTI is appropriate for LTACs and inpatient rehabilitation facilities (IRFs).

The purpose of validation is to assure high-quality surveillance data through accountability and by identifying, understanding, and correcting reporting problems. The focus of this document is external validation of facility-reported NHSN surveillance data conducted by state health departments or other oversight agencies. A separate guidance for facilities that seek to conduct internal validation (quality assurance) of their own NHSN data is also available at http://www.cdc.gov/nhsn/validation/index.html.

This document proposes standard methods for state health departments and other oversight agencies to conduct external validation of reported 2015 HAI data. Developing a standard approach to HAI data validation is important to assure nationwide data quality and to enhance fairness under current and planned reimbursement programs that use NHSN data. States may vary in their regulatory authorities and capacities for NHSN data validation but can best assure equivalent high data quality by striving to follow these standards. NHSN-specified external validation standards are intended to assure concordance of reported surveillance outcomes with those expected under NHSN surveillance definitions and methods, as determined and documented by trained auditors. Recommended sample sizes attempt to balance feasibility with adequate precision for HAI metrics at the facility level. Survey tools are provided to assess reporter knowledge and facility practices required to conduct adequate surveillance.

For 2015 data audits, the specified approach to facility and medical records sampling will be targeted external validation. Targeted validation provides an efficient approach to identify and correct likely reporting errors and their underlying processes in facilities with high volume of exposure to HAI risk, and thus to use limited validation resources as effectively as possible. Accuracy measures (e.g., sensitivity and specificity) derived from a targeted sample are likely to be reduced relative to a more representative sample. Although it may be a simpler and more efficient approach to begin the external validation process, targeted sampling has an important limitation in that representative information is not generated in this way. Future guidance is likely to focus on sampling methods that generate quantifiable representative information regarding NHSN data quality.

Comments and Feedback Welcome: NHSN validation approaches are a work-in-progress and will improve more quickly with the generous input and feedback of those implementing the methods. Please direct any comments or suggestions for improvement to the NHSN Helpdesk: NHSN@cdc.gov.



Acknowledgements and Thanks

Many aspects of this document were adapted from states conducting validation. In addition, many experts from state and local health departments and healthcare facilities collaborated to develop, review, and contribute to this document. The contributions of these individuals are gratefully acknowledged. However, the Guidance and Toolkit recommendations are the sole responsibility of the Centers for Disease Control and Prevention (CDC) and should not be regarded as having received the endorsement of any individuals or organizations outside of CDC.



Abbreviations, Terms, and Acronyms Used in this Document

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ABUTI*	(NHSN) Asymptomatic bacteremic urinary tract infection. This type of UTI may or may not be catheter-associated (CAUTI).						
ADT	Admissions/discharges/transfers (A core facility data system)						
BABY LOCATIONS*	(NHSN) Patient care locations housing a high proportion of infants aged <1 year, i.e. newborn nurseries, neonatal ICUs, and LDRP locations						
BSI	Bloodstream infection						
CAUTI*	(NHSN) Catheter-associated urinary tract infection. A primary UTI where an indwelling urinary catheter was in place for >2 calendar days when all elements of the UTI criteria were first present together AND indwelling urinary catheter was in place on the date of event or the day before.						
CCN	CMS Certification Number, i.e., a facility identifier						
CDC	Centers for Disease Control and Prevention						
CDI	Clostridium difficile Infection						
CEO	Chief executive officer						
CL	Central line						
CLABSI*	(NHSN) Central line-associated bloodstream infection. A primary laboratory-confirmed bloodstream infection (LCBI) where a central line was in place for >2 calendar days when all elements of the LCBI criteria were first present together AND central line was in place on the date of event or the day before.						
CMS	Centers for Medicare & Medicaid Services						
C-SUITE	Office for senior executives such as Chief Executive Officer (CEO) or Chief Medical Officer (CMO) of a healthcare facility						
DELTA COUNT*	(NHSN, as used in this guidance) The absolute difference between the number of expected events and observed events						
DI SSI*	(NHSN) Deep incisional surgical site infection						
DOB	Date of birth						
DOH	Department of health						
ED	Emergency department						
EMR	Electronic medical record						
EPISODE OF CARE	All medical services provided to a patient within a specific time period within a facility. For surveillance of HAIs, this term is used to indicate a single inpatient admission, and includes the ED visit leading to admission						
EXTERNAL VALIDATION	Survey and record review process by external agency to assure quality of NHSN surveillance and reporting						
FacWideIN*	(NHSN) Facility-Wide Inpatient, a type of surveillance used for LabID Event reporting						
FOLEY CATHETER	Indwelling urethral (urinary) catheter						
GI* HAI*	(NHSN) Gastrointestinal system healthcare-associated infection NHSN) Healthcare-associated infection: An infection is considered an HAI if all elements of a CDC/NHSN site-specific infection criterion were not present during the POA time period but were all present on or after the 3rd calendar day of admission to the facility (the day of hospital admission is calendar day 1). All elements used to meet the CDC/NHSN site-specific infection criterion must occur within a timeframe that does not exceed a gap of 1 calendar day between any two adjacent elements. The definition of a gap day is a calendar day during which no infection criterion elements are present.						
IAB*	(NHSN) Intra-abdominal healthcare-associated infection; a subset of GI*						
ICU	Intensive care unit						
	intensive care unit						



(NHSN) Drainage tube inserted through the urethra to the urinary bladder, left in place, and connected to a drainage bag. Also called a foley catheter. May be used for drainage and/or irrigation. Excludes condom catheters, straight in-and-out catheters, nephrostomy tubes, and suprapubic catheters. (NHSN) Surgery in a patient whose date of admission is different from date of discharge SURJERNAL ACTIVE efforts by a reporting facility to assure completeness and accuracy of NHSN data VALIDATION IP Infection preventionist IT Information technology LabiD Event* (NHSN) A measure developed for infection surveillance using laboratory results data without the requirement for extensive clinical documentation and intended for easy electronic reporting LCBI 1,2,2* (NHSN) laboratory-confirmed bloodstream infection criteria LDRP Labor, Delivery, Recovery, and Post-partum, a type of NHSN location in an acute care facility Length of stay (days) MEDICAL A record systematically documenting a single patient's medical history and care across time within a healthcare provider's jurisdiction. For the purpose of sampling, a medical record (which over time could include many healthcare encounters) refers to a single facility inpatient admission. MRN Medical record number MRSA, MSSA Methicillin-resistant <i>Stophylococcus aureus</i> , Methicillin-susceptible <i>Stophylococcus aureus</i> NICU Neonatal intensive care unit NP Nasopharyngeal NISN National Healthcare Safety Network OBSERVATION A Dedded patient care location designated for patients under observation, a form of outpatient status. The purpose of sobervation is to allow the physician time to make a decision about whether the patient should be admitted, and then rapidly move the patient to the most appropriate setting, i.e., admit to inpatient status or to send home. OBSERVATION Status for patients who are undergoing short-term treatment, assessment, and reassessment while a decision is made regarding the need for admission to the hospital. Observation patients may occupy beds								
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SECONDARY* INFECTION	(NHSN) Site affected by infection by dissemination from an alternative originating source (see PRIMARY)
SIR*	(NHSN) Standardized infection ratio
SI SSI*	(NHSN) Superficial incisional surgical site infection
SSI*	(NHSN) Surgical site infection
SUTI*	(NHSN) Symptomatic UTI
TARGETED SAMPLE	In this document, a purposive sample taken to target facilities at higher risk for HAI or medical records at higher risk for misclassification of HAI status (See also, purposive sample)
URINARY CATHETER*	(NHSN) See indwelling urinary (urethral) catheter.
UTI	Urinary tract infection
TERTILE	Lowest, middle, or highest one-third of a group
VAE*	(NHSN) Ventilator-associated event. An objective surveillance algorithm that can identify a broad range of conditions and complications (including but not limited to pneumonia) occurring in mechanically-ventilated adult patients, detailed in NHSN Manual Chapter 10.
VALIDATION	Assurance that reported NHSN surveillance data meet their pre-determined specifications and quality attributes as intended

^{*(}NHSN) indicates a term used and defined by NHSN



Chapter 1: Overview and 2015 Validation Standards

Validation can be defined as confirming or assuring that data meet pre-determined specifications and quality attributes. NHSN validation should assure high quality of three domains in reporting healthcare-associated infections (HAIs): denominators, numerators, and risk adjustment variables.

Why Validate?

NHSN was launched as a voluntary, confidential HAI reporting system for hospitals conducting surveillance, benchmarking, and quality improvement for HAIs. Since 2006, NHSN data have also been used by state and federal agencies for public reporting purposes and increasingly are used to incentivize quality improvement through payment mechanisms. These new uses have heightened the importance of the completeness and accuracy of the data. Hospital boards, administrators, and clinical leadership need to trust their own facility's data to assess performance, manage change in their facilities, and to know that other facilities are held to the same high standards when reporting. Consumers seeking to make informed decisions about their healthcare expect that publicly reported data are valid. These requirements are challenging because NHSN definitions are complex and may involve tracking and linking information from multiple hospital information systems (e.g., laboratory, admissions, and clinical data); coordinated data collection, interpretation, and entry by multiple staff members; and sometimes require subjective interpretation, all of which introduce opportunities for variation. This complex landscape will continue to change over time as NHSN methods evolve, use of electronic medical records increases, and reporting requirements expand.

In the context of powerful inducements for facilities to "look good," meaningful external validation is essential to assure that NHSN surveillance meets the requirements for which it was intended; that outcomes for reporting facilities are appropriate, that NHSN data are credible, and that the focus of NHSN surveillance will be better patient care and disease prevention. In the absence of meaningful external validation, healthcare facilities may fail to identify or report HAIs. This would not require overt gaming because variation in effort, resources, and practices between facilities can result in surveillance bias ("the harder you look the more you find") and in assessment bias ("we tend to see what we want to see"). For example, approaches to surveillance that create barriers to reporting, such as requiring the agreement of multiple reporters or permission from authorities before reporting can lead to lower measures of disease rates without improving patient safety. To provide for fair comparisons of facilities, standard surveillance and reporting methods must be adequately resourced and adhered to, data accuracy and completeness must be optimized, and risk adjustment for patient mix applied appropriately.

Validation is an important step toward assuring that reported NHSN data are actionable and motivate improved infection control efforts rather than strategies to avoid accounting for HAIs. Accurate, high quality NHSN data are important to infection prevention programs for setting priorities and measuring the impact of prevention efforts. Further, public health agencies at the local, state and federal levels need these data to identify HAI problems and to measure prevention program success. Each of these data users also has a role and a stake in assuring quality of NSHN data.



External Validation

External validation is a survey and audit process conducted by an agency outside the reporting facility (e.g. state health department), in which a facility's surveillance determinations and methods are investigated by one or more trained validators who work for the external agency, to evaluate surveillance program quality (e.g. knowledge and practices), and completeness and accuracy of reporting. Findings from external validation can be used to correct reporter misconceptions about NHSN definitions, criteria, and data requirements. As a result, external validation can help assure adherence to NHSN's specifications for HAI reporting by identifying and correcting shortcomings that would be difficult to address through internal validation alone. Data correction and completion should be required of reporters, and helping reporters understand what led to the errors enhances the likelihood of better reporting in the future. Common errors and challenging cases should be documented to derive information for teaching and to improve future reporting.

Sampling of hospitals and medical records for review can be done in a variety of ways to meet different goals. It is typically not possible or necessary for validators to visit every facility or review every patient record in search of candidate HAIs. Sampling is a practical necessity, and sampling methods should strike a balance between resource availability and programmatic objectives.

2015 Validation Guidance

For 2015 data validation, this guidance document specifies an algorithm for targeted sampling that provides for efficient investigation of potential surveillance and reporting problems in highly exposed facilities and medical records, where HAIs are most expected. Exposure risk derives from increased device days, surgical procedures, or specified positive laboratory test results, and targeting is driven by either high or low event reporting. In targeted samples, the ability to produce generalizable information about the population as a whole is constrained. A favorable outcome under targeted sampling suggests that success would be even more likely in a probability sample representing the entire population at risk. Because all facilities should be held accountable for accurate reporting, and smaller facilities that are unlikely to be targeted given low exposure risk may actually derive great benefits from validation, a 5% random sample of additional facilities should also be drawn after the targeted facility sample has been selected. States should not be constrained by the algorithm, and should seek adequate reporter training and internal quality assurance of all reporting facilities in their jurisdiction, even those that are not audited.



Chapter 2: Guidance for Conducting 2015 NHSN Validation

A targeted validation approach is recommended to use resources as efficiently as possible to identify reporting errors, particularly errors caused by correctable systematic surveillance problems or misconceptions. The recommended sample sizes and enriched sampling frames provide a reasonable chance to identify reporting errors if they exist. The scope of external validation includes six metrics: ICU CLABSI, ICU CAUTI, COLO SSI, HYST SSI, MRSA Bacteremia LabID Event and CDI LabID Event which are consistent with CMS Inpatient Quality Reporting Program requirements.

If unable to secure resources to complete the validation standard for all six HAIs listed above at the prescribed number of facilities, then narrow the scope of HAIs to be validated, while maintaining the sample sizes for chosen metrics and the recommended number of facilities to derive robust information about performance at facilities for selected metrics.

When selecting which HAIs to validate, oversight agencies may choose to use experience and/or data analysis to prioritize choices. For example, if validation of ICU CLABSI was completed as recommended in the 2014 Validation Guidance and Toolkit, agencies may seek to focus on other HAIs for 2015. Those with high rates of a particular HAI may wish to focus validation on this problem to assist facilities with prevention.

Facilities that will not be targeted for external validation audits using this suggested sampling method should still be held accountable for high quality surveillance and reporting programs and for conducting internal validation activities. Requesting evidence of up-to-date NHSN reporter training (such as a 2015 certificate of successful completion produced by each of NHSN's multimedia training modules from all facilities) is one way to assure appropriate reporter training without a site visit. Some may wish to administer surveillance process surveys or request documentation of internal validation activities by facilities.

For audited facilities, recommended external validation for 2015 includes assessment of numerators, denominators, and risk-adjustment variables, with medical records audit focused on outcomes (numerators). Numerator quality can be assured by a) adequate reporter knowledge (as demonstrated by completed certificates for 2015 online multimedia assessments), b) good surveillance practices (assessed by survey), and c) evidence of correct reporting (by an audit of medical records showing concordance of validator outcomes with events reported to NHSN). Denominators can be assessed by a) review of denominator data records, b) denominator collection practices surveys, and c) (for COLO and HYST procedures) comparison of crude monthly procedure counts in NHSN with ICD-10-PCScodes generated by the facility. Risk adjustment variables and documentation of internal validation work conducted by facilities should also be reviewed.

This external validation guidance and toolkit, informed by state and facility experience and the need for standardized validation methods, recommends on-site medical record reviews by trained validators using a medical record abstraction tool that follows 2015 NHSN methods and definitions, with CDC serving as adjudicator of discordant outcomes when necessary. On-site validation provides optimal opportunity for validators to gain full access to any documented information used by reporters when



conducting surveillance, and to strengthen relationships with reporting facilities through transparency. Use of electronic medical records systems that are made available at a distance to validators is a feasible, though perhaps a sub-optimal alternative way to audit medical records. This approach may require technical expertise and iterative work with facilities to assure validator access to all relevant documentation. In addition, without site visits the opportunities for interaction, education, and understanding of the overall HAI surveillance program are likely to be reduced. Remote review of copied medical records is discouraged for external validation program methodology, as potentially lacking complete data access and the interactivity that facilitates program capacity building. Ideally, validators will be either employed or contracted by agencies that have oversight responsibilities for patient safety and public health in the audited healthcare facilities, and across the continuum of healthcare.

CDC-Recommended Validation Elements and Preferred Approach

Validation Element	Off-site	On- or Off-site	On-site
Validator training and assessment	X		
NHSN Data analysis for completeness, timeliness, and quality	X		
Facility selection, request for line listings (CLABSI, CAUTI, MRSA	Х		
bacteremia, and CDI), and monthly surgical procedure counts			
(COLO, HYST)			
HAI Sampling Frame Development	Х		
Medical Record Selection, NHSN data download, and arrangements	Х		
for audit			
Facility surveillance Practices Surveys (Appendix 2)		Х	
Review of facility mapping, bed size			Х
Medical Record Reviews (Appendix 3)			Х
Post-review conference with IP re: surveillance practices and			Х
medical records audit discrepancies			
Administration of additional denominator counting surveys, as		Х	
needed			
Review of facility results, strengths, and weaknesses		Х	
Follow-up corrections and report to IP and administration	Х		



Chapter 3: Preparation for External Validation

1. Assure or update validator expertise in 2015 definitions

For CLABSI, be aware that important changes in NHSN require surveillance for, and reporting of, Mucosal Barrier Injury-Laboratory Confirmed Bloodstream Infection (MBI-LCBI) events introduced at the beginning of 2015. Additionally, the definition of neutropenia in the MBI-LCBI criteria was expanded in 2015. These additions and definitions can affect case-ascertainment and classification for CLABSI events. Validators MUST be familiar with these to correctly audit NHSN cases. The Medical Records Abstraction Tools are also designed to support these changes. Additional instructions for location mapping may affect location of attribution and risk adjustment for device-associated events, and should be part of the audit and survey process.

For SSI, be aware that important changes in definitions include the addition of: height and weight; diabetes status; incisional closure type (primary vs. non-primary); and a modified definition of procedure duration. The SSI definition changes can affect reporting of procedures (denominators) and SSI case-ascertainment (numerators). Validators need to be familiar with these changes to correctly audit procedures and SSIs in NHSN.

Surveillance and validation require rigorous adherence to standard NHSN protocols, surveillance methods, and NSHN definitions as written. Persons conducting audits must be trained in NHSN specifications, remain up-to-date when changes are made, and commit to using appropriate NHSN methods and definitions to validate HAI data reported to the system. In addition to reporter training resources, validator training resources are available on the NHSN website and will be expanded in the future (http://www.cdc.gov/nhsn/validation/2015/index.html). The following trainings are available on the training website. They are listed in order of recommendation for validators:

Type of NHSN	Recommended Validator Standard	Symbol Key for Online NHSN Training
Training		Types (Examples as below)
Interactive Online Multimedia Instruction Modules	Assure that all 2015 validators successfully complete these courses for any NHSN component they will validate, and provide copies of the certificates of completion	Self-paced, interactive trainings used to gain in-depth knowledge of NHSN HAI definitions
Slide sets	Highly recommended: Slide presentations include case-studies to help validators implement the basic content presented in HAI training webinars	Presentations and case studies used to walk through difficult cases to learn to apply the NHSN HAI definitions accurately
Webinars & Podcasts	Basic prerequisite for prospective validators; Basic training in HAI surveillance	Webinars and podcasts used to provide basic information on NHSN HAI definitions and surveillance protocols



Other opportunities for training include:

- CDC-sponsored trainings.
- NHSN blast emails, external partner calls, the quarterly NHSN newsletter, and the NHSN Manual, updated prior to each January with any changes to methods and definitions.

Even after training, willingness to seek help when needed from NHSN on definitions and criteria is important when cases are challenging. If facilities and auditors cannot agree on case-status using documented information and the NHSN case-definition as a gold standard, the case should be referred to CDC for adjudication. Forms for tracking cases that result in discrepancies and that require adjudication are found in <u>Appendices 4.1</u> and <u>4.3</u>.

Finally, although it is not required, duplicate abstraction of medical records by another auditor (early in the process and periodically repeated) may be a useful adjunct to validator training, in order to identify areas of difficulty and to achieve improved inter-rater reliability.^{3,4}

2. Select facilities

For 2015, as in 2014, CDC recommends targeted validation in order to investigate and correct potential deficiencies in an efficient manner, given the assumption of limited resources for validation. This approach also provides maximum opportunity to work with reporters to improve reporting.

The exercise for facility selection can also be used as a means of developing situational awareness about exposure risk and performance of facilities. This information may be useful for targeting prevention programs.

3. Establish a mechanism for secure data transfer between facilities and the state health department

To build a sampling frame for medical record selection, electronic files (spreadsheets) are required from laboratories that list positive blood cultures, positive quantitative urine cultures, and positive CDI toxin tests, with test dates, patient locations, identified pathogens and patient information to identify medical records for review. In addition, assistance may be needed from hospital medical records departments to identify hospital re-admissions within the surveillance window (3 months for COLO and HYST) of audited surgical procedures. Some agencies have established secure FTP sites for transfer of these sensitive data. Consider existing systems for secure data transfer and how to secure these data in both directions--to send line listings to characterize the sampling frame and to respond with the sample of medical records to be reviewed.

4. For each selected facility, know which HAIs you will be validating, based on information derived from the algorithms in Chapter 4, and your individual priorities and goals.

Before the validation process, for each selected facility and HAI to be validated, record the total number of HAI events reported by the facility for 2015 using <u>Appendix 4.3</u>, "Numerator Validation."



5. Develop and characterize the medical record sampling frame for each selected facility and each HAI to be validated, and for SSI assure a complete denominator:

For CLABSI, CAUTI, MRSA Bacteremia LabID Event and CDI LabID Event, sampling frames derive from positive laboratory (blood culture, urine culture, and CDI toxin-positive specimen) line-listings in surveillance locations. Hospitals should be encouraged to develop capacity to generate these lists electronically, because recurring need for this capability is expected, and creation of manual line-listings would present an excessive burden.

Facilities should report positive laboratory tests according to date of specimen collection, not date of result reporting.

In order to assure completeness of the laboratory line-listings, it is generally recommended that laboratory data derive directly from the laboratory information management system and not from vendor software (such as data-mining programs). However, if convincing evidence exists that vendor software can provide complete laboratory data, vendor systems may provide convenient linkage to ADT data that would otherwise need to be created. This issue may need to be explored through individual discussions with facilities, and by facilities with their vendors.

For SSI, sampling frames derive from procedures in NHSN. However, to assure that the NHSN procedure sampling frame is complete, a monthly tally from the facility for COLO procedures and HYST procedures performed, based on ICD-10 procedure/CPT codes in discharge data should be used. This data request may be made along with the line listing requests and the procedure numbers entered in Appendix 4.3, "Denominator Validation COLO" and "Denominator Validation HYST." If these numbers are reasonably close to the number of procedures listed in NHSN, the procedure denominator data are presumed to be relatively complete.

Structure of laboratory line listings

Validators need to be able to identify NHSN-reported HAIs on laboratory line listings. Facilities should be reporting HAIs to NHSN using the medical record number (MRN), and may also use patient name. In most cases, matching of reported HAIs will be based on MRN, gender, date of birth, and date of event. In some situations, more information may be needed from the IP about reported NHSN events to identify reported HAIs on the laboratory line listing, e.g. a request for additional personal identifiers of patients with NHSN-reported HAIs that can be linked to laboratory-reports.

The selected sample of positive laboratory tests also will need to be linked to patient medical records for review. The required patient MRN and laboratory test date from the line listing will be the primary identifiers for this purpose, but knowing patient date of birth, admission date, and possibly patient name may facilitate the request to medical records for record audits. If the facility can provide these fields with the line listing they should be requested.

CLABSI in ICU

From each selected facility, obtain a complete list of positive ICU blood cultures collected in 2015 to select the medical record sample before the site visit. A spreadsheet file (e.g. Excel) is recommended for ease of use.



Template positive ICU blood culture line listing (* indicates required data):

*MRN	*Facility	*Laboratory	*Specimen	*Blood	*Specific	*Gender	*Date	First	Last	l
	Admission	Specimen	Collection	Organism 1	ICU		of Birth	Name	Name	1
	Date	Number	Date	Genus and	Location					l
				Species						l

- For positive ICU blood cultures, the medical record number, admission date, laboratory specimen number, the date of specimen collection (not date of report), the resulting first organism ("Org 1") genus and species, specific ICU location, gender, and date of birth are required. Additional patient identifiers such as patient name may be helpful. If needed, ask the IP to translate specific patient location information on the laboratory line listings to mapped NHSN ICU locations, and assure that results for all ICUs are included. Be sure it is possible to distinguish NICU from adult/pediatric ICU locations on this line listing to stratify the CLABSI sample. No information about central line use should be requested; validators will screen for this information while reviewing records.
- Using the line listing, sort by MRN and facility admission date (which together characterize unique eligible admissions/episodes of care with possible ICU CLABSI), then enumerate the eligible episodes of care using the spreadsheet. Enter the number of unique episodes of care eligible for CLABSI review for the year in <u>Appendix 4.3</u>, "Numerator Validation, Sampling Frame Information."

CAUTI in ICU

From each selected facility, obtain a complete list of positive ICU urine cultures collected in 2015 to select the medical record sample before the site visit. A spreadsheet file (e.g., Excel) is recommended for ease of use. Limit positive ICU urine cultures to those with no more than 2 identified pathogens and at least 10⁵ CFU/ml organisms.

Template positive ICU urine culture line listing (* indicates required data; †second organism information is conditionally required):

*MRN	*Facility	*Laboratory	*Specimen	*Urine	*Urine	†Urine	†Urine	\ \ \
	Admission	Specimen	Collection	Organism 1	Colony	Organism 2	Colony	\
	Date	Number	Date	Genus and	Count 1	Genus and	Count 2	Continued
				species	(CFU/ml)	Species	(CFU/ml)	\

• For positive ICU urine cultures, the medical record number, facility admission date, laboratory specimen number, specimen collection date, identity of organisms (up to two) and colony counts (CFU/mI), specific ICU location, gender, and date of birth are needed. Additional patient identifiers such as patient name may be helpful. If needed, ask the IP to translate specific patient location information on the laboratory line listings to mapped NHSN ICU locations, and assure that results for all ICUs are included. Urine specimens with mixed flora, more than two organisms, or fewer than 10⁵ CFU/mI organisms will be rejected. No information about indwelling urinary (Foley) catheter status should be requested; validators will screen for this information while reviewing records.



Using the line listing, sort by MRN and facility admission date (which together characterize the
eligible admissions/episodes of care with possible ICU CAUTI), then enumerate unique eligible
episodes of care using the spreadsheet. Enter the number of episodes of care eligible for CAUTI
review for the year in <u>Appendix 4.3</u>, "Numerator Validation, Sampling Frame Information."

COLO Procedures

- For each selected facility, use NHSN to determine the number of reported COLO procedures conducted in 2015. Enter the number of NHSN-reported COLO procedures in Appendix 4.3, "Numerator Validation, Sampling Frame Information."
- Use NHSN to determine the <u>monthly</u> number of reported COLO procedures conducted in 2015.
 Record the results in <u>Appendix 4.3, "Denominator Validation COLO."</u> (These monthly data will be compared to the facility report generated below to assure that the procedure denominator is complete).
- Provide the list of ICD-10 procedure codes for NHSN COLO procedures and ask the facility to provide
 a monthly count of COLO procedures conducted in 2015, derived from hospital discharge data.
 Record the results in <u>Appendix 4.3, "Denominator Validation COLO,"</u> juxtaposed by month with the
 number of COLO procedures entered into NHSN for each month as determined above.

HYST Procedures

- For each selected facility, use NHSN to determine the number of reported HYST procedures conducted in 2015. Enter the number of NHSN-reported HYST procedures in <u>Appendix 4.3</u>, "Numerator Validation, Sampling Frame Information."
- Use NHSN to determine the <u>monthly</u> number of reported HYST procedures conducted in 2015.
 Record the results in <u>Appendix 4.3, "Denominator Validation HYST."</u> (These monthly data will be compared to the facility report generated below to assure that the procedure denominator is complete).
- Provide the list of ICD-10- procedure codes for NHSN HYST procedures and ask the facility to provide
 a monthly count of HYST procedures conducted in 2015, derived from hospital discharge data.
 Record the results in <u>Appendix 4.3, "Denominator Validation HYST,"</u> juxtaposed by month with the
 number of HYST procedures entered into NHSN for each month as determined above.

MRSA bacteremia LabID Event, facility-wide, inpatient (FacWideIN)

From each selected facility, obtain a complete list of blood cultures positive for methicillin-resistant *Staphylococcus aureus* (MRSA: includes *S. aureus* cultured from any specimen that tests oxacillin-, cefoxitin-, or methicillin-resistant by standard susceptibility testing methods or by a laboratory test that is FDA-approved for MRSA detection). Include those collected in 2015 for inpatient location/ED/ 24 hour observation unit facility-wide, to select the patient admissions/episodes of care for which review is planned. A spreadsheet format is recommended for ease of use. These laboratory line lists should include patient location at the time of specimen collection.



Template positive MRSA bacteremia, FacWideIN line listing (* indicates required data):

*Medical Record Number	*Facility Admission Date	*Laboratory Specimen Number	*Specimen Collection Date	*Blood Organism Genus and Species (Documenting S. aureus or MRSA)	*Documentation of Methicillin- Resistance (susceptibility test result or MRSA)	Continued
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{					
continued	*Specific Mapped	*Gender	*Date	First	Last
}	NHSN Location at		of Birth	Name	Name
\$	Specimen Collection				

- For positive MRSA bacteremia LabID Event (facility-wide, inpatient), the medical record number, facility admission date, laboratory specimen number, specimen collection date, documentation that specimen source was blood, genus and species, methicillin susceptibility information (organism ID may be shortened to MRSA, covering genus, species, and methicillin susceptibility requirements), specific inpatient or emergency department (ED) location/ 24 hour observation location, gender, and date of birth are required. Additional patient identifiers such as patient name may be helpful.
- Using the line listing, sort by MRN and facility admission date (which together characterize the
 eligible admissions/episodes of care with possible MRSA bacteremia LabID Event), then "count" the
 number of unique eligible episodes of care using the spreadsheet. Enter the number of episodes of
 care eligible for MRSA bacteremia LabID Event review for the year in Appendix 4.3, "Numerator
 Validation, Sampling Frame Information."

CDI LabID Event, facility-wide, inpatient (FacWideIN)

To create a sampling sample, obtain from each selected facility, a complete list of final *Clostridium difficile* toxin-positive laboratory results collected in 2015 for inpatients facility-wide [excluding NICU, skilled care nursery, babies in labor/delivery/recovery/post-partum (LDRP) locations, or well-baby nurseries]/ ED/ 24 hour observation units. Laboratories may conduct one- two- or three-step testing for toxigenic *C. difficile* on unformed stool specimens; regardless of the testing approach, only final positive results indicating the presence of toxin-producing *C. difficile* should be included.

A spreadsheet format is recommended for ease of use. These laboratory line lists should include patient location at the time of specimen collection.

Template positive *C. difficile* assay FacWideIN line listing (* indicates required data):

*MRN	*Facility	*Laboratory	*Specimen	*Result of Final CDI	* Specific Mapped	
	Admission	Stool	Collection	Toxin Test (assure	NHSN Location at	Continued
	Date	Specimen	Date	test is toxin-	Specimen	
		Number		positive for CDI)	Collection	

continued	*Gender	*Date of	First	Last
}		Birth	Name	Name



- For positive CDI LabID Event (facility-wide, inpatient), the medical record number, facility admission date, stool specimen number, specimen collection date, result of final CDI toxin test, specific inpatient (or ED/ 24 hour observation) location, gender, and date of birth are required. Additional patient identifiers such as patient name may be helpful.
- Using the line listing, sort by MRN and facility admission date (which together characterize unique eligible admissions/episodes of care with possible CDI LabID Event), then enumerate the eligible episodes of care using the spreadsheet. Enter the number of episodes of care eligible for CDI LabID Event review for the year in <u>Appendix 4.3</u>, "Numerator Validation, Sampling Frame Information."

6. Notify facilities of the planned audit and request the required laboratory line listings

The request may include:

- a) 2 separate positive blood culture line listings (positive ICU blood cultures for CLABSI validation and positive MRSA bacteremia for LabID event validation in facility-wide inpatients/ ED/ 24 hour observations.)
- b) Positive ICU urine cultures for CAUTI validation
- c) CDI toxin-positive specimens for LabID Event validation in facility-wide inpatients/ ED/ 24 hour observations.
- d) Monthly totals for COLO and HYST procedures from medical records-based monthly ICD-10-PCS procedure totals

For chosen facilities, contact the IP and discuss the audit process, including the likely scope of the audit and how the audit sample will be drawn from eligible medical records. Discuss the current request for blood culture, urine culture, and C. difficile toxin-positive line listings for appropriate patient populations (with structures described above). If all six HAIs will be validated, up to 60 specific medical records will be requested each for ICU CLABSI and ICU CAUTI, up to 60 medical records each for COLO and HYST procedures with any subsequent admissions within 3 months following the procedure, and for LabID Event, access to either a) ADT data and complete inpatient and outpatient laboratory records for 60 specified episodes of care each for MRSA bacteremia and CDI LabID Event auditing OR b) corresponding medical records that include these elements during on-site validation. Ask about the lead-time for the facility to generate the required line listings and how much lead-time the medical records department will need to arrange for medical record access. Ask how patient medical records can best be accessed onsite and how they are organized; this can affect the time required to abstract the records. Disorganized records on microfilm may be particularly difficult and time-consuming to abstract. Discuss the anticipated number of days and reviewers needed to complete the audit, based on experience or the guidance to follow. Request documentation that the facility's NHSN reporters have completed training on 2015 NHSN reporting methods and definitions. In addition, a monthly breakdown of how many COLO and HYST procedures were conducted using ICD-10-PCS coded data should be requested if these will be validated.

Consider a mutually agreeable <u>due date for the laboratory line listings</u>, <u>dates for the medical record</u> <u>request</u>, and proposed <u>date(s)</u> <u>for the onsite audit</u>. For the audit, request arrangements for medical



records access including e.g., workspace, computer systems, terminals and passwords, microfilm readers, and (eventually) specific medical records.

The laboratory line listings should be provided by the facility through a secure file transfer (for example, encrypted email, secure FTP site, or encrypted file by courier, or snail mail) as a sortable and searchable (e.g., .csv, Excel) file, and should include facility information (identity and NHSN facID), hospital contact name, hospital contact phone, hospital contact email, date of report, and timeframe of laboratory results.

Compose a letter notifying the facility CEO and copied to the IP that provides an overview of your authority to conduct validation (if applicable) or requesting voluntary access to medical records for the audit process, the purpose of the audit, proposed dates for the audit, and specific data and accommodations needed from hospital staff (see Appendix 1.2 for an example letter). Explain the purpose of the audit (i.e., to assure accountability of all hospitals in complete and accurate reporting of HAIs according to NHSN methods and definitions) and how validation results will be used and/or reported.

- 7. Select medical records (to be discussed in the next chapter)
- 8. Download ("freeze") the facility's reported data from NHSN before disclosing which medical records were selected for the audit.

Do this after selecting the medical records sample to minimize downloads, using NHSN analysis. We suggest using CDC-defined output with the modifications below for freezing and exporting reported 2015 NHSN data.

NOTE: All output options should be exported using the "Export Output Dataset" option at the bottom of the modification screen within NHSN. For more information about how to make modifications to these output options, please see the Analysis Quick Reference Guide library at: http://www.cdc.gov/nhsn/PS-Analysis-resources/reference-guides.html.

Output Option: Line Listing – All CLAB Events

Found within: Device-associated Module > Central Line Associated BSI **Purpose:** Obtain a line listing of all CLABSI events in ICU and NICU locations

Suggested Modifications:

- Change the output title to "<Facility ID > Freeze Data <Freeze Date > ICU CLABSI, 2015"
- Specify time period as: specDateYr 2015 to 2015
- Specify other selection criteria: locationType IN ('CC CC', 'CC_N CC_N')
- Indicate "Sort" variables (optional)

Output Option: Line Listing – All CAU Events

Found within: Device-associated Module > Urinary Catheter-Associated UTI

Purpose: Obtain a line listing of all CAUTI events in ICU locations

Suggested Modifications:

- Change the output title to "<Facility ID > Freeze Data <Freeze Date > ICU CAUTI, 2015"
- Specify time period as: specDateYr 2015 to 2015
- Specify other selection criteria: **locationType** IN ('CC CC')



Indicate "Sort" variables (optional)

Output Option: Line Listing – All Procedures **Found within:** Advanced > Procedure-level Data

Purpose: Obtain a line listing of all COLO and HYST procedures, with associated surgical risk-adjustment

variables

Suggested Modifications:

Change the output title to "<Facility ID > Freeze Data <Freeze Date > COLO procedures (or HYST procedures), 2015"

• Specify time period as: procDateYr 2015 to 2015

Specify other selection criteria: procCode = COLO (or procCode=HYST)

 Indicate "Select Available Variables" including (optional) procID, procCode, dob, patID, gender, procDate, modelRiskAll, asa, anesthesia, scope, emergency, trauma, ageAtProc, swClass, procDurationHr, procDurationMin

Output Option: Line Listing – All SSI Events

Found within: Procedure-associated Module > SSI

Purpose: Obtain a line listing of all COLO (or HYST) SSI events

Suggested Modifications:

- Change the output title to "<Facility ID > Freeze Data <Freeze Date> <procedure> SSI, 2015"
- Specify time period as: specDateYr 2015 to 2015
- Specify other selection criteria: procCode = COLO (or procCode=HYST)
- Indicate "Sort" variables (optional)

Output Option: Line Listing for All CDIF LabID Events

Found within: MDRO/CDI Module - LABID Event Reporting > All C. difficile LabID Events

Purpose: Obtain a line listing of all C. difficile LabID Events

Suggested Modifications:

- Change the output title to "<Facility ID > Freeze Data <Freeze Date > CDIF LabID Events, FacWidelN, 2015"
- Specify time period as: specDateYr 2015 to 2015
- Specify other selection criteria: "cdif" =Y,
- Indicate "Sort" variables (optional)

Output Option: Line Listing for All MRSA LabID Events

Found within: MDRO/CDI Module - LABID Event Reporting > All MRSA LabID Events

Purpose: Obtain a line listing of all All MRSA Blood LabID Events FacWideIN

Suggested Modifications:

- Change the output title to "<Facility ID > Freeze Data <Freeze Date> MRSA Blood LabID Events FacWidelN, 2015"
- Specify time period as: specDateYr 2015 to 2015
- Specify other selection criteria: "mrsa"=Y, "SpecimenSource"= (BLDSPC)
- Indicate "Sort" variables (optional)

9. Request selected medical records in advance of the facility site-visit

Submit the medical records request to the facility in a secure fashion so they can arrange for access to the information for your visit.



Chapter 4: Targeted Sampling of Facilities and Medical Records

Targeted Facility Sampling Overview (see detailed algorithm in **Appendix 1.1**)

Validators are encouraged to complete the facility ranking algorithms in Appendix 1.1 for the six HAI types. If CLABSI and CAUTI will be validated in facility types other than acute care hospitals, separate rankings should be completed for acute care hospitals, long-term acute care hospitals (LTACs), and inpatient rehabilitation facilities (IRFs). This will provide a system for assigning relative priority to each facility for each HAI. Even for those not planning to conduct validation, this ranking activity provides awareness of which facilities are highly exposed to HAI risk and those reporting high or low event outcomes. Additional analyses to evaluate data completeness, timeliness, and quality also are encouraged. In particular, targeted sampling of hospitals performing the surgical procedures to be audited and of the surgical procedures themselves requires that risk-adjustment variables (e.g., ASA score, anesthesia, procedure duration) are complete. Analysis to assure completeness of these variables is recommended before facilities are ranked for SSI validation.

Ultimately, validation resources must be weighed and decisions made as to which HAIs will be validated based on past validation work, need for information on data quality and training needs, unrealized disease prevention, and perceived utility for prevention activities. The facility rankings should help with logistical planning when these considerations are weighed.

- The recommended approach to facility selection for 2015 is targeted (as was recommended for 2014 validation) to prioritize validation of facilities where HAIs are most expected. A recommended minimum number of facilities should be validated (with a recommended minimum number of medical records) for each selected HAI:
 - o Smaller states/jurisdictions with 20 or fewer facilities should validate them all
 - Medium states with 21 to 149 facilities should select at least 18 targeted facilities plus a 5% random sample of remaining facilities
 - Larger states with 150 or more facilities should select at least 21 targeted facilities plus a 5% random sample of remaining facilities

Ranking Algorithm

- For each HAI, sort facilities based on predicted/expected number of events.
- After sorting, the top tertile (33%) of facilities will undergo further targeting and prioritization, based on performance, using the facility SIR relative to the median SIR for the top tertile group of facilities. Detailed guidance for this process is found in Appendix 1.1.
- If the minimum number of targeted facilities is not reached within the top tertile alone, the process should be repeated by targeting the second tertile, and (if necessary) the third.
- If additional facilities are needed to achieve the recommended minimum number, facilities without a calculated SIR may be considered for validation based on the "delta count", defined as the absolute difference between expected and observed NHSN Events reported to NHSN.
- For each HAI, all unselected facilities from all 3 tertiles will be subject to a 5% random sample in order to assure accountability for facilities that are not highly exposed.
- If you choose to validate multiple HAIs at your facilities then you will need to evaluate the facilities to be chosen based on where they rank after you've completed the ranking algorithm for each HAI individually.



Targeted Medical Record Sampling Overview (see detailed algorithms in **Appendix 1.3**)

For sampling, a medical record refers to the record of a single facility inpatient admission, also referred to as an episode of care. For surgical procedures, the episode of care refers to the procedure and all associated medical encounters documented during the surveillance follow-up window. For each HAI to be validated, a sample size of 60 Medical Records/Episodes of Care per facility is recommended as a goal.

For CLABSI, CAUTI, COLO and HYST validation, up to 20 reported NHSN infection events will be reviewed. If more than 20 events have been reported to NHSN, 20 should be selected by random sampling. If less than 20 are reported, all events should be reviewed. In addition, a sampling frame of eligible (candidate) medical records will be developed for each HAI and from these 40 unreported "candidate events" will be selected, by targeting those with increased risk of event occurrence, where this is possible. Definitions of candidate events for each type of HAI and methods for targeting candidate events at increased risk for HAI are described below. Thus a total of (up to) 60 episodes of care containing reported or candidate events will be reviewed for each HAI per facility.

For MRSA bacteremia and CDI LabID Event validation, candidate events are defined by a positive laboratory test. Sixty (60) episodes of care will be selected based on presence of one or more qualifying laboratory tests during an episode of inpatient care, and information from the hospital laboratory and ADT system will be reviewed. Twenty (20) episodes of care will be reviewed to identify the FIRST reportable NHSN LabID Event, and 40 episodes of care will be reviewed to determine whether the SELECTED (non-first) laboratory event should have been reported to NHSN. If less than 20 are reported, all events should be reviewed.

Sample structure

- o (Up to) 60 medical records each for ICU CLABSI, ICU CAUTI, COLO, and HYST, including
 - (Up to) 20 reported HAIs
 - (Goal of) 40 non-reported candidate HAIs. For ICU CLABSIs, these will be stratified by NICU
 and adult/pediatric ICU locations, and will prioritize targeted pathogens. For CLABSI and
 CAUTI, many of these will be eliminated early because they do not have a device (central
 line or urinary catheter). For COLO and HYST, the medical record at the time of the surgical
 procedure will be reviewed, as well as any additional records during the surveillance
 window.
- (Goal of) 60 episodes of care each for <u>candidate MRSA bacteremia LabID Events and candidate CDI LabID Events</u>, including
 - (Up to) 20 "first" positive laboratory tests of the episode of care
 - (Up to) 40 "non-first" positive laboratory tests of the episode of care

Line listings required from facility

To identify unreported "candidate" CLABSI, CAUTI, MRSA bacteremia LabID Events and CDI LabID Events, a sampling frame of medical records and/or positive laboratory tests is needed, and will require assistance from the facility being validated before the audit (see table below and specific instructions for medical records selection in <u>Appendix 1.3: Step-by-Step Targeted Medical Record Selection</u>). For COLO



and HYST SSIs the required sampling frame is derived from COLO and HYST procedures already entered and available in NHSN, however completeness of surgical risk-adjustment variables should be assured before sampling is conducted, because these variables are used for targeting.

Line Listings Required from Facilities for Sampling of CLABSI, CAUTI, MRSA Bacteremia and CDI LabID Events

HAI Event to be	Request to Facility for Line Listing	Line Listing Will Define the Following
Validated	(detailed in Chapter 3)	Sampling Frame Elements
ICU CLABSI	Line listing of positive ICU and NICU	Episodes of care (identified by patient ID
	blood cultures, with patient ID and	and unique admission date) with one or
	admission date	more positive ICU blood culture(s)
		(include NICUs)
ICU CAUTI	Line listing of positive ICU (non-NICU)	Episodes of care (identified by patient ID
	urine cultures ^a with patient ID and	and unique admission date) with one or
	admission date	more positive ICU urine culture(s) ^a
		(exclude NICUs)
MRSA bacteremia	Inpatient ^b blood cultures positive for	Episodes of care with one or more
LabID Event	MRSA	inpatient ^b blood cultures positive for
		MRSA
CDI LabID Event	Inpatient ^b stools ^c toxin-positive for <i>C</i> .	Episodes of care with one or more
	difficile, excluding those from baby	inpatient ^b stools ^c toxin-positive for <i>C</i> .
	locations ^d	difficile, excluding those from baby
		locations ^d

^aPositive ICU urine cultures with no more than 2 identified pathogens and at least 10⁵ CFU/ml organisms



^bFor LabID Event, emergency department (ED) and 24 hour observation location specimens are considered facwideIN. Specimens collected from other affiliated outpatient locations on the day of admission are considered inpatient specimens.

^cSurveillance guidance for laboratories recommends that *C. difficile* toxin testing be done only on unformed stool specimens, and formed stool should be rejected

^dBaby locations include those with 80% or more infants (≤1 year); typically NICU, newborn nursery, and special care nursery. Babies in LDRP locations should also be excluded.

Chapter 5: Activities During and After the Facility Site Visit

Suggested Tools to bring along for validation site-visits

- Letter of introduction, state ID badge or other authorization
- 2015 NHSN Manual
 - Before visit: Tag/highlight case definitions
 - Tag/highlight location descriptions for patient location mapping
- Information about the facility:
 - o Facility's most recent NHSN Annual Survey
 - List of surveillance locations with demographics
 - o List of medical records requested for screening
 - Confidential list of HAIs reported by facility to NHSN (assure that validators are blinded until after review is completed).
- Copies of Methods Surveys (<u>Appendix 2</u>) and form to collect contact information (<u>Appendix 2.3</u>)
- Multiple copies of blank medical record abstraction tools (Appendix 3)
- Copies of 2015 Tennessee checklists (available at http://health.state.tn.us/ceds/hai/)
- Blank audit discrepancies reports (<u>Appendix 4.1</u>)
- External Validation Documentation Form (Appendix 4.3)
- Miscellaneous tools: Straight edge (e.g.: ruler) for reading data printouts, stapler, binder clips, pens, highlighters, sticky notes, tape flags

Please note that some of the listed tools are templates that should be adapted to the facility and state before copies are made.

Request documentation of current NHSN reporter training

NHSN reporters should have documentation of successful completion of the online, self-paced multimedia training modules for HAIs they oversee. This is an opportunity to establish or reinforce state expectations for this annual update. Consider recording the results in Appendix 4.3, custom field.

Review risk adjustment variables:

For CLABSI and CAUTI, review ICU location mapping, location bed size, and teaching hospital status. For MRSA bacteremia and CDI LabID Event reporting, review location mapping facility-wide if this has not been done to the state's satisfaction in the past 3 years. Otherwise, review changes since the last facility-wide review.

Bring a copy of the facility NHSN Annual Survey, and review the ICU location mapping and bed size information with the IP, along with an up-to-date list of CDC locations and descriptions (see http://www.cdc.gov/nhsn/forms/instr/57 103-TOI.pdf and http://www.cdc.gov/nhsn/PDFs/pscManual/validation/pcsManual-2015-valid.pdf). If there is insufficient time to complete this onsite, consider arranging a conference call to review location mapping when data are readily accessible.

Review NHSN definitions for teaching hospital types (under Key Terms, http://www.cdc.gov/nhsn/PDFs/pscManual/validation/pcsManual-2015-valid.pdf), and assure that facility teaching hospital status is accurate in the NHSN Annual Survey.



For COLO and HYST, many risk adjustment variables can be validated as part of the medical record review process. The medical record abstraction forms for COLO and HYST include fields for ASA score, patient age, and other risk adjustment variables, as well as SSI outcome. Validation of risk adjustment variables is recommended to assure that sampling has appropriately targeted high-risk procedures.

Review denominator methods and documentation

CLABSI and CAUTI denominator counting methods

Surveillance and denominator data collection surveys found in <u>Appendices 2.1</u> and <u>2.4</u> may be administered to the IP contact before or during the site visit; however it may be impractical to interview multiple denominator data collectors during the site visit. In this case, collecting contact information during the site visit may be advisable for subsequent administration of surveys by telephone (<u>Appendix 2.3</u>). This allows time at the facility to be used efficiently and accommodates interviews with individuals who may work at other times (e.g. the night shift).

In many facilities, the same person will collect denominator data for device-associated infections (including CLABSI and CAUTI) concurrently. Because of this, the denominator counting survey for CLABSI and CAUTI in Appendix 2.4 may be administered for each metric separately or for both combined. Knowledge of definitions and counting methods is important even in facilities where denominators are reported electronically in order that spot-checks can be conducted periodically. A form for facilities to document required internal validation of electronic denominator counting is provided in Appendix 2.2.

Facilities may have already administered denominator counting surveys for internal validation purposes. If this is the case, validators may choose to accept their evidence or conduct this survey among a more limited sample of denominator counters.

CLABSI and **CAUTI** denominator records

While visiting, request original records of denominator data collection paperwork, which can provide insight into the frequency, reliability, and consistency of this task and how omissions are handled (NHSN provided guidance for missing device-associated denominator data in September 2013 http://www.cdc.gov/nhsn/PDFs/NHSNMissingDenomData_Sep2013.pdf). Consider whether patient days and central-line days data appear as anticipated when manually counted each day: different ink, different but similar numbers. Determine for what percent of days data are missing and what was done for reporting on those days. Findings should be documented in Appendix 4.3.

Electronically collected CLABSI and CAUTI denominators

Unexamined electronic denominator counting may be a source of error in HAI reporting.^{5,6} If the facility uses electronic denominator data collection, obtain documentation of their denominator validation process and any periodic spot checks. NHSN specifies that electronic denominator counts should fall within 5% of manual counts for three consecutive months before electronic counts can be used (See <u>Appendix 2.2</u>).

If documentation of electronic denominator validation is not available, the facility should resume manual counting (and assure staff training), to re-validate electronic counts, and to retain evidence of valid electronic counting (within 5% for 3 months). Facilities should conduct periodic spot checks even after formal validation to prevent lost information due to changing medical records systems or other



disruptions. Accurate electronic denominator reporting may require iterative programming corrections in consultation with IT support until accuracy is established.^{7,8}

Completeness and accuracy of SSI (COLO and HYST) denominators

Evaluate the information in Appendix 4.3, "Denominator Validation COLO" and "Denominator Validation HYST" (this information was gathered during preparation for the facility site visit). If there appear to be large differences in the number of procedures identified by these two data sources, discuss this with the IP. Consider matching a subset of records between the two systems and examining un-matched records to explore potential reasons for this discrepancy. In particular beginning in 2013, all procedures meeting the NHSN procedure definition should be entered, regardless of pre-existing infection / wound class or incision closure method. If the two systems generate roughly similar data, the NHSN procedure denominator should be considered complete.

Electronically collected MRSA bacteremia and CDI facility-wide inpatient (FacWideIN) denominators

"FacWideIN" surveillance data includes all patient days counted at the same time each day for all inpatient locations, including any patients housed for the day in inpatient locations, whether or not the facility considers them "admitted patients" or "observation" patients, but excluding any patients housed for the day in outpatient "observation" locations. This information is often collected electronically. Because the task of validating "FacWideIN" patient days and admissions is daunting, denominator data validation can be accomplished using manual counting of patient days and admissions in three specified location types for one month each: one ICU, one Labor/Delivery/Recovery/Post-Partum (LDRP) location (if available), and one or more wards where "observation" patients are frequently located. Manual counts should be within 5% of the referent (usual) electronic counts, or an evaluation of why they differ should be conducted. One consideration is the facility's ability to capture "observation" patients within inpatient locations electronically. Electronic ADT data often are found to be more accurate than electronic billing data in this regard. Note that patient counts should differ for MRSA bacteremia LabID Event and CDI LabID Event denominators because CDI denominators exclude infants (< 1 year old). This internal validation process can be conducted by facilities when requested or required.

Structured Medical Records Review

Validator blinding and consultation at the facility site-visit

Validator blinding as to HAI status is required and is normally accomplished by mixing and reviewing the selected medical records before determining which have been reported to NHSN with HAIs.

Medical records should be reviewed in a blinded manner using 2015 Medical Records Abstraction Tool processes (Appendix 3). These tools include algorithms and logic designed to establish presence or absence of required criteria for case definitions and to provide support to avoid common errors.

For CLABSI validation, when consideration is given to an alternative primary site infection leading to secondary bloodstream infection, use of an appropriate Tennessee checklist (available at http://health.state.tn.us/ceds/hai/) is highly recommended. These checklists provide a structure to record required elements from the NHSN Manual's Chapter 17 criteria. The Tennessee checklists are also useful for surgical site infection (SSI) validation when documenting organ/space SSIs. The



checklists exist for multiple infection types (derived from the NHSN manual Chapter 17), and in multiple dated versions. Be sure the selected version is for 2015 definitions.

If working on paper, bring enough copies of the medical records abstraction tools to complete a separate form for each medical record. After all medical records have been abstracted by validators, events reported to NHSN should be revealed and a meeting arranged with IPs / NHSN reporters to discuss any discrepancies between validator outcomes and reported outcomes, while medical records are readily available.

Discussion of audit results with IP

Whether or not reporting errors are identified, review the data with the IP to assure transparency and provide opportunity for discussion and feedback. If case-determinations are discordant, determine whether reporters or auditors missed any documented information that would affect the correct result (undocumented information should not be considered). Use NHSN criteria as the gold standard. For difficult cases, seek adjudication from CDC.

Look carefully for systematic reporting errors or misconceptions that could affect reporting beyond the reviewed medical records. If systematic errors are found, the facility should be asked to rereview and correct affected data, not just those records reviewed by auditors. These errors should be re-assessed during the next audit to evaluate improvement.

Use errors as learning opportunities for reporters and validators. These discussions may provide insight into the soundness of the facility's surveillance processes and competencies, and topics where additional training may be useful. Leave a copy of expected changes to NHSN data with the IP and agree to a deadline for changes to be made (see Appendix 4.1). An exit interview with a facility C-suite administrator (e.g., CEO or CMO) would rarely be needed, unless a process improvement plan is indicated.

Post-visit

Denominator data collection surveys (Appendix 2.4) may be completed after the visit.

Document validation findings (e.g., using Appendix 4.3) to create a facility summary report.

A follow-up letter to the IP and facility C-suite administrator will close the communication loop and provide valuable feedback. Send a letter thanking them, recognizing all participants in the audit, and documenting results, necessary corrections, and recommendations. When appropriate, identify systematic strengths as well as problems with resources and support for surveillance, data collection, and reporting (Appendix 4.2).

If the facility was required to change data in NHSN or to re-review information due to systematic errors, follow-up with the facility and assure corrections are made by the agreed upon deadline.



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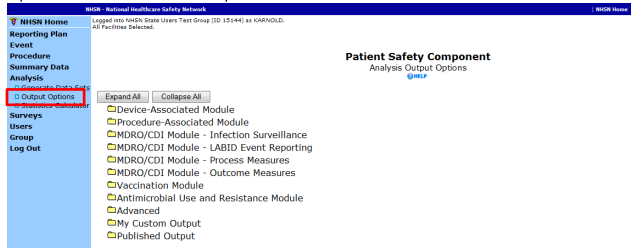
Appendix 1: Preparation Tools for External Validation

Appendix 1.1: Step-by-Step Targeted Facility Ranking

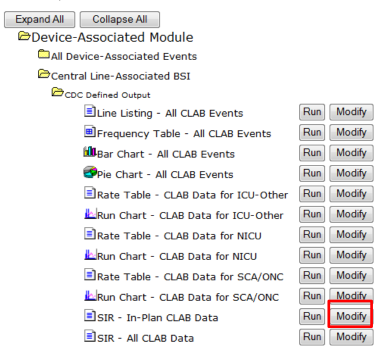
1. Generate new datasets in NHSN to ensure any data updates are included for analysis. On the NHSN Landing Page, navigate to Patient Safety Component -->[YOUR State Users' Group]. Select the "Analysis" tab and click "Generate Datasets." Click the Generate New button. Allow the dataset generation process to complete; you are able to leave NHSN during the generation process.



2. After successful dataset generation, navigate to Analysis → Output Options to display the tree view list of all analysis reports available within NHSN's analysis tool.



3. Use the tree view structure to navigate to the SIR report of interest. In this example (targeting for ICU CLABSI validation), we will select the Device Associated Module, -> Central Line-Associated BSI, -> CDC Defined Output, -> SIR for In-Plan CLABSI data. This uses data reported to NHSN that has been shared with the group. Click the Modify button to proceed to the modification screen, which can be used to filter and export data from NHSN.



- 4. A modification screen will open titled "Analysis SIR." On the modification screen, there are two key areas to modify, one that controls the time interval of data that are analyzed and displayed and one that controls the level of aggregation of that data.
 - a. Use the "Select a time period" option to limit the time period of data that is included in the report to be exported. Set "Date Variable" to SummaryYr, "Beginning" to 2015 and "Ending" to 2015:

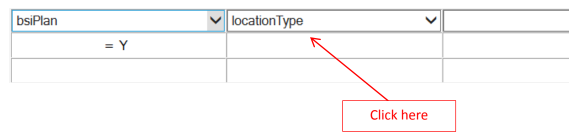
Select a time period or Leave Blank for Cumulative Time Period: OHELP



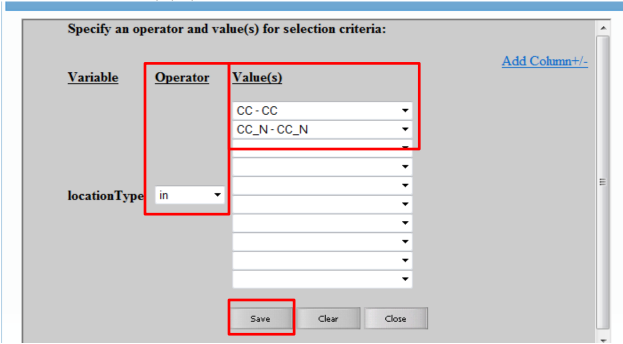
b. Scroll down to "Specify Other Selection Criteria." Retain bsiPlan = Y. In column 2, select "Location type" from the dropdown list, Click in the space below "location Type."

Specify Other Selection Criteria: OHELP

<u>Show Criteria</u> <u>Column + Row + Clear Criteria</u>



c. Doing this will pop up a new window (shown below in gray). Set "Operator" = "in" and Value(s) to "CC-CC" and "CC_N-CC_N" to specify all ICU locations, adult and neonatal. Scroll to the bottom of the pop-up screen and select "Save" to close the pop-up:



d. Under the "Other Options" section, use the "Group by" option to view the data at a particular level of aggregation. By default, this is set to SummaryYH, (half-years). Change the Group by option to "SummaryYr".



- 5. After making these modifications, scroll to the bottom of the modification screen. Click the Export Output Data Set button to export the data selected by your modifications to a different file format.
- 6. Clicking the Export Output Data Set button will take you to the Export Output Options screen. Use the dropdown menu to select the file format to export the data. In this example, we will export to an Excel spreadsheet (*.xls). Click the Export button to begin the export process. NHSN will create a .zip file with your data export in it and prompt you to specify a location to save the file on your computer.

Export Output Options

Excel spreadsheet (*.xls)	•		
		Export	Back

For CLABSI data, the exported SIR report file will be displayed at multiple levels of aggregation, which are outlined and displayed in the screenshot to follow.

8. In Excel, <u>select the aggregation level that provides a facility-specific SIR for all ICUs</u> (shown below in black). This level of aggregation will allow you to explore the level of exposure risk for ICU CLABSIs and measured performance at each facility. Select these rows and copy this information to a new spreadsheet. (Also, insert a row above your data

and copy the header row so you can identify the variables on the new page). Arrange the facilities in rank order according to "exposure;" the expected/predicted number of CLABSIs [numExp], (high to low), and create three new columns titled "Delta count", "Stratum", and "Targeted Selection Number".

- 9. Use Excel to calculate the Delta count for each facility/row. The formula in Excel is (=ABS[row cell under InfCount]— [row cell under numExp]). (You will use Delta count only if an SIR is not calculated by NHSN).
- 10. Select the top tertile (33%) of facilities by predicted number of ICU CLABSIs. This "Top Tertile" of facilities where ICU CLABSIs are most expected, may have the greatest potential for surveillance and prevention impact.
- 11. Within the top tertile, sort by SIR from highest to lowest, and identify the current median SIR for the top Tertile. (Recall that median is the "middle" value for the group). To sort just the top tertile, highlight the entire row for each facility in the top tertile, and click "Data, "Sort"; Column "Sort by" (select SIR), "Sort On" (values), and "Order" (highest to lowest).
- 12. Within the top tertile, assign stratum A to facilities with SIR above the current median SIR, stratum B for remaining facilities with SIR less than or equal to the median and above zero, and stratum C for facilities with SIR = zero (but not missing). Note that some facilities will not have a calculated SIR; do not include these in the strata (see step 15 below).
- 13. Re-sort within each stratum A, B, and C, by numExp from highest to lowest. To sort just one stratum at a time, highlight the entire row for each facility in the first stratum, and click "Data," "Sort;" Column "Sort by" (select numExp), "Sort On" (values), and "Order" (highest to lowest). Repeat this process for the next two strata, one-by-one.
- 14. Assign sequential Targeted Selection Numbers to facilities, by selecting the highest available numExp from each stratum alternating A, B, and C. For example, facility #1 will be the facility with the highest numExp from stratum A, facility#2 the facility with the highest numExp from stratum B, and #3 the facility with the highest numExp from stratum C. Return to stratum A and assign#4) to the next facility in stratum A, assign #5 to the next facility in stratum B, and facility #6 will be the next facility in stratum C. Continue alternating strata until no facilities remain or the target number of facilities (18 or 21) is reached. If additional facilities are needed, repeat this process (steps 11-14) using the second and then third tertile based on exposure.
- 15. Once all hospitals with an assigned SIR have been prioritized, evaluate facilities with fewer expected events. In hospitals where NHSN does not calculate an SIR (because the predicted number of infections is less than one), a different method rather than the above method of stratifying by SIR should be used. This is because the value of a calculated SIR is exceedingly imprecise when the expected number of infections is less than one, and a single infection can result in a very high SIR. If additional facilities are needed to complete the targeted number, prioritize them based on the highest and descending delta count (only for facilities without a calculated SIR).
- 16. After the targeted selection is complete, ALL remaining facilities from ALL tertiles will be subject to random selection under the 5% rule.
- 17. This basic process can be followed with minor modifications for each of the six HAI metrics, to identify facilities that are highly exposed (and therefore at risk for HAIs) and to characterize their performance using the SIR to rank them for validation.

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3	5	2344	4.012	1.246	0.3735	0.405, 2.908	1/1/2014	ICU-OTHER	SIR for all adult/ped	diatric IC	CUs in all fac	cilities in g	roup		
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17	1	10	0.019				1/1/2014	ICU-OTHER		17775					
18	0	10	0.02				1/1/2014		IN:ACUTE:CC:C	10000					
19	2	10	0.019				1/1/2014		IN:ACUTE:CC:M	10000					
20	0	368	0.552				1/1/2014		IN:ACUTE:CC:MS	10000					
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25	1	2	0.006				1/1/2014		IN:ACUTE:CC:NURS	15164					
26	0	10	0.023				1/1/2014		IN:ACUTE:CC:S	15164					
27	1	10	0.019				1/1/2014		IN:ACUTE:CC:M	17775					
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Targeted Facility Ranking for ICU CAUTI:

Note: See "Step-by-Step Targeted Facility Ranking Method, using ICU CLABSI" as an example; a similar process will be used for ranking of facilities for ICU CAUTI, with the following exceptions:

Follow parts 1 and 2, as shown above.

In part 3, select the Device Associated Module, -> Urinary Catheter-Associated UTI, -> CDC-defined Output, -> SIR—In-Plan CAUTI Data. Select the modify button to proceed to the modification screen as before.

Follow part 4a, as shown above.

In part 4b, scroll down to "Specify Other Selection Criteria. Retain utiPlan=Y. In column 2, select location Type and click in the space below to pop up the new selection window. Set "Operator" to "=" and Value(s) to "CC-CC". (Omit "CC_N-CC_N", because you do not want to include NICU locations in the exposure calculations for CAUTI). Scroll to the bottom of the gray pop-up and select "SAVE".

The selection box should resemble the screen shot below.

Specify Other Selection Criteria: QHELP

Show Criteria Column + Row + Clear Criteria

utiPlan ▼	locationType ▼	
= Y	= CC	

Follow steps 4d, 5, 6, and 7 as shown above. The exported SIR report Excel file will be displayed with multiple aggregation levels similar to the CLABSI data shown above.

In Part 8, using Excel, <u>select the aggregation level that provides a facility-specific SIR for all ICUs</u>. This level of aggregation will allow you to explore the level of exposure risk for ICU CAUTIs and measured performance at each facility. Copy this information to a new spreadsheet. Arrange the facilities in rank order according to "exposure"; the expected/predicted number of CAUTIs [numExp], (high to low), and create three new columns titled "Delta count," "Stratum," and "Targeted Selection Number."

Complete steps 9-16 to assign a sequential Targeted Selection Number for ICU CAUTI to facilities and to draw a 5% random sample as before.

Targeted Facility Ranking for COLO:

Note: Targeting surgical procedures requires that risk-adjustment variables in NHSN are complete. Please work with facilities to assure acceptable data quality and completeness before attempting to select facilities and records.

Note: See "Step-by-Step Targeted Facility Ranking Method, using ICU CLABSI" as an example; a similar process will be used for ranking of facilities for COLO validation, with the following exceptions:

Follow parts 1 and 2, as shown above.

In part 3, select the Procedure-Associated Module, ->SSI, ->CDC-defined Output, ->SIR-In-Plan All SSI Data by Procedure. Select the modify button to proceed to the modification screen as before.

Follow part 4a, as shown above.

In part 4b, scroll down to "Specify Other Selection Criteria. Retain ssiPlan = Y. In column 2, select procCODE and click in the space below to pop up the new selection window. Set "Operator" to "=" and Value(s) to "COLO." Select "SAVE."

The selection box should resemble the screen shot below:

Specify Other Selection Criteria: OHELP

Show Criteria Column + Row + Clear Criteria

ssiPlan ▼	procCode ▼	
= Y	= COLO	

Follow steps 4d, 5, 6, and 7 as shown above. The exported SIR report Excel file will be displayed with multiple aggregation levels. A screen shot of an Excel spreadsheet is provided below to illustrate:

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	C16	▼ (=	f _x									
	А	В	С	D	Е	F	G	Н	I	J	K	L
1	summaryYr	procCount	infCountAll	numExpAll	SIRAII	SIRAII_pval	SIRAII95CI	procCode	orgid	outpatient	months	
2	1/1/2014	39	3	1.77	1.695	0.2614	0.350, 4.953	SIR for all f	acilities,	all procedui	res	
3	1/1/2014	39	3	1.77	1.695	0.2614	0.350, 4.953	COLO	SIR for	all faciilties,	specific	procedures
4	1/1/2014	37	2	1.638	1.221	0.4873	0.148, 4.411		10000			
5	1/1/2014	2	1	0.132					15164	SIR for each	facility,	all procedures
6	1/1/2014	37	2	1.638	1.221	0.4873	0.148, 4.411	COLO	10000	*THIS IS TH	E LEVEL 1	O EVALUATE*
7	1/1/2014	2	1	0.132				COLO	15164	SIR for each	facility,	specific procedures
8	1/1/2014	37	2	1.638	1.221	0.4873	0.148, 4.411	COLO	10000	N	6	
9	1/1/2014	2	1	0.132				COLO	15164	N	2	
4.0												

In Part 8, Using Excel, select the aggregation level that provides a facility-specific SIR for COLO SSIs (shown in black in the above screenshot). This level of aggregation will allow you to explore the level of exposure risk for COLO SSIs and measured performance at each facility. Copy this information to a new spreadsheet. Arrange the facilities in rank order according to "exposure"; the expected/predicted number of SSIs [numExp], (high to low), and create three new columns titled "Deltacount," "Stratum," and "Targeted Selection Number."

Complete steps 9-16 to assign a sequential Targeted Selection Number for COLO SSI to facilities and to draw a 5% random sample as before.

Targeted Facility Ranking for HYST:

Note: Targeting surgical procedures requires that risk-adjustment variables in NHSN are complete. Please work with facilities to assure acceptable data quality and completeness before attempting to select facilities and records.

Note: See "Step-by-Step Targeted Facility Ranking Method, using ICU CLABSI" as an example; a similar process will be used for ranking of facilities for HYST validation, with the following exceptions:

Follow parts 1 and 2, as shown above.

In part 3, select the Procedure-Associated Module, ->SSI, ->CDC-defined Output, ->SIR-In-Plan All SSI Data by Procedure. Select the modify button to proceed to the modification screen as before.

Follow part 4a, as shown above.

In part 4b, scroll down to "Specify Other Selection Criteria. Retain ssiPlan = Y. In column 2, select procCODE and click in the space below to pop up the new selection window. Set "Operator" to "=" and Value(s) to "HYST." Select "SAVE."

The selection box should resemble the screen shot below.

Specify Other Selection Criteria: QHELP

<u>Show Criteria</u> <u>Column + Row + Clear Criteria</u>

ssiPlan ▼	procCode ▼	
= Y	= HYST	

Follow steps 4d, 5, 6, and 7 as shown above. The exported SIR report Excel file will be displayed with multiple aggregation levels similar to the COLO data spreadsheet shown above.

In Part 8, Using Excel, <u>select the aggregation level that provides a facility-specific SIR for HYST SSIs</u>. This level of aggregation will allow you to explore the level of exposure risk for HYST SSIs and measured performance at each facility. Copy this information to a new spreadsheet. Arrange the facilities in rank order according to "exposure"; the expected/predicted number of SSIs [numExp], (high to low), and create three new columns titled "Delta count," "Stratum," and "Targeted Selection Number."

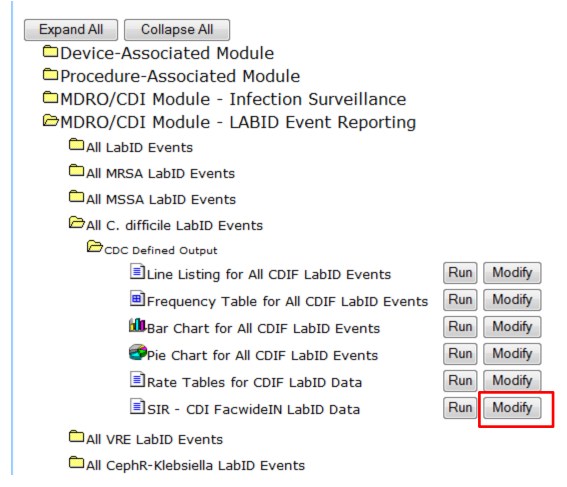
Complete steps 9-16 to assign a sequential Targeted Selection Number for HYST SSI to facilities and to draw a 5% random sample as before.

Targeted Facility Ranking for CDI LabID Event:

Note: See "Step-by-Step Targeted Facility Ranking Method, using ICU CLABSI" as an example; a similar process will be used for ranking of facilities for CDI LabID Event, with the following exceptions:

Follow parts 1 and 2, as shown above.

In part 3, select the MDRO/CDI Module-LabID Event Reporting, -> All C. difficile LabID events, -> CDC-defined Output, -> SIR-CDI FacwideIN LabID Data. Select the modify button as shown in the screen shot below.



Follow part 4a, as was shown for ICU CLABSI.

In part 4b, modify the selection criteria grid to analyze only IN-PLAN, FacWideIN data. Click the first box in the top row, and select the variable "cdifLabIDPlan"; click on the empty cell directly below this variable, to open a gray pop-up box, and type "Y" in "Value(s)" cell. Click SAVE. Next, click in the second box in the top row, and select the variable "location"; click on the empty cell directly below this variable to open another gray pop-up box. Select location = FACWIDEIN, and SAVE. See screen shot below.

Specify Other Selection Criteria: ②HELP Show Criteria Column + Row + Clear Criteria cdifLabIDPlan ▼ location ▼ = Y = FACWIDEIN

Follow steps 4d, 5 and 6, as shown for ICU CLABSI.

For part 7, the exported SIR report Excel file will be displayed at several levels, as illustrated in the screenshot below:

Fi	e Home	Insert	Page I	Layout Formulas I	Data Review	View									♡ ?
	017	▼ (f _x											
	А	В	С	D	Е	F	G	Н		J	K	L	М	N	0
1	summaryYr	_TYPE_	_FREQ_	CDIF_facIncHOCount	numExpCDI	numpatdays	sir_l	sir_u	SIR_pval	SIR	spcOrgType	orgID			
2	1/1/2014	0	175	9941	8981.348	11944462	1.085	1.129	0	1.107	SIR for all fa	cilities	in the grou	р	
3	1/1/2014			322	235.998	254200	1.219	1.522	0	1.364	CDIF	100			
4	1/1/2014			8	57.016	68795	0.061	0.276	0	0.14	CDIF	101			
5	1/1/2014			135	95.419	158536	1.186	1.675	0.0001	1.415	CDIF	102			
6	1/1/2014			183	168.317	183307	0.935	1.257	0.1376	1.087	CDIF	103			
7	1/1/2014			61	129.912	186976	0.359	0.603	0	0.47	CDIF	104			
8	1/1/2014			59	67.143	83769	0.669	1.134	0.176	0.879	CDIF	105			
9	1/1/2014			61	104.33	151973	0.447	0.751	0	0.585	CDIF	106			
10	1/1/2014			39	65.961	99509	0.42	0.808	0.0002	0.591	CDIF	107	*THIS THE	LEVEL TO I	NVESTIGATE*
11	1/1/2014			127	116.421	124068	0.909	1.298	0.1745	1.091	CDIF	108	SIR for eac	h facility in	the group
12	1/1/2014			140	156.752	229709	0.751	1.054	0.0954	0.893	CDIF	109			
13	1/1/2014			91	40.829	68914	1.794	2.737	0	2.229	CDIF	110			
14	1/1/2014			6	54.44	81964	0.04	0.24	0	0.11	CDIF	111			
15	1/1/2014			144	134.459	168483	0.903	1.261	0.2161	1.071	CDIF	112			
16	1/1/2014			38	63.655	95871	0.422	0.819	0.0004	0.597	CDIF	113			
17	1/1/2014			52	64.913	76570	0.598	1.051	0.0579	0.801	CDIF	114			
18	1/1/2014			13	30.273	49980	0.229	0.734	0.0003	0.429	CDIF	115			
19	1/1/2014			29	72.694	107924	0.267	0.573	0	0.399	CDIF	116			
20	1/1/2014			57	80.046	115823	0.539	0.923	0.0042	0.712	CDIF	117			

In Part 8, Using Excel, select the aggregation level that provides a facility-specific SIR for CDI LabID Event (shown in black in the above screenshot). This level of aggregation will allow you to explore the level of exposure risk for LabID Event and measured performance at each facility. Copy this information to a new spreadsheet. Arrange the facilities in rank order according to "exposure"; the expected/predicted number of LabID Events [numExpCDI], (high to low), and create three new columns titled "Delta count," "Stratum," and "Targeted Selection Number.-"

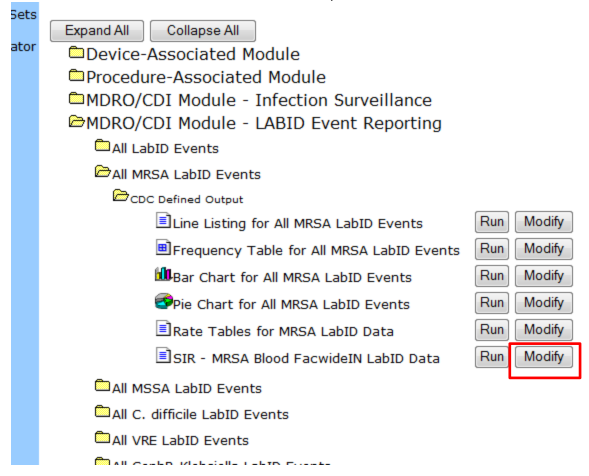
Complete steps 9-16 to assign a sequential Targeted Selection Number for LabID Events to facilities and to draw a 5% random sample as before.

Targeted Facility Ranking for MRSA Bacteremia LabID Event:

Note: See "Step-by-Step Targeted Facility Ranking Method, using ICU CLABSI" as an example; a similar process will be used for ranking of facilities for MRSA Bacteremia LabID Event, with the following exceptions:

Follow parts 1 and 2, as shown above.

In part 3, select the MDRO/CDI Module-LabID Event Reporting, -> All MRSA LabID events, -> CDC-defined Output, -> SIR-MRSA Blood FacwideIN LabID Data. Select the modify button as shown in the screen shot below.



Follow part 4a, as shown for ICU CLABSI above.

In part 4b, modify the selection criteria grid to analyze only IN-PLAN data. Click the first box in the top row, and select the variable "mrsaLabIDBIdPlan"; click on the empty cell directly below this variable, to open a gray pop-up box, and type "Y" in "Value(s)" cell. Click SAVE.

NOTE: facilities that are conducting IN-PLAN MRSA all specimen surveillance are ALSO conducting IN-PLAN MRSA Bacteremia surveillance as a subset. NHSN includes these facilities under "mrsaLabIDBIdPlan'=Y. Any surveillance that is not IN-PLAN will be excluded. Next, click in the second box in the top row, and select the variable "location"; click on the empty cell directly below this variable to open another gray pop-up box. Select location = FACWIDEIN, and SAVE. See screen shot below.

Specify Other Selection Criteria: OHELP

Show Criteria Column + Row + Clear Criteria

mrsaLablDBldPlan •	location •	
= Y	= FACWIDEIN	

Follow steps 4d, 5 and 6, as shown for ICU CLABSI above.

For part 7, the exported SIR report Excel file for MRSA Bacteremia LabID Event will be displayed at several levels, and should look similar to the screenshot (for CDI LabID Event FACWIDEIN) shown above.

In Part 8, Using Excel, <u>select the aggregation level that provides a facility-specific SIR for MRSA Bacteremia LabID Event</u>. This level of aggregation will allow you to explore the level of exposure risk for LabID Event and measured performance at each facility. Copy this information to a new spreadsheet. Arrange the facilities in rank order according to "exposure"; the expected/predicted number of LabID Events [numExpMRSA], (high to low), and create three new columns titled "Delta count," "Stratum," and "Targeted Selection Number."

Complete steps 9-16 to assign a sequential Targeted Selection Number for LabID Events to facilities and to draw a 5% random sample as before.

Appendix 1.2: Sample Letter Requesting Site Visit and Line Listings for External Validation

Please customize this template to meet your state's needs

Dear [Name of CEO]

Cc: [Name of IP]

The [Health Department] will conduct an audit of surveillance practices and reporting of healthcare-associated infections in [multiple/all] hospitals statewide, focusing on 6 different metrics for 2015 data. These include the metrics designated by the CMS Inpatient Quality Reporting Program: central line-associated bloodstream infections (CLABSI) and catheter-associated urinary tract infections (CAUTI) in ICUs, surgical site infections (SSI) following colon (COLO) and abdominal hysterectomy (HYST) procedures, and proxy measures for MRSA bacteremia (MRSA bacteremia LabID Event) and Clostridium difficile infection (CDI LabID Event). [Modify metrics as indicated] Participation in the audit is

[select as appropriate]

- [obligatory, to assure compliance with state healthcare-associated infection (HAI) reporting legislation and assure that facilities are accurately identifying and reporting healthcare-associated infections]. OR
- [voluntary, but may be of value to you in preparation for CMS validation activities, and by assuring
 that all state facilities are held to a high standard of accountability]. [Facilities that participate will be
 acknowledged by the SHD in the following way_________. Facilities that choose not to
 participate will also be identified in the following way_______.]
- [Modify as per state decision]: The individual results of SHD validation will be shared with your infection prevention staff and you [but will / will not be shared in the following additional ways]. Pooled results of SHD validation will be shared publically, but will not identify individual facilities.

A site visit has been tentatively scheduled for [Day and Date] with [Name of IP], Infection Preventionist, who has also been asked to assist with generating 4 line listings (described below) of eligible medical records for review, and two reports of monthly surgical procedures. Successful preparation for the audit will require the assistance of the microbiology laboratory, medical records system, and IT to generate specified line listings ahead of time that will be used to select medical records for review, and later assistance from medical records personnel to make medical records available for review at the time of the audit.

At this time, we request your support for production of the following 4 microbiology laboratory-based line listings, coordinated through the IP, and transmitted to us securely via FTP [FTP site] in a spreadsheet (e.g. Excel) file format. Please note that these lists must include information about facility admission date, which may require coordination of microbiology data with another hospital data system. The line listings will be due by [Date]. If questions arise, we can be reached at the following number [XXX-XXX-XXXX]:

Requested Line Listings

1) A complete list of positive ICU blood cultures for 2015, with additional variables based on the template below. NICUs should be included.

Template positive ICU blood culture line listing (* indicates required data):

*MRN	*Facility	*Laboratory	*Specimen	*Blood	*Specific	*Gender	*Date	First	Last
	Admission	Specimen	Collection	Organism 1	ICU		of	Name	Name
	Date	Number	Date	Genus and	patient		Birth		
				Species	Location				

2) A complete list of positive ICU urine cultures for 2015, with additional variables based on the template below. NICUs <u>should not</u> be included. If possible, limit positive ICU urine cultures to those with no more than 2 identified pathogens and at least 10³ CFU/ml organisms.

Template positive ICU urine culture line listing (* indicates required data, †indicates conditionally required data):

*MRN	*Facility	*Laboratory	*Specimen	*Urine	*Urine	†Urine	†Urine	*Specific	*Gender	*Date	First	Last
	Admission	Specimen	Collection	Organism 1	Colony	Organism 2	Colony	ICU		of	Name	Name
	Date	Number	Date	Genus and	Count 1	Genus and	Count 2	Location		Birth		
				Species	(CFU/ml)	Species	(CFU/ml)					

3) A complete list of blood cultures positive for methicillin-resistant *Staphylococcus aureus* (MRSA), among inpatients facility wide for 2015, with additional variables based on the template below.

Template positive MRSA bacteremia, FacWideIN line listing (* indicates required data):

*	'MRN	*Facility	*Laboratory	*Specimen	*Blood Organism	*Documentation	*Specific	*Gender	*Date of	First	Last
		Admission	Specimen	Collection	Genus and	of Methicillin-	Mapped NHSN		Birth	Name	Name
		Date	Number	Date	Species	Resistance	Location				
					(documenting S.	(susceptibility test					
					aureus or MRSA)	result or MRSA)					

4) A complete list of toxin-positive *Clostridium difficile* stool specimens among inpatients facility-wide for 2015, with additional variables based on the template below. Please include only final results for toxin testing that is conducted following multiple steps.

Template positive C. difficile assay FacWideIN line listing (* indicates required data):

*MRN	*Facility	*Laboratory	*Specimen	*Result of	* Specific	*Gender	*Date	First	Last
	Admission	Specimen	Collection	CDI Toxin	Mapped NHSN		of Birth	Name	Name
	Date	Number	Date	Test	Location				

The line listings will be due by [day and date <u>in advance of</u> site visit] so that we may select medical records for review from among candidate records. We will then communicate our selected records to infection prevention so that they can be made available for the audit.

5) In addition, we request a monthly count of selected 2015 inpatient surgical procedures performed in your facility based on the following ICD-9-CM/ICD-10-PCS procedure codes:

Procedure	COLO Procedures	HYST Procedures
Class	Colo i loccuares	Titoccadies
ICD-9-CM/		
ICD-10-		
PCSProcedure		
Codes:		
Codes.		
2015 Month	Number of Procedures	Number of Procedures
January		
February		
March		
April		
May		
June		
July		
August		
September		
October		
November		
December		

During our visit, we will be available to describe the process and evaluation tools, as well as answer any questions you may have about the state health department's HAI data validation program.

If your healthcare facility has initiated or completed conversion to an electronic medical record system, we will need a means of accessing these records during our visit, including any diagnostic/laboratory results, clinical documentation and ICD-9-CM codes related to these patients.

Should there be any scheduling difficulties, please contact me directly, either by phone [phone number] or email [email].

HAI Program Director / Regional Representative

cc: IP name

enc.

Appendix 1.3: Step-by-Step Targeted Medical Record Selection

ICU CLABSI Targeted Medical Record Selection Process (Note: this is the same process recommended for 2013 ICU CLABSI validation)

- 1. From each selected facility, request a securely transmitted line listing of all positive ICU blood cultures, from all ICUs reporting to NHSN, for the entire year, with required additional variables used for medical record identification and matching to NHSN reports (See Chapter 3 for recommended line listing structure).
- 2. Assure the line listing includes positive blood cultures from all ICU locations required to report CLABSIs to NHSN, using location mapping information in NHSN
- 3. Assign a random number to each positive blood culture
- 4. Sort the list of blood cultures by MRN and admission date to generate clusters of blood cultures associated with recognizable patient records
- 5. Identify reported CLABSIs on the blood culture line listing
 - a. Using the NHSN CLABSI list and available patient information on blood culture line listing, flag and mark blood cultures reported as CLABSIs. Create a new variable, "stratum" and assign these blood cultures and all other blood cultures in the same medical record to stratum 1.
 - b. If reported CLABSIs are missing from the blood culture line listing, the list may be incomplete.

 Investigate and correct this problem. Add omitted CLABSI records to the medical record review list.
- 6. Select simple random sample of (up to) 20 reported ICU CLABSIs for review
 - a. Select stratum = 1
 - b. Sort by random number, MRN, and hospital admission date
 - c. Select the first 20 random numbers with unique episodes of care (defined by MRN and admission date) as the sample of reported CLABSI records
- 7. Identify unreported candidate CLABSI events and stratify by targeted pathogens
 - a. Select stratum not equal to 1
 - b. Sort non-stratum 1 blood cultures by pathogen (focusing on Organism 1 only)
 - i. If the organism (Org 1) is a "Targeted Pathogen" (see list below), assign the positive blood culture to stratum 2. If the organism (Org 1) is not a "Targeted Pathogen," assign the positive blood culture to stratum 3.
 - ii. Targeted Pathogens:
 - 1. Candida spp., Torulopsis spp. (yeast)
 - 2. Enterococcus spp.
 - 3. Staphylococcus aureus (includes MRSA, MSSA)
 - 4. Coagulase-negative staphylococcus (includes most staphylococcus spp. other than *S. aureus*, MRSA, MSSA)
 - 5. Klebsiella spp., E. coli, or Pseudomonas spp. (common gram negatives)
- 8. Among unreported candidate CLABSI events, use location information to identify NICU vs. adult/pediatric ICU records (If facility has no NICU, skip to step 10 below, and select 10 additional medical records from adult/pediatric ICUs for screening sample.).
 - a. Re-sort blood cultures by ICU type (NICU vs. adult/pediatric ICU) and create a variable NICU (Yes/No). Assign NICU status to each blood culture as appropriate.
- 9. Select the NICU screening sample
 - a. Select NICU= Yes, and stratum = 2 (targeted pathogens)
 - b. Sort by random number, MRN, and admission date

- c. Select the first 10 random numbers with unique episodes of care (defined by MRN and admission date) as the sample of NICU records containing candidate CLABSIs.
- d. If 10 NICU medical records with stratum 2 blood cultures are not available, supplement the NICU sample with NICU records with stratum 3 blood cultures (where NICU = Yes, and stratum = 3); take the initial medical records (lowest random numbers with unique MRNs) to total 10 selected medical records from NICU.
- 10. Select the non-NICU screening sample
 - a. Select NICU = No, and stratum = 2 (targeted pathogens)
 - b. Sort by random number, MRN, and admission date
 - c. Select the first 30 random numbers with unique episodes of care (defined by MRN and admission date) as the sample of adult/pediatric ICU medical records with candidate CLABSIs.
 - d. If 30 adult/pediatric ICU medical records with stratum 2 blood cultures are not available, supplement the non-NICU medical record sample with stratum 3 blood cultures (where NICU= No, and stratum = 3); take the initial medical records (lowest random numbers with unique MRNs) to total 30 selected medical records from adult/pediatric ICUs.
- 11. The final screening sample should contain: (up to) 20 medical records with reported CLABSIs, (up to) 40 medical records divided among NICU (if available) and adult/pediatric ICUs.
- 12. If medical records are not well balanced among <u>different</u> targeted pathogens, consider post-selection adjustment to include a variety of these organisms, in order to evaluate a variety of surveillance skills, as noted below.

Before requesting medical records for the audit, download ("freeze") the facility's reported data from NHSN

Why Target CLABSI Pathogens?

The targeted pathogens provide an opportunity to assess a facility's competency in correctly using different components of the NHSN CLABSI definition. For example:

- Candida and torulopsis (yeast) spp. are commonly seen in sputum samples, but infrequently
 cause true healthcare-associated pneumonia. NHSN cautions against reporting candida
 pneumonia in immunocompetent patients, unless there is evidence of invasive infection on
 lung biopsy or in pleural fluid under the definitions for PNU. These restrictions are further
 codified (as prohibitions) under ventilator-associated event (VAE). Candida BSI is common in
 ICU patients receiving parenteral nutrition. Reviewing medical records with candida BSI may
 provide an opportunity to look for misclassification.
- Some facilities that do MRSA active surveillance testing on admission incorrectly assume that MRSA colonization on admission means that a MRSA bloodstream infection would not need to be reviewed for CLABSI.
- Including enteric organisms such as enterococcus and gram negative rods can demonstrate a
 facility's ability to distinguish primary bloodstream infection vs. an alternative primary
 infection like UTI, GIT, or IAB with secondary bloodstream infection. Interested states can
 also assess use of the mucosal barrier injury reporting definitions, although these are not
 included in the Toolkit.

Facilities need to know how to correctly report single and confirmed isolates of common commensal organisms like coagulase-negative staphylococcus, and should be able to recognize synonyms (e.g. Staphylococcus epidermidis), used by the microbiology laboratory.

ICU CAUTI Medical Record Selection Process

- 1. From each selected facility, request a securely transmitted line listing of all positive ICU urine cultures, from all ICUs reporting to NHSN, for the entire year, with required additional variables used for medical record identification and matching to NHSN reports (See Chapter 3 for recommended line listing structure).
- 2. Assure the line listing includes appropriate positive urine cultures from all ICU locations required to report CAUTIs to NHSN, using location mapping information in NHSN
- 3. Assign a random number to each positive urine culture
- 4. Sort the list of urine cultures by MRN and admission date to generate clusters of urine cultures associated with recognizable episodes of care
- 5. Identify reported CAUTIs on the urine culture line listing
 - a. Using the NHSN CAUTI list and available patient information on urine culture line listing, flag and mark urine cultures reported as CAUTIs. Create a new variable, "stratum," and assign these urine cultures and all other urine cultures in the same medical record to stratum 1.
 - b. If reported CAUTIs are missing from the urine culture line listing, the list may be incomplete. Investigate and correct this problem. Add omitted CAUTI records to the medical record review list.
- 6. Select simple random sample of (up to) 20 reported ICU CAUTIs for review
 - a. Select stratum = 1
 - b. Sort by random number, MRN, and hospital admission date
 - c. Select the first 20 random numbers with unique patient episodes of care (defined by MRN and admission date) as the sample of reported CAUTI records
- 7. Identify unreported candidate CAUTI events
 - a. Select stratum not equal to 1
- 8. Select the screening sample
 - a. Sort by random number, MRN, and admission date (if available)
 - b. Select the first 40 random numbers with unique medical records (defined by MRN and admission date)
- 9. The final screening sample should contain: (up to) 20 medical records with reported CAUTIs, and (up to) 40 medical records without reported CAUTIs from adult/pediatric ICUs.

Before requesting medical records for the audit, download ("freeze") the facility's reported data from NHSN

COLO Procedure Targeted Medical Record Selection Process

- 1. Using NHSN, download a line listing of all COLO procedures for 2015, following these steps:
 - a. Log In to NHSN for the facility being validated and the Patient Safety Module.
 - b. From the left hand Nav Bar, Click "Analysis" then "Output Options."
 - c. Select the folder titled "Advanced," then "Procedure-level Data," then "CDC Defined Output."
 - d. Select the "Modify" button for "Line Listing All Procedures."
 - e. Under Modify Attributes of the Output, change the Output Name to "Line Listing COLO Procedures 2015," and the Output Title to "Line Listing for COLO Procedures 2015."
 - f. Option: Under "Select output format" retain "Output Format" as HTML (this will allow you to download and manipulate the file in Excel), and consider whether you want to check the box for "Use Variable Labels." This option will make the variable names longer (and more explicit), but is often not necessary if you know the variable names.
 - g. Under "Select a time period or Leave Blank...etc" for "Date Variable," select <u>ProcDateYr</u>, for "Beginning" enter 2015, and for "Ending" enter 2015.
 - h. Under "Specify Other Selection Criteria" do the following:
 - i. Column 1, row 1: select "procCode"
 - ii. Column 1, click row 2 to pop-up a gray dialog box, where Variable= "procCode", Operator= "=" and Value(s) = "COLO-Colon surgery"
 - iii. Click Save
 - iv. Column 2, row 1: select "outpatient"
 - v. Column 2, click row 2 to pop-up a gray dialog box, where Variable= "outpatient", Operator= "=" and Value(s) = "N-No"
 - vi. Click Save
 - vii. Column 3, row 1: select "ageAtProc"
 - viii. Column 3, click row 2 to pop-up a gray dialog box, where Variable= "ageAtProc", Operator= ">=" and Value(s) = "18"
 - ix. Click Save
 - i. Under "Modify Variables to Display by Clicking" select "Modify List"; retain the default Selected Variables: orgID, patID, dob, gender, procID, procDate, and procCode. Add variables by double clicking from the left hand list: ProcDateYr, outpatient, ageAtProc (to assure that you have selected 2015 inpatient adult COLO procedures), anesthesia, asa, procDurationHr, procDurationMin, Scope, medAff, numBeds, swClass, and modelRiskAll (variable that will be used to select procedures at higher risk to result in SSI). Click Save.
 - j. Under "Specify Sort Variables by Clicking" select "Modify List"; remove procCode from the right hand list by double clicking (all procedures will be COLO). Add procID by double clicking the variable in the left hand box; it will move to the right hand box. Click Save.
 - k. Select Run. You should see a line listing sorted by procID from lowest to highest. Click the box "Save As" to save your Template. The template will save under the name you specify, e.g., "Line Listing for COLO Procedures 2015."
 - I. Select Export Output DataSet. Under Export Output Options, select Excel Spreadsheet (*.xls). Select Export. An Excel file will be produced titled "LineListing_COLOProcedures2015."
- 2. Next, you will identify any of these procedures that have been reported to NHSN with an SSI. For this step, return to NHSN Analysis Output Options. This time, select the folders titled "Procedure-Associated Module," "SSI," and "CDC Defined Output."
 - a. Select the "Modify" button for "Line Listing All SSI Events"
 - b. Under "Modify Attributes of the Output" change the Output Name to "Line Listing COLO SSI Events 2015," and the Output Title to "Line Listing for COLO Surgical Site Infection Events 2015."
 - c. Optional: decide if you want to use Variable Labels.
 - d. Under "Select a time period or Leave Blank...etc" for "Date Variable," select <u>ProcDateYr</u>, for "Beginning" enter <u>2015</u>, and for "Ending" enter <u>2015</u>.
 - e. Under "Specify Other Selection Criteria" do the following:
 - i. Column 1, row 1: select "procCode"

- ii. Column 1, click row 2 to pop-up a gray dialog box, where Variable= "procCode," Operator= "=" and Value(s) = "COLO-Colon surgery"
- iii. Click Save
- iv. Column 2, row 1: select "outpatient"
- v. Column 2, click row 2 to pop-up a gray dialog box, where Variable= "outpatient", Operator= "=" and Value(s) = "N-No"
- vi. Click Save
- f. Under "Modify Variables to Display by Clicking" select "Modify List"; retain the default Selected Variables: orgID, patID, dob, gender, admitDate (this is date of admission for the procedure), eventID, eventDate, eventType, spcEvent, and procDate and procCode. Remove the remaining variables by double clicking.
- g. Under "Specify Sort Variables by Clicking" select "Modify List"; select linkedproc. This is the same variable as procID in the procedures file. NOTE: if you do not find a linked procedure, this SSI has probably been entered off-plan. You can use the other variable (procDate, patID, etc.) to investigate this.
- 3. Returning to the procedures file; mark any procedure that has been reported with an SSI as a reported case. All others are considered Candidate SSIs. Select the 40 candidate SSIs with the highest SSI risk ("modelRiskAll") for review.

Before requesting medical records for the audit, download ("freeze") the facility's reported data from NHSN

HYST Procedure Targeted Medical Record Selection Process

- 1. Using NHSN, download a line listing of all HYST procedures for 2015, following the steps outlined above for COLO.
 - a. If you have saved your template for downloading the line list of COLO procedures, you can make a few small modifications to download the HYST procedures rather than starting over (where you have entered "COLO" replace it with "HYST).
- 2. (As for COLO above), using NHSN, download a line listing of all HYST SSIs for 2015, following the steps outlines above, and replacing "COLO" with "HYST."
- 3. (As for COLOs above) return to the HYST procedures file; mark any HYST procedure that has been reported with an SSI as a reported case. All others are considered candidate SSIs. Select the 40 candidate SSIs with the highest SSI risk ("modelRiskAll") for review.

Before requesting medical records for the audit, download ("freeze") the facility's reported data from NHSN

Strategy for Selection of MRSA Bacteremia LabID Events for Validation

- 1. From each selected facility, request a securely transmitted line listing of all positive MRSA blood cultures from all inpatient locations /ED/ 24 hour observations for the entire year, with required additional variables used for medical record identification and possible matching to NHSN reports (See Chapter 3 for recommended line listing structure). Facilities should be STRONGLY encouraged to provide this in a spreadsheet (e.g. Excel) format.
- 2. Sort the line listing by specimen date. Assign a sequential number [1 to X] to each positive MRSA blood culture in the list. This will be used for random specimen selection.
- 3. Next sort the list by patientID, admission date, and specimen date. This allows you to identify individual episodes of patient care (a unique admission date and patientID) and to determine whether there is only one MRSA blood culture or multiple MRSA blood cultures during an episode of care.
- 4. Divide the original list into two lists: [A] first patient specimens (created by separating out all first specimens during a unique episode of care) and [B] non-first specimens (by separating out all remaining specimens). This may require some manual sorting.
- 5. Begin with list [B] (non-first specimens) to draw a random sample of 40 specimens that will be used to evaluate the SELECTED specimen and whether it should have been reported to NHSN. Sample only once from any episode of care
- 6. Use list [A] (first patient specimens) to draw a random sample of 20 specimens that will be used to identify the FIRST REPORTABLE LabID Event during an episode of care. In this case, validators are looking for evidence of positive MRSA blood cultures that are not on the inpatient list, but which were collected on the date of admission from an affiliated outpatient location other than ED/ 24 hour observations, or during a recent admission with an eligible specimen from the same inpatient location within the prior 14 days.

Before requesting medical records or other data for the audit, download ("freeze") the facility's reported data from NHSN

Strategy for Selection of *C. difficile* Infection (CDI) LabID Events for Validation

- From each selected facility, request a securely transmitted line listing of all toxin-positive Clostridium difficile stool specimens from all inpatient locations/ED/ 24 hour observations for the entire year, with required additional variables used for medical record identification and possible matching to NHSN reports (See Chapter 3 for recommended line listing structure). Facilities should be STRONGLY encouraged to provide this in a spreadsheet (e.g. Excel) format.
- 2. Sort the line listing by specimen date. Assign a sequential number [1 to X] to each toxin-positive CDI result in the list. This will be used for random specimen selection.
- 3. Next sort the list by patientID, admission date, and specimen date. This allows you to identify individual episodes of patient care (a unique admission date and patientID) and to determine whether there is only one inpatient CDI specimen or multiple inpatient CDI specimens during an episode of care.
- 4. Divide the original list into two lists: [A] first specimens (created by separating out all first specimens during a unique episode of care) and [B] non-first specimens (by separating out all remaining specimens). This may require some manual sorting.
- 5. Begin with list [B] (non-first specimens) to draw a random sample of 40 specimens that will be used to evaluate the SELECTED specimen and whether it should have been reported to NHSN. Sample only once from any episode of care.
- 6. Use list [A] (firstpatient specimens) to draw a random sample of 20 specimens that will be used to identify the FIRST REPORTABLE LabID Event during an episode of care. In this case, validators are looking for evidence of toxin-positive CDI results that are not on the inpatient list but which were collected on the date of admission from an affiliated outpatient location other than ED/ 24 hour observation or during a recent admission with an eligible specimen from the same inpatient location within the prior 14 days.

Before requesting medical records or other data for the audit, download ("freeze") the facility's reported data from NHSN

Appendix 1.4: Sample Letter Requesting Availability of Medical Records for Audit

Please customize this template to meet your state's needs

Dear [Name of IP]

As we discussed in our letter of [date], the [Name of Health Department] plans to audit surveillance practices and reporting of healthcare-associated infections for 2015 in multiple hospitals including your own. Thank you for your recent assistance in procuring the required line listings for medical record selection.

In the list below, we have identified the [XXX] medical records we would like to review during the audit, scheduled for [date(s)]. We appreciate your assistance in assuring that our team of [X] reviewers will have access to adequate working space, any necessary system passwords, and to these records when we visit. If your healthcare facility has initiated or completed conversion to an electronic medical record system, we will need a means of accessing these records including any diagnostic/laboratory results, clinical documentation, and ICD-9-CM codes related to these patients during our visit.

We look forward to visiting your facility and working with you in person. If questions arise, we can be reached at the following number [XXX-XXX-XXXX]:

Appendix 2: Surveillance Surveys

(Designed for External Validation of Surveillance Processes)

Appendix 2.1: CLABSI/CAUTI Surveillance Coordinator Survey

orgID /	Name of Hospital	_ Date of Survey
Instruc	tions: Administer this survey to the person who oversees NSHN surveillar	nce and denominator counting
1. W	hich best describes your facility's training for CLABSI and CAUTI Denomir	nator counters? (select all that apply)
	No specific training is provided or required	
	Peer training (person who previously counted) trains new staff	
	Training is provided by IP	
	Training by NHSN (e.g. online training) is required	
	Annual training updates are required / provided	
	Other (describe):	
	you conduct periodic spot-checks or otherwise validate CLABSI and CAL ply)	JTI denominator counts? (select all that
·	Not at this time	
	Yes, when we have a new denominator counter	
	Yes, when I have concerns	
	Yes, routinely	
3. W	hich best describes your own training for 2015 NHSN surveillance? (selec	et all that apply)
	No specific training for 2015	Select Training Modules Taken
	CDC-sponsored 2015 training webinar (live or on-line)	□CLABSI □ CAUTI
		□SSI □LabID Event
	CDC-sponsored 2015 on-line case-studies	□CLABSI □ CAUTI
		□SSI □LabID Event
	CDC-sponsored 2015 online self-paced interactive multimedia	□CLABSI □ CAUTI
	instruction trainings	□SSI □LabID Event
	State-sponsored 2015 NHSN training event(s)	□CLABSI □ CAUTI
		□SSI □LabID Event
	Other (describe):	
4. W	hich staff member(s) is/are responsible for entering CLABSI (numerator	□ IP
ev	ents) data into NHSN?	Clerical support
		Other
5. W	hich staff member(s) is/are responsible for entering CAUTI (numerator	☐ IP
	ents) data into NHSN?	Clerical support
		Other
6. Is	entered data checked for errors or validated by analysis?	Yes
		∏No
		Unk
	a. If yes, describe what is done:	·
	,,	

7.	How many persons typically review a medical record before an event is reported to NHSN?	☐ One reviewer typically decides, with internal (e.g. second reviewer) adjudication when needed ☐ Two or more persons typically review and agree before reporting ☐ One reviewer typically decides, with external (e.g. CDC) adjudication when needed ☐ Approval is required (e.g. from physician or administrator) before events are reported ☐ Other (explain):
8.	Is there ever pressure (e.g.; from administrators or physicians) to not report a CLABSI, CAUTI (or other NHSN) event?	Yes No Unsure Comment:
9.	In cases of ambiguity, who makes the final decision regarding the determination of whether an infection should be reported?	

Appendix 2.2: Documentation of Electronic CLABSI/CAUTI Denominator Validation

OrgID/ Name of Hos	spital:		Date of Survey	/:	
Instructions: NHSN	I requires that the mon	thly electronic deno	minator count falls within	n a 5% tolerance interval	
of the monthly ma	nual denominator cou	nt for 3 months befo	re reporting electronic d	enominator counts for	
CLABSI/CAUTI. If t	here is no electronic de	nominator counting	at this facility, skip this s	urvey.	
If electronic device	denominator counting	is used for reporting	g at this facility, documen	nt the NHSN-required	
validation results b	pelow:				
Initial electronic d	enominator validation	(when electronic d	enominator reporting be	gan):	
Location name:		Manual count	*Calculated 5%	Electronic count	
			tolerance interval		
Month/year:	Patient days				
	Central line days				
	Indwelling urinary				
	catheter days				
Location name:				·	
	Patient days				
Month/year:	Central line days				
	Indwelling urinary				
	catheter days				
Location name:		-			
	Patient days				
Month/year:	Central line days				
	Indwelling urinary				
	catheter days				
If available, please	e document additional	information for any	more recent electronic	denominator validation:	
Location name:		Manual count	*Calculated 5%	Electronic count	
			tolerance interval		
Month/year	Patient days				
.,	Central line days				
	Indwelling urinary				
	catheter days				
Location name:	,				
	Patient days				
Month/year	Central line days				
	Indwelling urinary				
	catheter days				
Location name:	,				
	Patient days				
Month/year:	Central line days				
• •	Indwelling urinary				
	catheter days				
*Equation for calcu	, , , , , , , , , , , , , , , , , , ,	terval is: manual cou	ınt ± (manual count * 0.0	5).	
	ons where manual cour			•	
· •	ce interval = [164±(164				
_	Electronic count 178 falls outside the tolerance interval.				

Appendix 2.3: Contact Information for Manual CLABSI / CAUTI Denominator Counters

Please feel free to adapt this template to meet your state's needs

NOTE: If facility assures annual training updates for denominator counters, and three or more denominator counters show proficiency on the survey in part 4, or if facility has already internally surveyed denominator counter proficiency, this can serve as evidence of proficiency.

OrgID	/ Name of Hospi	tal	_Date of Survey			
Instr	<i>Instructions:</i> Collect <u>contact information</u> for persons directly responsible for denominator collection in surveillance					
locat	tions and <u>adminis</u>	ster the survey (in p	art 4 below) late	er, by telephone.		
	Name of data	Surveillance	CLABSI	Work hours/	Phone number(s)	Supervisor
	collection	locations	CAUTI	Preferred time		
ID	professional	covered	Both	for telephone		
				survey		
1						
2						
3						
4						
5						
6						
7						
8						
9						
10						

To be expanded as needed....

Appendix 2.4: CLABSI and CAUTI Denominator Counting Survey (with Key)

divide	Instructions: Administer in person or by telephone, directly to individuals responsible for denominator counting. This form is color-coded so that it can be divided into a CLABSI denominator collection form (yellow and orange) in facilities where these tasks are performed by different persons. Orange indicates questions applicable to both CLABSI and CAUTI denominator collection.								
Facility OrgID:			of individual interviewed:		Name/ID of individual interviewed:		Position: ☐ IP ☐ Clerical ☐ Nursing ☐ Other (explain)	Interviewer initials:	Date of survey:
(circle): CLABSI, CAUTI, BOTH NHSN location(s) covered:									
PATIEN	IT DAYS (for both CLABSI and CA	AUTI denomi	inator counters)	Answer Key:					
1. Ho	ow are patient days usually colle	cted? (choos	se one)						
	Electronically (document the so	oftware							
	system utilized and skip to Q8)	:							
Manually									
	Some units electronic and som	e units manı	ual						
Comment:									
2. Is there a specified time when the denominator count is taken? ☐ Yes ☐ No		or	The answer should be Yes						
3. When is it done?		Counts should be done at a specific time daily, preferably at nearly the same time throughout the facility to avoid errors when patients transfer							
4. De	scribe the method used to cour	nt patient da	ays:	(from NHSN) "To calculate patient days, for each day of the month at the same time each day, record the number of patients. At the end of the month, sum the daily counts and enter the total into NHSN."					
	Count the number of patients	assigned to a	a unit bed <u>at the time counts are conducted</u>						
	Other (specify)								
Other (specify)									

	5. When reporting monthly patient day total, what is done if there are missing patient day data? (choose one)			NHSN issued specific guidance on imputing values for missing data in September 2013	
	,			gov/nhsn/PDFs/NHSNMissingDenomData_Sep2013.pdf)	
	Estimate or re-create missing data from existing information using our own metho	ods			
	Impute missing values using recent CDC/NHSN guidance				
	Other (specify):				
6. W	hich best describes your training for denominator (patient days and central line or	catheter days	s) counting? (sele	ect all that apply)	
	No specific training was provided			Formal training by NHSN or NHSN-trained IP is recommended due to technical aspects of definitions	
	Peer training (person who previously counted explained their approach to new staff)			(e.g., central line, permanent line, temporary line) and	
	Formal training by IP			methods (e.g., when to count lines, how many to count).	
	Formal training by NHSN (e.g., online training)			county.	
	Annual training updates				
	Other (describe):				
	hich staff member counts patient days and central line or catheter days when be "regular" data collector(s) is/are not working?	□ IP □	Another trained	counter	
8. D	oes your facility have a mechanism in place for quality control of denominator data	? (Select one)):		
	(Electronic data) Yes, data submitted electronically is periodically checked using	hods			
	(Manual data) Yes, manually collected data are periodically counted by more the				
	Yes, other (explain)				
	No formal quality control process				
9. Which staff member(s) is/are responsible for entering ICU patient days and central line or catheter day data into NHSN? □ IP □ Counter □				Clerical □ Other (specify)	

CENTRAL	CENTRAL LINE DAYS (for CLABSI denominator counters only)				
10. How	10. How are central line days collected for the unit(s) you oversee? (<i>choose one</i>)				
	Electronically (specify software system				
	utilized and skip to Q13):				
	Manually				
	Some units electronic and some units manua	I			
	Comment:				

11. ld	dentify the method used to count central line days : (choose one)		A daily count of the number of patients with a central	
	Count the number of patients with at least one central line at the time surveillance round	s are conducted	line in the patient care location during a time period, which is summed for the monthly total	
	Count the number of central lines that are in place at the time surveillance rounds are con	nducted	which is summed for the monthly total	
	Count the number of central lines that are in use at the time surveillance rounds are conc	lucted		
	Other (specify):			
12. When reporting monthly patient day total, what is done if there are missing central line day data? (choose one)			ific guidance on imputing values for missing data in	
	Report sum of available daily counts with no adjustment for missing data	(http://www.cdc.gov/nhsn/PDFs/NHSNMissingDenomData_Sep2013.pdf)		
	Estimate or re-create missing data using existing information (e.g.: medical records), then sum			
	Impute missing values using recent CDC/NHSN guidance for missing denominator data			
	patient has a radial arterial line and a peripheral IV. How many central line days are punted for this patient on this day?	Zero. The radial o	arterial line and peripheral IV are not central lines.	
	patient has a temporary central line and a permanent central line that are both in use. ow many central line days are counted for this patient on this day?	One. Although the patient has two central lines, a device day is defined as the number of patients who have the device, not the number of devices.		
О	he patient above with the temporary central line and the permanent central line is on an ncology ward. Should you report one temporary line day, one permanent line day, or both temporary and a permanent line day?	permanent lines,	n an oncology location has both temporary and the line day is reported as a temporary line day. This tailed in the NHSN Manual, Instructions for Form 57.117I)	
a	patient has a long-term port-a-cath that has not been accessed during this hospital stay, nd a peripheral IV that is in use. How many central line days are counted for this patient on his day?	Zero. The port-a-cath was not inserted during this visit and thus is not counted until accessed. The peripheral IV is not a central line. If the por a-cath was inserted during this admission it would be counted each day thereafter, whether in use or not		
	port-a-cath was inserted during this admission for planned chemotherapy. It is not in use. ow many central line days are counted for this patient on this day?	_	line was inserted during this admission it would be that it remains in place, whether in use or not	

18. A patient has a long-term central line that was accessed yesterday but is not currently in use, and a peripheral IN line days are counted for this patient on this day?	One. The port-a-cath was accessed during this stay and subsequently the line will be counted for each daily count until discharge, unless removed.	
19. A patient has a long-term central line that was accessed during evaluation leading to admission, but the line is n central line days are counted for this patient on this day	Zero. Brief access in an outpatient location does not count toward linedays during an admission. If the line had been accessed after admission or remained in use after admission following first access in the ED, it would be considered accessed for the purpose of counting line-days.	
20. If a central line is removed at 2PM and replaced at 8PM. The central line day count is done at 5PM, should the line be counted?	Yes No Unk	No. Central line must be in place at time of count
NICU-Specific Central Line Questions (Optional: Check here	and skip section if NICU questions do r	not apply to your job) 🔲)
21. When reporting central line (CL) days, in neonates, which neonatal weight is used for reporting? (select one)	☐ Birth weight ☐ Current weight	Birth weight
22. Neonates with both a CL and an umbilical catheter (UC) are included in the daily count as: (select one)	UC only CL only	CL only. No separate reporting of UCs; UCs are considered CLs, and reporting is for one or more CL, stratified by birth weight.

Indw	relling Urinary Catheter Days (for indwelling urinary catheter counters only)	
23. I	How are indwelling urinary catheter-days collected for the units you oversee? (choose one)	
	Electronically (specify software system utilized and skip to Q26):	
	Manually	
	Some units electronic and some units manual	
	Comment:	
24. I	dentify the method used to count indwelling urinary catheter days: (choose one)	7-2: Indwelling urinary catheter (AKA Foley catheter): A drainage tube
	Count the number of patients on the unit with a urine collection bag	that is inserted into the bladder through the urethra, left in place, and connected to a drainage bag, including urinary catheters that are used
	Count the number of patients on the unit with a Foley catheter or condom catheter	for intermittent or continuous irrigation, but excluding suprapubic,
	Count the number of patients on the unit with a Foley catheter, condom catheter, or suprapubic catheter	condom, or straight in-and-out catheters.
	Count the number of patients on the unit with a Foley catheter or indwelling urethral three-way (infusion) catheter used for bladder washes	
	Other (specify):	
	When reporting monthly patient day total, what is done if there are missing catheter day data? (choose one)	NHSN issued specific guidance on imputing values for missing data in September 2013
	Report the sum of available daily counts with no adjustment for missing data	(http://www.cdc.gov/nhsn/PDFs/NHSNMissingDenomData_Sep2013.pdf)
	Estimate or re-create missing data using patient information (e.g.: medical record), then sum	
	Impute missing values using recent CDC/NHSN guidance for missing denominator data	
	A patient has a draining ureteral stent and a Foley catheter; each one connected to a collection bag. How many urinary catheter days are counted for this patient on this day?	One. Ureteral stents are not counted because they are not urethral catheters
ı	A patient has a three-way indwelling urinary catheter used for irrigation after surgery to prevent blood in the bladder from clotting, and to provide for urinary drainage. How many urinary catheter days are counted for this patient on this day?	One. Catheters to be counted include indwelling urethral catheters used for intermittent or continuous irrigation, as well as those used for drainage.
	A patient on the unit has a supra-pubic urinary catheter. How many urinary catheter days are counted for this patient on this day?	Zero. Supra-pubic catheters are not urethral catheters because they enter the bladder through the abdominal wall.
i	A patient's indwelling urinary catheter is removed at noon and replaced at 5PM. Daily indwelling urinary catheter counts take place at 2PM. How many urinary catheter days are reported for this patient on this day?	None. There was no indwelling urinary catheter at the time of the daily denominator count. NOTE: However, If this patient develops a bloodstream infection attributable to a urinary tract infection, this day will count as one of two required catheter days to establish CLABSI criteria, because the catheter need only be in place for part of the two days to meet this criterion.

Appendix 2.5: Surgical Procedure and SSI Surveillance Methods Survey (with Key)

Instructi	Instructions: Administer this survey to the person who oversees NSHN SSI surveillance and reporting of surgical denominator (surgical procedure) data						
Facility or	g ID:	Name / ID of individual interviewed:	Position: □IP □Other (explain):		Interviewer initials:		Date of survey:
Procedu	re (Denominator) Data				_	
2)	data electronic entered manua If manual, who	ity normally upload surgical ally to NHSN, or is procedurally? (choose one): has primary responsibility as entry to NHSN? (choose of	re data for surgical	□ Electronic (skip to Q3) □ Manual □ Other (comment): □ IP □ Clerical/support staff □ Clerical/support staff	O oversight	and un	responsible for entering denominator data able to fully meet other responsibilities, recommend clerical support for this task
21	What source(s)	of information does your f	acility	□ Other		pieuse	Discussion for Q 3 and 4: Medical records
procedures? (choose all that apply): CPT codes assigned by ICD-9-CM procedure collaboration Vendor system using Collaboration Vendor system using Identification Vendor system using be procedure codes assigned Other		□ Selected flagged/filtered OR re □ CPT codes assigned by surgeon □ ICD-9-CM procedure codes ass □ Vendor system using OR record □ Vendor system using ICD-9-CM after discharge (specify) □ Vendor system using both OR is procedure codes assigned after discharge of the codes as a code after discharge of the code as a code after discharge of the code after disc	cords/reports is igned by coders after dis ds (specify) I procedure codes assign		coder opinion is regarded as technical gold standard for identifying NHSN procedures, but may be questioned if other sources are inconsistent, and is often not as timely as OR systems. Presence of designated ICD-9-CM procedure code is considered a requirement of NHSN procedure. Planned OR schedules are often inaccurate due to inability to predict procedures. OR records systems may be imprecise (e.g., may record XLAP rather than specifying that XLAP led to COLO, APPY, or SB). OR notes may be coded inaccurately; e.g.; surgeon may call procedure VHYS based on route of extraction whereas coder may classify as HYST based on route of detachment.		
		☐ Extra scrutiny to XLAPs ☐ Cross-reference data sources (explain):			Cross-referencing of sources (e.g.: OR records plus ICD-9-CM procedure codes assigned after discharge) is probably the best way to assure complete denominator. In general, XLAPs should be scrutinized by IPs conducting surveillance for COLO and HYST.		

5)	Under what circumstances do you remove COLO and/or HYST procedures from NHSN? (choose all that apply):	□ COLO or HYST ICD-9-CM procedure code was not assigned for the procedure □ COLO or HYST ICD-9-CM procedure code was assigned, but IP believes coder assigned COLO or HYST code in error □ Incision not primarily closed in OR □ Patient did not stay overnight □ Infection was present at the time of surgery (wound class = CO or D) □ ASA score was high □ Other	Although questioning of ICD-9-CM procedure codes is acceptable, removal of procedures with designated ICD-9-CM procedure code is only acceptable if procedure does not meet other aspects of NHSN procedure definition. Therefore it would be appropriate to remove procedure if there is 1) no appropriate ICD-9-CM procedure code, 2) no primary closure (note: new definition of primary closure for 2015), 3) not an inpatient (no overnight stay), 4) no incision/scope (Correct answers 1,3,4)
6)	If the OR record does not match the listed ICD-9-CM procedure codes, what should you do?		For validation purposes, NHSN recommends that IPs should bring coding mismatches to coders for review, and should not over-ride coders' decisions.
7)	Which of the following are consistent with the definition of primary closure for 2015 (clarified as of April 1)? (check ALL that apply)	 □ Complete closure of skin with suture □ Partial closure of skin with staples □ Closure of skin except for wick/drain through incision □ Closed fascia with incision loosely closed at the skin level □ Closed fascia, with skin layer left open 	All but the last option are considered primary closure in 2015.
8)	Does your facility conduct NHSN analysis to look at longitudinal trends for COLO or HYST SSIs and procedures?		This is recommended practice for facility use of NHSN data
9)	What would you do if your procedure denominator this month was dramatically higher from one month to the next?		Recommended: investigate this aggregate data by exploring the data at a patient/procedure level to identify the reason.

Surgical site Infection (Numerator) Data Collection Questions					
Instructions: Interview individual(s) directly respo	ng SSI data	Date of survey:	Date of survey:		
Name/ID of individual interviewed:	Position		(circle): COLO, HYST, BOTH		
Numerator (SSI Event) Data:					
10) If a patient with an SSI is admitted to your facility but the surgical procedure was performed in another hospital ("hospital A"), what do you do? (choose all that apply)		☐ Report the SSI to NHSN ☐ Report the SSI to "hospital A" ☐ Report the SSI to the health department ☐ No external reporting Comment:		Best practice is to report to "hospital A" and (if required by the state) to health department. Hospital A should report to NHSN.	
11) If you do not report the SSI to "hospital A", why not? (choose all that apply)		☐ HIPAA concerns ☐ Not a priority for IP program ☐ Logistically difficult (which hospital, who to contact) ☐ Not required Comments:		If facility cites HIPAA concerns, consider sharing Appendix 7, or CSTE position statement 13-ID-09, which contains information from the Office of Civil Rights assuring that sharing SSI information with the originating facility does not violate HIPAA.	
12) If you are contacted by the IP from another hospital regarding a patient with an SSI who underwent a procedure in your facility, what do you do? (choose all that apply)		□ Ask the IP for help completing the NHSN report □ Document in your tracking records □ Report the SSI to NHSN □ Ask the IP to report the SSI to NHSN □ No internal reporting or documentation Comment:		The other IP can best document the depth of infection, but cannot report the event to NHSN because it has to be linked. Suggest asking the other IP to help complete the NHSN report form, include a note or a copy in the patient record, and report to NHSN.	

13) What methods are routinely and systematically used to identify possible SSI? (Check all that apply)	Reports/Rounds: □ Emergency department line lists with diagnoses □ Admissions line lists with diagnoses □ Surgical ward rounds □ Positive laboratory cultures from inpatients □ Positive laboratory cultures from ED
	□ Pharmacy reports (antibiotic starts or continuations) □ Other
	Surgical service information: □ Inpatient returns to surgery □ Surgical service readmissions
	ADT/Medical Records Data Mining: □ Readmissions within one month of discharge □ Extended LOS □ Discharge diagnostic coding □ Other
14) How does your facility conduct post-discharge surveillance for SSIs? (check all that apply)	□ IP does not have a formal post-discharge surveillance plan □ IP conducts patient survey by mail □ IP conducts patient survey by telephone □ IP provides line list of patients to surgeon for response □ Surgeon indicates SSIs identified at surgical follow-up □ Surgeon surveys patient by mail □ Surgeon surveys patient by telephone □ IP reviews surgical clinic / wound clinic information □ IP reviews surgical patient records 30-60 days after procedures Other/ Comment:
15) During one trip to the operating room, both a COLO procedure and a HYST procedure are done. A deep-incisional SSI develops. To which procedure should you attribute the SSI?	 □ COLO □ HYST □ Both □ Whichever is higher on the procedure hierarchy (this would be COLO), because you cannot determine which procedure led to the SSI □ Neither

16) During one trip to the operating room, both a COLO procedure and a HYST procedure are done. The patient later meets criteria for a GI-IAB with peritonitis (an organ-space SSI). To which procedure should you attribute the SSI?	 □ COLO □ HYST □ Both □ Whichever is higher on the procedure hierarchy □ hierarchy □ Neither Two answers are correct(a and d): The procedure which is higher on the 2015 procedure hierarchy (this would be COLO) because you cannot determine which procedure led to the SSI
17) During one trip to the operating room, both a COLO procedure and a HYST procedure are done. An abscess of the vaginal cuff (organ-space SSI) develops. To which procedure should you attribute the SSI?	 □ COLO □ HYST □ Both □ Whichever is higher on the procedure hierarchy □ Neither The vaginal cuff is the operative site of the HYST, and the hierarchy is not needed; this SSI is attributable to the HYST (answer b).
18) During one trip to the operating room, both a SB procedure and a HYST procedure are done. An abscess of the small-bowel anastomosis site (organ-space SSI) develops. To which procedure should you attribute the SSI?	 □ SB □ HYST □ Both □ Whichever is higher on the procedure hierarchy □ Neither □ The SSI is localized to the operative site of the SB, and the hierarchy is not needed; this □ SSI is attributable to the SB (answer a). SB is higher on the hierarchy, but the hierarchy is only used when attribution cannot be determined by localized infection.

Appendix 2.6: LabID Event Surveillance Methods Survey (with Key)

OrgID / Name of Hospital

0.6.5	Trame of Hospital								
	ID Event Surveillance Methods Su ructions: Administer this survey to	•	es NH.	SN LabID Event reporting					
Denominator Data Collection Questions									
Nan	e of individual interviewed: Position: FacWideIN MRSA bacteremia FacWideIN CDI				Interviewer Da initials:			of survey:	
1)	For FacWidelN reporting, denom facility-wide level	inator data are entered	into I	NHSN once a month at the		True False		Т	
2)	For CDI reporting, the denominat	or should include all co	mplet	ted CDI toxin tests		True False		F (denominator = admissions and patient days)	
3)	Patient days include only admitte located on inpatient wards are ex		ward	s; observation patients		True False		F (all patients housed in inpatient locations)	
4) For CDI reporting pediatric locations should be excluded from FacWideIN reporting						☐ True ☐ False		F (NICU and well- baby locations and babies on LDRP are excluded for CDI)	
5)	For MRSA bacteremia reporting be excluded from the denominator	paby locations (NICU, ne	ewbor	n nursery, etc) should be		True False		F (no location exclusions for MRSA)	
Nur	merator Data Collection Questions	5							
Nan	ne of individual interviewed:	Position:		FacWideIN MRSA bacteremia FacWideIN CDI	Interviewer Date initials:			of survey:	
6)	For FacWideIN reporting, one mowide level	onthly numerator for Ev	ents i	s reported at the facility-		True False		F (events are reported by location)	
 For CDI reporting, the numerator should include toxin-positive CDI results conducted on formed stool specimens 						True False		F (laboratories should only process and report results for unformed stools)	
8)	A second event is always reporte MRSA bacteremia or toxin-positiv		ed fro	m the most recent positive		True False		Т	
9) A second event is only reported if >14 days have passed from the most recently reported labID event						True False		F (If the patient changes location, a second event is reported even within 14 days of prior event)	
10) A second event is only reported if the patient changes location OR >14 days have passed since the most recent positive MRSA bacteremia or toxin-positive CDI test in the same location						True False		Т	
11) Only reportable CDI LabID Events should be entered into NHSN						True False		Т	
Poli	Policy Question								
12) Does your facility laboratory limit CDI testing and reporting to unformed stool specimens only, or does the laboratory process all stool specimens for CDI if ordered?						Unformed specimens All stool specimens		Recommended policy is to only process unformed stool specimens for CDI	

Appendix 2.7: Template for Internal Validation of LabID Event Denominator (FacWideIN)

Please feel free to adapt this template to meet your state's needs

Electronically collected MRSA bacteremia and CDI FacWideIN denominators

"FacWideIN" includes all patient days counted at the same time each day for all inpatient locations, including any patients located for the day in inpatient locations, whether or not the facility considers them admitted patients or observation patients, but excluding any patients located for the day in outpatient observation locations. This information is typically collected electronically. Because the task of validating electronic patient days and admissions facility-wide is daunting, denominator validation can be accomplished using manual counting of patient days and admissions in three specified location types for three months each: one ICU, one Labor/Delivery/Recovery/Post-Partum (LDRP) location (if available), and one or more inpatient wards where observation patients are frequently located. Electronic counts should be within 5% of manual counts or an evaluation of why they differ should be conducted.

MRSA Bacteremia LabID Event Denominator Validation										
Location of	Month of	Admissions			Patient Days					
Validation*	Validation	Usual	5%	Manual	Usual	5% Tolerance	Manual			
	(specify)	Count	Tolerance	Count	Count	interval†	Count			
			interval†							
	1									
	2									
	3									
	1									
	2									
	3									
	1									
	2									
	3									

^{*}Select one ICU, one Labor/Delivery/Recovery/Post-Partum (LDRP) location if available, and one or more inpatient ward location where observation patients are frequently located and conduct manual (patient level) validation of admissions and patients days for three months, according to NHSN definitions

(http://www.cdc.gov/nhsn/PDFs/pscManual/validation/pcsManual-2015-valid.pdf, and http://www.cdc.gov/nhsn/forms/instr/57 127.pdf).

Remember that for MRSA bacteremia both mothers and babies are counted in LDRP locations.

†Equation for 5% tolerance interval is: Usual Count ± (Usual Count * 0.05).

Example calculations where Usual Count = 164 and Manual Count = 178:

Eligible 5% tolerance interval = $[164\pm(164*0.05)]=155.8$ to 172.2

Manual Count 178 falls outside the tolerance interval, suggesting that Usual Count is inaccurate and should be investigated.

CDI LabID Event Denominator Validation									
Location of	Month of		Admissions		Patient Days				
Validation*	Validation	Usual	Usual 5% Tolerance Manual			5% Tolerance	Manual		
	(specify)	Count	interval†	Count	Count	interval†	Count		
	1								
	2								
	3								
	1								
	2								
	3								
	1								
	2								
	3								

^{*}Select one ICU, one Labor/Delivery/Recovery/Post-Partum (LDRP) location if available, and one or more inpatient ward location where observation patients are frequently located and conduct manual (patient level) validation of admissions and patients days for three months, according to NHSN definitions

(http://www.cdc.gov/nhsn/PDFs/pscManual/validation/pcsManual-2015-valid.pdf, and http://www.cdc.gov/nhsn/forms/instr/57 127.pdf).

Remember that for CDI, only mothers (and not babies) are counted in LDRP locations.

†Equation for 5% tolerance interval is: Usual Count ± (Usual Count * 0.05).

Example calculations where Usual Count = 164 and Manual Count = 178:

Eligible 5% tolerance interval = $[164\pm(164*0.05)]=155.8$ to 172.2

Manual Count 178 falls outside the tolerance interval, suggesting that Usual Count is inaccurate and should be investigated.

Appendix 3: Medical Record Abstraction Tools

Note: Criteria, logic, and order of questions in the Medical Records Abstraction Tools should NOT be modified by state health departments; they have been designed and piloted to facilitate correct auditing using NHSN definitions. Please bring any problems to the attention of NHSN.

2015 CLABSI Medical Record Abstraction Tool

Part A: Determination of CLABSI								
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 Date of Audit/								
Patient ID Patient DOB/ Reviewer Initials								
Review Start Time: End Time: Time spent reviewing this record (minutes):								
FACILITY Admission Date:/ FACILITY Discharge Date:/								
Positive Blood Cultures / Repeat Infection Timeframe:								
Pre-screening Question: Were any positive blood cultures drawn on or after facility day 3 or was the DOE the day of transfer or discharge, or the next day? If Yes, continue If No, (all positive blood cultures were drawn before facility day 3) there was no HAI/CLABSI Event. STOP, record outcome (a) No candidate VL CLABSI								
a. Document <u>ALL positive blood cultures</u> sequentially below in Table 1aThen using information from "Table 1b. Locations" below, indicate which were "Validation Location (VL) blood cultures", defined as those collected <u>during VL stays</u> , or on day of or day after VL discharge. Note: <u>These VL blood cultures are eliqible for possible VL CLABSI</u> . (Non VL blood cultures may also be important to establish BSI repeat infection timeframe and other location of attribution.). Complete Table 1c. which documents presence placed/accessed.								
b. For each organism, indicate whether it is a pathogen (P) or common commensal (cc); the list of common commensals is available in LCBI Criteria. Note: Common commensals should only be evaluated as matched pairs/multiples if they were drawn on same/consecutive days; otherwise they are considered contaminants.								
The matching common commensals represent a single element; therefore, the collection date of the first common								
determine the Date of Event. If signs/symptoms occur within the 3 days prior to date of element, the first sign/symptoms RIT dates.								
KTT dates. c. Using clinical information (which can include signs/symptoms, and test results), divide listed blood cultures into distinct "RITs" and assign na RIT Number.								
Repeatedly positive blood cultures during previous BSIRIT (regardless of possible change in organism) are considered a single Infection Event.								
Note: Infection Window Period (IWP): The NHSN Infection Window Period is defined as the 7-days during which all sitespecific infection criteria must be met. It includes the day the first positive diagnostic test that is used as an element of								
the site-specific infection criterion, was obtained, the 3 calendar days before and the 3 calendar days after. The RIT is a 14-day timeframe during which no new infections of the same type are reported. The date of event is Day 1 of the 14-day RIT. If criteria for the same type of infection are met within the 14 day RIT, a new event is not identified or reported. Additional pathogens recovered during the RIT from the same type of infection are added to the event.								
Table 1a. List of positive blood cultures								

Positive BC	Date BC Collection	Location BC?	this date or day before?	Organism genus/species			P or CC*	Infectio n DOE#	RIT Dates and RIT number	
1	, ,			0.80	8000, 0000				1	, ,
2	//	Y/N	Y/N							
2	//	Y/N	Y/N							
3	//	Y/N	Y/N							
4	//	Y/N	Y/N							
5		Y/N	Y/N							
6		Y/N	Y/N							
7		Y/N	Y/N							
8		Y/N	Y/N							
9	//	Y/N	Y/N							
10	//	Y/N	Y/N							//
*BC=blood	culture, P=patho	gen, CC=commoi	n commensal, RIT=	Repeat Inf	ection Timefro	ате,	DOE=Date of Even	t. Add rows	if needed.	
Table 1b. Locations: Table 1c. Central Lines*:										
Document	all facility loca	tions and date	s sequentially for	r this	Document	t time	e periods below	with ANY ce	entral line i	n place for at least part
			itions being valid	lated for						ment individual lines
	circling Yes or						eplaced on same			
Facility Location	Physically Admit/	Discharge/ Transfer	Location Name	Pt in				•		great vessel (aorta, PA,
Order	Transfer IN	OUT	(include ED)	VL?						n, external iliac, common
			,		_					or accessed and used for NHSN Manual 4-2)
1	/ /	/ /		Y/N	CL placed		CL removed	aymanne me	meoring (r	
2				Y/N	accessed		without replacement	Locations h	oused with (CL
3	//	//		Y/N						
4	//			Y/N			//			
5				Y/N	, ,		, ,			
6 7				Y/N			//			
8		//		Y/N Y/N	/ /		, ,			
Add rows if	needed	1		1711	Add rows i	f nee				
		STIONS (ma	y be answered	in any o		,				
			ken during ANY	-	-	Sele	ct one:			
	lay of, or day at		_			□ Y	es -> Proceed			
						□ N	o -> STOP, rec	ord outcom	ie (a) No ca	ndidate VL CLABSI
S2. Was C	CL in place** fo	r >2 calendar d	ays AND in place	Select	one:					
during a VL stay for any period of time? — Yes -> Proceed										
□ No -> STOP, record outcome (a) No candidate VL CLABSI										
If yes to both screening questions: there is a candidate VL CLABSI.										
**In place: day of CL placement is considered CL day 1, unless patient admitted to facility with CL in place, where first central line access is CL Day 1.										
3. LABORATORY CONFIRMED BLOODSTREAM INFECTION (LCBI) CRITERIA										
3. LAD	ORATORY C	ONFIRMED I	BLOODSTREA	M INFE	CTION (LC	BI) (RITERIA			

- a. Evaluate all positive blood cultures in order as potential Laboratory Confirmed Bloodstream Infection (LCBI), using table columns below to determine if there was a LCBI, and which type (LCBI 1, LCBI 2 or LCBI 3) was met, if any. All elements listed in a column are required to meet the LCBI definition.
- b. If an LCBI definition is met, determine whether the LCBI also meets the corresponding definition of mucosal-barrier injury (MBI-LCBI), which is a subset of LCBI. Each positive blood culture reviewed should result in a reported outcome on page 4.
- c. <u>ONLY</u> IF Infection Event is related to infection at another primary site, document the alternative primary site and specific type of infection on page 4, <u>attach completed 2015 Tennessee checklist for alternative primary site</u>, and cite evidence (e.g.; required cultures, test results, logical and DOE dates) documenting that alternative primary site infection definition was met within a timeframe that does not exceed 1 calendar day between adjacent required elements, and that there was evidence of infection at the specific site at the time of positive blood culture collection and documentation of the secondary BSI attribution period. <u>Correct assignment also requires review of 2015 NHSN</u>

 Manual Ch. 4, Appendix 1 "Secondary Bloodstream Infection Guide", pp. 4-16 to 4-19, with documentation on page 4.

LCBI type:	LCBI 1 (any age)	LCBI 2 (any age)	LCBI 3 (age ≤1 year only)
Organism(s) in blood element	Recognized pathogen (P) cultured from one or more blood cultures	Matching common commensal* (CC) cultured from two or more blood cultures drawn on separate occasions on same or consecutive days (this is one element and can bridge to other elements either forward or backward).	Matching common commensal* (CC) cultured from two or more blood cultures drawn on separate occasions on same or consecutive days (this is one element and can bridge to other elements either forward or backward).
Other site exclusion	☐ Organism cultured from blood is not related to an infection at another site. ► If alternative primary site is likely, completed 2015 Tennessee checklist is required, with review of NHSN Manual Appendix 1 (pp 4-22 through 4-27). Type of alternative primary site infection, date of alternative primary event, and Appendix 1 criterion should be recorded under outcomes on p 4.	 □ Organism cultured from blood is not related to an infection at another site. ► If alternative primary site is likely, completed 2015 Tennessee checklist is required, with review of NHSN Manual Appendix 1 (pp 4-22 through 4-27). Type of alternative primary site infection, date of alternative primary event, and Appendix 1 criterion should be recorded under outcomes on p 4. 	Organism cultured from blood is not related to an infection at another site. If alternative primary site is likely, completed 2015 Tennessee checklist is required, with review of NHSN Manual Appendix 1 (pp 4-22 through 4-27). Type of alternative primary site infection, date of alternative primary event, and Appendix 1 criterion should be recorded under outcomes on p 4.
Age and Symptoms/ Signs element	(Any Age) (Any symptom or No Symptoms/Signs)	(Any Age) At least ONE of: Fever >38.0°C Chills, or Hypotension	(Infant ≤1 year of age) ☐ At least ONE of: ☐ Fever >38.0°C ☐ Hypothermia <36.0° ☐ Apnea, or ☐ Bradycardia
Timeframe	(NA)	All LCBI 2 elements must occur within the Infection Window Period, the seven-day time period which includes the date the positive blood culture was collected, the 3 calendar days before and the 3 calendar days after.	All LCBI 3 elements must occur within a the Infection Window Period, the seven-day time period which includes the date the positive blood culture was collected, the 3 calendar days before and the 3 calendar days after.
coagulase-neg			anthracis], Propionibacterium spp., spp., and Micrococcus spp.) See complete list
For any ever	nt meeting LCBI criteria above, determ	nine whether event is an MBI-LCBI usin	g criteria below.
	ets at least one of the following:		
_	·		vith one of the following documented
_	the same hospitalization as positive ide III or IV gastrointestinal graft vs		
		or ≥20 mL/kg in a 24-hour period for	natients <18 years of age) with
	•	fore the date the positive blood cult	
OR		•	
	· ·	ate days with values of absolute neu	· · · · · · · · · · · · · · · · · · ·
	-	thin a 7-day time period which inclu	•
Culture	AND-	lar days before, and the 3 calendar (- (select appropriate LCBI column)	uays arter.
	Organism is one of the following	Organism is viridans group	Organism is viridans group
МВІ	intestinal organisms and no other organism is isolated: Bacteroides spp., Candida spp., Clostridium spp., Enterococcus spp., Fusobacterium spp., Peptostreptococcus spp., Prevotella spp.,	streptococcus with no other organisms isolated	streptococcus with no other organisms isolated

		Veillonella spp., Enterobacteriaceae*,							
*Partial		CBI eligible <i>Enter</i>	obacteracia	genera:					
Citrobac	Citrobacter, Enterobacter, Escherichia, Klebsiella, Proteus, Providencia, Salmonella, Serratia, Shigella, Yersinia								
4. Did	4. Did Infection Episode Qualify as LCBI Event? (begin loop)								
	If LCBI definition was NOT met, record outcome ([b] No LCBI, and reason (e.g. unmatched common commensal or								
∐ No	asymptom Infection E		nmensais or	aiternative pi	rimary site	infection with s	econdary BSI), and continue to next		
	-	positive blood cu	ltures STOP						
		document type o			below				
Yes		may be more th				re.			
			- (Date of LCBI Event (date FIRST of required		
First LCD	I CDI 1	MADI LODI 1		BI (circle one):			elements was met):		
First LCB Second L	-		LCBI 2	MBI LCBI 2	LCBI 3	MBI LCBI 3 MBI LCBI 3			
Third LCI			LCBI 2	MBI LCBI 2	LCBI 3	MBI LCBI 3			
	s if needed	IVIDI ECDI I	LCDI Z	IVIDI LCDI Z	LCDI 3	IVIDI ECDI 3			
		thcare-Associat	tad Dracar	t on Admics	ion or N	oithar?			
							to the day after facility admission (POA)?		
	ect Yes or No	_	the time pe	1100 01 2 days	before far	cinty duringsion	to the day after facility admission (1 OA):		
							documented by a healthcare professional [e.g., acceptable. Physician diagnosis of LCBI		
		entation cannot be		,			,		
☐ Yes	If Yes, LCE	I was POA; doc	ument (c) F	OA LCBI typ	e and eva	luate next pos	itive blood culture outside of the		
	event LCB	I RIT.				-			
	If no more	blood cultures	, STOP						
□No	If no, prod	eed to b.							
	s the Date o	of Event on or a	fter the fac	ility Day 3?	Select Yes	or No):			
☐ Yes	If Yes, the	LCBI was HAI; ¡	proceed to	6.					
□ No	If No, LCB	was not HAI; a	locument (d) non-HAI L	CBI type o	ınd evaluate ne	ext positive blood culture .		
	If no more	blood cultures	, STOP						
6. Wa	s this HAI-L	CBI a CLABSI?							
			•	for >2 calen	dar days	present or rem	noved on the date of LCBI event or the		
		I event? (Select							
					e, day of fii	st line access is co	onsidered line Day 1.		
□ Yes	• •	-LCBI is CLABSI;	•						
□No	-	ıment e) HAI-LO		BSI and eval	uate next	positive blood	culture .		
		blood cultures				(, , , ,) ,			
		ON LOCATION				-			
		a VL on date of	LCBI Even	* or day bef	ore Even	t? (Select Yes or	r No):		
□ Yes	If yes, pro								
□ No	If no, document (f) CLABSI not attributable to VL and evaluate next positive blood culture outside the previous LCBI RIT.								
	If no more blood cultures, STOP								
*Date of	LCBI Event i	s date when first	of required	LBCI elements	occurred.				
	s patient tr		from anot	her bedded	inpatient	location, on da	ate of LCBI Event or day before Event?		
☐ Yes		ntion of attribut	tion was th	e <u>transferrin</u>	g locatior	. Proceed to c			
□No	If no loca	tion of attributi	ion was los	ation at time	of infact	ion: STOP reco	rd outcome (g) VL CLABSI		
	-	erring location					TO OUTCOME (9) VE CLADSI		
c. Was	s me transt	erring location	a vanuali0i	1 10CaU0N (V	∟ı: ıSeiec	i res or NOI:			

☐ Yes	If yes, location of attribution (transferring location) WAS a validation location; STOP record outcome (g) VL
	CLABSI
□ No	If no, location of attribution (transferring location) was NOT a validation location; record outcome (f) CLABSI
	not VL attributable

Positive Blood Culture Number	Outcome (a-g)	Detail for	Date of Primary Event/ Secondary BSI Attribution Period					
1								
2								
3								
4								
5								
(a) No candidate validation location (VL) CLABSI (b) No LCBI Reason, Select one): □ Contaminant (unmatched cc) □ Matching ccs with no symptoms □ Alternative primary source of BSI (complete box): -Primary source of BSIDate of alternative primary eventAttach TN checklist with elements abstracted -Circle correct NHSN Manual Appendix 1 criterion: 1. Blood/site-specific cxs match for ≥1 organism 2. Blood/site-specific cxs do NOT match but blood								
(c) POA LO	CBI CBI, Select one:	: LCBI1	MBI-LCBI1	LCBI2	MBI-LCBI2	LCBI3	MBI-LCBI3	
(d) non-HA	AI LCBI							
• •	CBI <i>, Select one</i> : BI not CLABSI		MBI-LCBI1	LCBI2	MBI-LCBI2	LCBI3	MBI-LCBI3	
Type of LC	BI, Select one	: LCBI1	MBI-LCBI1	LCBI2	MBI-LCBI2	LCBI3	MBI-LCBI3	
* *	not VL attributes. Select one		MBI-LCBI1	LCBI2	MBI-LCBI2	LCBI3	MBI-LCBI3	
	BI, Select one	: LCBI1	MBI-LCBI1	LCBI2	MBI-LCBI2	LCBI3	MBI-LCBI3	
Location of attribution Note: Each infection episode should have an assigned outcome a-g. There may be multiple LCBIs, or multiple CLABSIs								

Note: Each infection episode should have an assigned outcome a-g. There may be multiple LCBIs, or multiple CLABSIS during a single episode of care.

This tool requires that the episode of care be reviewed only until the first <u>validation</u> <u>location CLABSI</u> is found (option g above), or the end of the medical record is reached.

Don't forget to record the abstraction end time on page 1

2015 CAUTI Medical Record Abstraction tool

Fill in de	FIERS AND ABSTRACTED DATA emographic (white) section and ation as needed to answer ques	then complet	te the Section 2, sc	reening qu	estions. Fill in Tables 2a, 2b, 2c, and 3 to document
State	Facility (NHSN) orgID	•	<i>e</i>): ACH / LTACH / C Other	ancerH /	Date of Audit//
HICNO:	Patient ID	<u> </u>	Patient DOB		Reviewer Initials
Review Start Tir	ne:	End Time:		Time sper	nt reviewing this record (minutes):
FACILITY Admiss	sion Date//		FACILITY Disc	charge Date	e//
2. SCREEN	IING QUESTIONS (may be answ	vered in any o	rder)		
facility day 3 (Dallocation is Facili		npatient	Select one: ☐ Yes -> Procee ☐ No? -> STOP candidate VL CA	(a) Not a	Note: The complete list of UTI pathogens and common
S2. Were any positive urine cultures* taken during ANY validation location (VL) stay, the day of, or day after VL discharge?			Select one: ☐ Yes -> Proceed ☐ No? -> STOP (a) Not a candidate VL CAUTI		commensals are provided on the supporting documents sublink of NHSN website (http://www.cdc.gov/nhsn/acute-care-hospital/cauti/index.html)
•	catheter in place for >2 calend a VL stay for any period of time	•	Select one: □ Yes -> Procee □ No? -> STOP candidate VL CA	(a) Not a	
 Enter all Urine Cu Docume evidence infection HAI, whe location. 	ening questions: there is a candidal qualifying positive urine cultures of ltures, and indicate those collected at presence of Foley catheter (Tables, (Positive Blood Cultures Tables of pepisode sequentially for a UTI. You will be the HAI-UTI was a CAUTI, and will have the found by the country of the country of the sequentially for a UTI.	h as			
Laboratory Cult	ures				

*Positive urine culture = at least 10^5 CFU/ml of 2 or fewer organisms, one of which must be a bacterium with at least 10^5 CFU/ml. DO NOT LIST cultures with more than 2 species or those classified as "mixed" flora; these cannot be used to meet UTI criteria. Exclude urine cultures that are positive only for yeast, mold, dimorphic fungi, or parasites. Note: 10^5 =100,000

Document ALL positive urine cultures* sequentially below and using information from "Locations" below (Table 2b), indicate which were "VL urine cultures", defined as those collected during VL stays, or on day of or day after VL discharge. Note: These VL urine cultures should be evaluated for possible VL CAUTI. (Non VL urine cultures may also be important to establish prior onset of UTI Repeat Infection Timeframe (RIT) and another location of attribution.)

Columns 3, 4, and 7 (in red) are optional, but some validators may prefer to use these columns to organize their investigation

Table 1. Positive Urine Cultures

Candidate	Date UC	VL	Foley on this date	CFU/ml	Organism	Matched uropathogen in blood
UTI	Collection	UC?	or day before?	(≥10 ⁵)	genus/species	Within UTI IWP?**
					(maximum 2)	
1	/	Y/N	Y/N			Y/N or NA (sx)
2	/	Y/N	Y/N			Y/N or NA (sx)
3	/	Y/N	Y/N			Y/N or NA (sx)
4	/	Y/N	Y/N			Y/N or NA (sx)
5	/	Y/N	Y/N			Y/N or NA (sx)
6	/	Y/N	Y/N			Y/N or NA (sx)

Add rows if needed

Table 2a. Locations

Document all facility locations and dates for this episode of care chronologically below, and indicate locations being validated for CAUTI by circling Yes or No (VL=validation location).

Facility	Admit/Transfer	Discharge/	Location Name (include ED)	Validation Location (VL)?
Location Order	IN	Transfer OUT		
1	/			Y/N
2	/			Y/N
3	/			Y/N
4				Y/N
5				Y/N
6				Y/N

Add rows if needed

Table 2b. Foley Catheters

Document time periods with ANY Foley catheter in place for at least part of each day below (do NOT document individual catheters removed and replaced on same/consecutive days).

^{**}If colony counts are high (CFU/ml $\geq 10^5$), circle Y or N and document matching organism(s) isolated from blood in "positive blood cultures" Table 2d below; if patient with UTI symptoms ("sx") in UTI Infection Window Period (IWP) circle NA.

This document is outdated. Reference only.							
Foley placed or in	Foley removed without	Locations with Foley		Foley in validation location			
place	replacement						
	//_			Y/N			
	//_			Y/N			
	//_			Y/N			
	//_			Y/N			
	//_			Y/N			
	//_			Y/N			
Table 2c. Positive Blood	Cultures						
IF urine culture above cont	ains ≥10 ⁵ CFU/ml and patient is ASY	MPTOMATIC, document any p	oositive blood culture(s). This	information is needed to document ABUTI,			
-	=	commensal organisms in blood	d). At least one of the blood of	organisms must have been collected within			
	blood cultures, indicate below.						
□ No positive blood cult	ure(s) OR						
Candidate UTI	Blood culture collection dat	e Matching organism(s)	Matching common comm	nensal(s)			
(from Table above)							
1	_/_/_						
2	_/_/_						
3	_//_						
4	_/_/						
5	_/_/						
6							

3. S y	3. Symptoms* (Check one or more as required, or note date)											
* Sympto	* Symptoms required to meet UTI definition, within the IWP.											
No UTI sx	Candidate	Apnea	Bradycardia	CVA pain	Dysuria	Fever	Frequency	Hypothermia	Lethargy	SP	Urgency	Vomiting
	UTI									Tenderness		
	1											
	2											
	3											
	4											
Add rows i	f needed					•						

4. URINARY TRACT INFECTION (UTI) CRITERIA Starting with Candidate UTI #1, determine which type of UTI criteria [ABUTI, SUTI1a, SUTI1b, SUTI2] were met (if any). Required elements for UTI are highlighted in color. All elements listed in a column are required within the infection window time frame. UTI SUTI1a SUTI1b SUTI2 ABUTI type: (Symptomatic, any age) (Symptomatic, any age) (Symptomatic, infants only) (Asymptomatic, any age) □ ≥ 10⁵ CFU/ml urine ≥10⁵ CFU/ml urine ≥ 10⁵ CFU/ml urine (2 or fewer ☐ ≥ 10⁵ CFU/ml urine (2 or fewer microorganisms, (2 or fewer microorganisms, microorganisms) (2 or fewer microorganisms, at least one of which is a at least one of which is a urine culture at least one of which is a at least one of which is a bacterium of ≥105 element bacterium of ≥105 CFU/ml bacterium of ≥105 CFU/ml bacterium of ≥105 CFU/ml 0 Matching Blood organisms/matching common culture(s) element commensals Patient has/had an indwelling urinary catheter but it has/had not been in place >2 calendar days on the date of event+ OR (Any age, Foley present) At least ONE of: \square Patient did not have a Fever >38.0°C urinary catheter in place Suprapubic tenderness* on the date of event nor (With or without a Foley) (With or without a Foley) CVA pain or tenderness* the day before the date of \square Age ≤ 1 year AND (Any age) Age, AND event ☐ No listed symptoms ☐ At least ONE of: Appropriate Foley for >2 days and in place At least **ONE** of: allowed within time Fever >38.0°C symptoms (*= when last required element fever (>38°C) in a patient frame no other Hypothermia <36.0°C recognized documented that is \leq 65 years of age Apnea* (Foley optional) cause) and suprapubic tenderness* Bradycardia* Note: Patients > 65 years of Foley catheter • costovertebral angle Dysuria* age with a non-catheterstatus pain or tenderness* Lethargy* associated ABUTI may have element • urinary frequency * Vomiting* a fever and still meet the urinary urgency* (Foley optional) ABUTI criterion) • dysuria* --OR----OR---(Any age, Foley recently removed) ☐ At least ONE of: o fever (>38°C) in a patient that is ≤ 65 years of age Urgency*

	This do	cument is	outdated	. Refere	ence only.		
			 Dysuria* Suprapubic tenderness* CVA pain or tenderness* AND 				
5 Did candidate UTL	qualify as a UTI event, usir	ng criteria shown on	nage 32 (hegin lo	on)			
	If Yes, document type of UTI		 	• •	roceed to 6.		
□Yes	Note: there may be more th						
□ No		record outcome (<mark>b) no</mark> e.			with no matching pathogen in blood,). Loop to	next	
	Type of UTI		Date of UTI (date FIRST required element was met)		UTI RIT dates		
First candidate UTI			/ /		/to/		
Second candidate UTI			_		// to//		
Third candidate UTI		/	//		// to/		
Add rows if needed							
day RIT. If date of event for l the event.	JTI occurs within a previous 14 (IWP): The NHSN UTI Infection	4 day UTI RIT, no new L Window Period is defi	ITI is identified nor i ned as the 7-days di	reported. Additio	eterminations. The date of UTI event is Day 1 of the onal positive urine cultures during the UTI are add	ded to	
6. Was UTI Healthcar	e-Associated (HAI), Preser	t on Admission (PO	A), or Neither?				
a. Did the date of eve	ent of UTI occur during the	time period of 2 days	before admission	to the day aft	ter admission (i.e., POA)? (Select one):		
Note: Date of Event: The Date of Event is the date the first element used to meet an NHSN UTI criterion occurs for the first time within the seven-day infection window period. Acceptable documentation includes patient-reported signs or symptoms documented in the chart by a healthcare professional (e.g., patients states measured fever > 38.0° C or 100.4° F, nursing home documents fever prior to arrival to the hospital, patient complains of dysuria).							
□ Yes	If Yes, this UTI was POA; of the RIT If no more urine cultures,	STOP			luate next positive urine culture collected ou	tside	
□ No	If no, UTI was a HAI. Proceed to 7.						

7. Was this HA	-UTI a CAUTI?								
a. Was a Foley catheter in place for > 2 days on the date of event AND was either present for any portion of the calendar day on the date of event, OR removed the									
	day before the date of event?								
Note: If the patient w	Note: If the patient was admitted to a facility/ED with a Foley in place, date of admission to inpatient location is considered to be device day 1								
☐ Yes	If yes, HAI-UTI is CAUTI; proceed to 8.								
	If no, HAI-UTI was not CAUTI; document outcome (d) HAI-UTI not CAUTI and a UTI RIT is set. Evaluate next positive urine culture								
□No	outside the UTI RIT.								
	If no more urine cultures, STOP.								
8. Was VALIDA	TION LOCATION (VL) the Location of Attribution (LOA)								
a. Was patier	nt in a VL on the date of UTI Event* or day before UTI event? (Select Yes or No)								
□ Yes	If Yes, proceed to b.								
	If No, CAUTI was not attributable to VL; document outcome (e) CAUTI not VL attributable and a UTI RIT is set. Evaluate next positive								
□No	urine culture outside the UTI RIT.								
	If no more urine cultures, STOP								
*Date of UTI Event is	date when first of required UTI elements occurred during the UTI IWP.								
b. Was patien	t transferred to VL from another institution or bedded inpatient location, on date of UTI Event or day before UTI Event? (Select Yes or								
No):									
☐ Yes	If yes, location of attribution was the transferring location**; Proceed to c.								
□ No	If no, location of attribution was location at time of UTI Event; STOP, record outcome (f) VL CAUTI								
c. Was the tra	nsferring location** a validation location (VL)? (Select one):								
☐ Yes	If yes, location of attribution (transferring location) WAS a validation location; STOP, record outcome (f) VL CAUTI								
	If no, location of attribution (transferring location) was NOT a validation location; record outcome (e) CAUTI not VL attributable and a								
□ No	UTI RIT is set. Evaluate next positive urine culture outside the UTI RIT.								
	If no more urine cultures, STOP								
**If patient is transfe	rred more than once on the day of /day before the UTI Event, the FIRST transferring location from that time period is location of attribution.								

Outcom	Outcome of 2015 CAUTI audit:									
Candidate	Outcome	Detail for outcomes (b) through (g)	(a) Not a candidate VL CAUTI							
UTI*	(a-g)	(See key to right)	(b) No UTI; reason:							
1			☐ Asymptomatic but no matching blood pathogen							
T			(c) POA UTI (not HAI)							
2			(d) HAI-UTI not CAUTI							
			□ Type of UTI							
3			□ Date of Event							

4	(e) CAUTI not VL attributable
	Type of UTI
5	□ Date of Event
	Location of Attribution
	(f) VL CAUTI
	□ Type of UTI
	□ Date of Event
	□ Validation location of attribution

Note:

This tool requires that the episode of care be reviewed only until the first Validation Location (VL) CAUTI is found (outcome g above), or all positive urine cultures have been reviewed.

^{*}Report only those candidate UTIs which are outside previous UTI RITs

⁻ There may be multiple UTIs or multiple CAUTIs during a single episode of care.

2015 COLO Procedure/SSI Medical Record Abstraction Tool

For use in acute care hospital SSI validation following inpatient COLO procedures performed during Q1-Q4, 2015

1. Patient and Medical Record I	DENTIFIERS					
State Facility OrgID		Date of Audit		Reviewer Initials		
HICNO:	Patient ID		Patient DOB/_		Gender: F N	Л
Facility Admission Date 1 (for index C	OLO Procedure)		Facility Discharge Date 1			
Review Start Time:	End Time:		Time spent reviewing record	d (minutos):		
			Time spent reviewing record	<u> </u>	d during index C	2010 surgani
COLO Procedure Date://2 (USE THIS TOOL ONLY FOR COLOS PERFO	2015 DRMED IN 2015)		 Describe in words all procedures (e.g. colon resection, colon 		-	
 Circle ICD-9-CM COLO procedure 	,		(e.g. colon resection) colo	scomy jonnacion, a	ipperiacecomy).	
	41, 45.49, 45.52, 45.71-45.76, 45.79, 4	5 81-45 83 45 92-				
	13, 46.14, 46.43, 46.52, 46.75, 46.76,					
, , , , ,	-, - ,,,,,					
Note to validators: For O1-O3 2015 t	he presence of one or more of the abo	ve ICD-9-CM codes is				
•	ure. IPs are advised not to over-ride m					
decisions, but may discuss errors with		g				
·						
(Criteria for Q4 2015 will include ICD-10-F	PCS codes). Link to SSI section for ICD-10-PC	CS and CPT codes can be				
found in the "Supporting materials" section	on of the link below: http://www.cdc.gov	//nhsn/acute-care-				
hospital/ssi/index.html						
 "ModelRiskAll" Score (derived from 	om NHSN line list):					
Record later admission dates below o	only if they occur within 30 days of COL	O Procedure (Procedur	e date = day 1 of 30).			
Facility Admission Date 2://_		Facility Discharg	ge Date 2://			
Facility Admission Date 3://_		Facility Discharg	ge Date 3://			
2. NHSN Operative Procedure Cri	teria					
■ Did COLO operative procedure	COLO procedure performed on NH:					sh skin/mucous
meet NHSN definition for inpatient	membrane (including laparoscopic a					
procedure? (NHSN Manual 9-9 and 9-10)*	(Note incisional closure is NO longer	r an eiement of the NHSN	Operative Procedure definition, but	t is adaressed under i	risk-aajustment)	
3 10)						
□ No	If No, STOP, record (a) Not a cand	lidata COLO SSI: did i	not most NHSN Operative Pro	ocoduro dofinitio	n	
	· · · · · · · · · · · · · · · · · · ·	iluate COLO 331, ulu 1	Tot meet Nh3N Operative Pro	ocedure definitio	II	
□ Yes	If Yes, proceed to 3.					
*Notes to validator:						
Do not report procedure if ASA so						
 "NHSN Inpatient" requires that adm 	nission date is different from discharge date	2				
 "Inpatient procedure" requires use o criteria; see Manual 9-3 for details. 	f inpatient O.R./equivalent. "O.R. equivale	nt" may include C-section	room, interventional radiology roo	om, or cardiac cathete	erization lab mee	ting FGI or AIA

•	Regardless of wound class at the time of procedure or closure method (primary vs non-primary) all inpatient NHSN COLO procedures should be reported to the NHSN denominator, and all
	infections meeting COLO SSI criteria during the surveillance window should be reported.

3. Document COLO Procedure F	Risk-Adjustment V	ariables in Medical Record	at Time of Pro	cedure for	r Comparison to NHSN	
Type of closure: Definitions: Primary Closure: "defined of the presence of wires, wicks, drains category includes surgeries where the is closed at the skin level, by any man surgery. "(See NHSN Manual Ch 9-10)	s, or other devices or skin is closed by son ner, a designation of	objects extruding through the need the need the need to be need to	incision. This of the incision	Primary	Other than primary	
Diabetes: Definition: The NHSN SSI surveillance definition of diabetes indicates that the patient has a diagnosis of diabetes requiring management with insulin or a non-insulin anti-diabetic agent. This includes patients with "insulin resistance" who are on management with anti-diabetic agents. This also includes patients with a diagnosis of diabetes who are noncompliant with their diabetes medications. The discharge ICD-9-CM codes in the 250 to 250.93 range are also acceptable for use to answer YES to the diabetes field question.						
. ICD-10 CM Codes that may be used t with dates of surgery on October 1, 20 http://www.cdc.gov/nhsn/acute-care	015 or later, can be f	ound under Supporting Materio		Yes No		
	The NHSN definition excludes patients with no diagnosis of diabetes. The definition excludes patients who receive insulin for perioperative control of hyperglycemia but have no diagnosis of diabetes.					
Note: gestational diabetes is a type of	f diabetes.	464	(: ()	4 2 2	4 F /D / L'CACA C\	
			e (circle one):	1 2 3	4 5 (Do not report if ASA=6)	
		General anesthesia Scope (includes robotic)		Yes No		
	Emergency2 (no	n-elective and unscheduled		Yes No		
		blunt or penetrating injury)		Yes No		
	Tradina. ((Select one):	M F		
		Centre	Age (years):	.,,,,,		
			Height:	feet/	inches OR meters	
			Weight:	pou	inds OR kilograms	
Wound class (Select one)					CO D	
Was there evidence of	infection in the surg	ical area <u>at the time of the inde</u>	ex procedure*-	Yes No		
COLO procedure duration*:	Procedure date:	Procedure start time (mil***):	Procedure finis	sh* date:	Procedure finish* time (mil):	
Index procedure						
2 nd Procedure within 24 hours**						
Procedure duration (derived from		<u> </u>	minutes			
*Procedure finish time is when all inst				ays in OR a	re done, all dressings and drains	
are secured, and physicians/surgeons **If pt goes to OR again and another				of the origin	al procedure finish time and	
during the same admission, count as a						
ASA scores	,	, , , , , , , , , , , , , , , , , , , ,		J	•	

***	' minutes i	in length				
4.	Document Subsequent Surgery /Invasive Procedure During SSI Surveillance Window					
•	Was a s	ubsequent surgery performed through the primary incis	ion beyond 24 hours after the original procedure finish time but			
	within the 30-day surveillance window following the original procedure, OR was the surgical organ/space otherwise entered or					
	manipulated invasively (e.g., to drain a hematoma) at any time during the 30-day surveillance window [Date of procedure=Day 1]?					
	No	If no, skip to 5.				
	Yes	If yes, document additional procedure(s) and dates for a	onsideration and proceed to 5.			
Inv	asive pro	cedure 1:	Date 1:			
Inv	asive pro	cedure 2:	Date 2:			
Do	cument a	ny evidence of infection during invasive procedures abov	re:			
	Document surgical infection during surveillance window period					
5.	Docum	ent surgical infection during surveillance window perio	d			
5. ■			d urveillance window, including while hospitalized or post-discharge,			
	Was the	ere any documentation of surgical infection within the su				
	Was the	ere any documentation of surgical infection within the su	urveillance window, including while hospitalized or post-discharge,			
	Was the	ere any documentation of surgical infection within the sun nmunication from patient or other hospital, visits to the	urveillance window, including while hospitalized or post-discharge,			
•	Was the e.g. con require	ere any documentation of surgical infection within the sun nmunication from patient or other hospital, visits to the d when SSI is detected at a different facility).	urveillance window, including while hospitalized or post-discharge, ED or clinic? (NOTE: Reporting an SSI to the surgical facility IP is			
•	Was the e.g. con require No Yes	ere any documentation of surgical infection within the sum nmunication from patient or other hospital, visits to the d when SSI is detected at a different facility). If No, proceed to 7.	urveillance window, including while hospitalized or post-discharge, ED or clinic? (NOTE: Reporting an SSI to the surgical facility IP is			
• -	Was the e.g. con require No Yes	ere any documentation of surgical infection within the summunication from patient or other hospital, visits to the d when SSI is detected at a different facility). If No, proceed to 7. If Yes, abstract information regarding infection status is ent SSI Definition Criteria	urveillance window, including while hospitalized or post-discharge, ED or clinic? (NOTE: Reporting an SSI to the surgical facility IP is			
□□6.	Was the e.g. con require No Yes Docume Using the	ere any documentation of surgical infection within the summunication from patient or other hospital, visits to the d when SSI is detected at a different facility). If No, proceed to 7. If Yes, abstract information regarding infection status is ent SSI Definition Criteria BY NEW SSI Definitions criteria (see following), document we	urveillance window, including while hospitalized or post-discharge, ED or clinic? (NOTE: Reporting an SSI to the surgical facility IP is not the space below, and proceed to 6.			
□□6.	Was the e.g. con require No Yes Docume Using the (date was	ere any documentation of surgical infection within the summunication from patient or other hospital, visits to the d when SSI is detected at a different facility). If No, proceed to 7. If Yes, abstract information regarding infection status is ent SSI Definition Criteria BY NHSN SSI Definitions criteria (see following), document when the last element used to meet NHSN infection criterion.	ED or clinic? (NOTE: Reporting an SSI to the surgical facility IP is In the space below, and proceed to 6. This is the space below of infection criteria were met and the date of infection			
□ □ 6.	Was the e.g. con require No Yes Docume Using the (date we to deep,	ere any documentation of surgical infection within the summunication from patient or other hospital, visits to the d when SSI is detected at a different facility). If No, proceed to 7. If Yes, abstract information regarding infection status is ent SSI Definition Criteria when the last element used to meet NHSN infection criterion of the process o	ED or clinic? (NOTE: Reporting an SSI to the surgical facility IP is In the space below, and proceed to 6. In the space below, and proceed to 6. In thich depth of infection criteria were met and the date of infection occurred). Note: Available criteria for SSI may progress (e.g. superficial			

2015 HYST Procedure/SSI Medical Record Abstraction Tool

For use in acute care hospital SSI validation following inpatient HYST procedures performed during Q1-Q3, 2015

1. Patient	1. Patient and Medical Record IDENTIFIERS								
State	Facility	OrgID		Date of Au	ıdit	Review	er Initials		
HICNO			Patient ID	·	Patient DOB		Gender	F M	1
Facility Admission Date 1 (for index HYST procedure):					Facility Discharge Date 1:				
Review Start	Time:		End Time:		Time spent reviewing this re	cord (min	utes):		
HYST Proced	dure Date:	//2015	;)		Describe in words all proce	dure(s) du	ring index H	IYST	
(USE THIS TOO	OL ONLY FOR I	HYSTs PERFORM	ED IN 2015)		surgery (e.g. hysterectomy,			orec	tomy
■ Circle IC	D-9-CM pro	ocedure codel	s) for index H	YST:	(BSO), cesarean section, ap	pendector	ny)		
	 Circle ICD-9-CM procedure code(s) for index HYST: 68.31, 68.39, 68.41, 68.49, 68.61, 68.69 								
•		1-Q3 2015, the		e or more of					
			•	OLO procedure.					
		r-ride medical	_	•					
may discuss e			J	,					
(Criteria for Q4	4 2015 will inc	lude ICD-10-PCS	codes).Link to S	SI section for					
ICD-10-PCS and	d CPT codes c	an be found in th	e "Supporting m						
-		w.cdc.gov/nhsn/	'acute-care-						
hospital/ssi/ind	dex.html								
"ModelRickA	II" Score (de	rived from NHS	M line list\:						
Modeliniska	ii Score (de	iivea iioiii iviis	on intenst).						
Record later o	admission do	ates below only	if they occur v	vithin 30 days of	HYST Procedure (Procedure d	late = day	1 of 30).		
Facility Admis	ssion Date 2	:/			Facility Discharge Date 2:	_//_			
Facility Admis					Facility Discharge Date 3:	_//_			
2. NHSN	l Operative	Procedure C							
■ Did HYST					inpatient during trip to hospital				
•	e meet NHSN			-	nucous membrane (including		c approach	, or	during
definition for inpatient reoperation via an incision that was left open during a prior procedure.							ut is		
	procedure? (NHSN Manual (Note incisional closure is NO longer an element of the NHSN Operative Procedure definition, but is addressed under risk-adjustment)						аетіпітіоп. п		
procedure						riocedure	aejinition, b		
procedure 9-9 and 9-	-10)*	add	ressed under risk	r-adjustment)	ot meet NHSN Operative P				
procedure 9-9 and 9-	-10)*	add record (a) No	ressed under risk	r-adjustment)					
procedure 9-9 and 9-	No, STOP, Yes, procee	add record (a) No	ressed under risk	r-adjustment)					
procedure 9-9 and 9- No If Yes If *Notes to valid	No, STOP, Yes, proced	add record (a) No	ressed under risk t a candidate	r-adjustment)					
procedure 9-9 and 9- No If Yes If *Notes to valid Do not re "NHSN In	No, STOP, Yes, proceedator: eport procea	add record (a) Noted to 3. The second secon	ressed under risk t a candidate e=6 ay ("admission d	r-adjustment) HSYT SSI; did n		rocedure	definition	of inpo	

Regardless of wound class at the time of procedure or closure method (primary vs non-primary) all inpatient NHSN HSYT procedures should be reported to the NHSN denominator, and all infections meeting HYST SSI criteria during the surveillance window should be reported.

3. Document HYST Procedure Risk-Adjustment Variables in Medical Record for Comparison to NHSN					
Type of closure: Definitions: Primary Closure: "defined regardless of the presence of wires, wi incision. This category includes surgeri portion of the incision is closed at the should be assigned to the surgery. "(Se	Primary	Other than Primary			
Diabetes: Definition: The NHSN SSI surveillance definition of diabetes indicates that the patient has a diagnosis of diabetes requiring management with insulin or a non-insulin anti-diabetic agent. This includes patients with "insulin resistance" who are on management with anti-diabetic agents. This also includes patients with a diagnosis of diabetes who are noncompliant with their diabetes medications. The discharge ICD-9-CM codes in the 250 to 250.93 range are also acceptable for use to answer YES to the diabetes field question. ICD-10 CM Codes that may be used to answer the question about diabetes beginning with events with dates of surgery on October 1, 2015 or later, can be found under Supporting Materials at this site: http://www.cdc.gov/nhsn/acute-care-hospital/ssi/index.html The NHSN definition excludes patients with no diagnosis of diabetes. The definition excludes patients who receive insulin for perioperative control of hyperglycemia but have no diagnosis of diabetes is a type of diabetes. Note: gestational diabetes is a type of diabetes.					
		ASA score	(circle one):	1 2 2 / 5	5 (Do not report if ASA=6)
		General anesthesia		Yes No	b (Do not report ij ASA-o)
			(Select one):	Yes No	
	Emergency	? (non-elective, unscheduled)	(Select one):	Yes No	
	Trauma?	(blunt or penetrating injury)		Yes No	
			(Select one):	F	
		Patient	: Age (years):	£+/	inches OD masters
			Height: Weight:	feet/ pounds	inches OR meters OR kilograms
		Wound class	(Select one):	C CC CO [
Was there evidence of inf	ection in the surgica	al area <u>at the time of the index</u>	·	Yes No	
HYST procedure duration*:	Procedure date:	Procedure start time (mil):	Procedure fi	nish* date:	Procedure finish* time (mil):
Index procedure					
2 nd Procedure within 24 hours**					
Procedure duration (derived from	above informatio	n):hours and	minutes		
*Procedure finish time is when all instr	ruments and sponge	counts are completed and ve	rified, post-op	x-rays in OR are	e done, all dressings and
drains are secured, and physicians/sur	•	·			
**If pt goes to OR again and another p during the same admission, count as o ASA scores.					

		This document is out	<u>luateu. Neference only.</u>			
4. D	ocumen	t Subsequent Surgery /Invasive Procedure During (HYST) SSI Surveilland	ce Window			
•	Was a subsequent surgery performed through the primary incision beyond 24 hours after the original procedure finish time but within the 30-day surveillance window following the original procedure, OR was the surgical organ/space otherwise entered or manipulated invasively (e.g., to drain a hematoma) at any time during the 30-day surveillance window [Date of procedure is Day 1]?					
	No No	If No, skip to 5.				
	Yes	If yes, document additional procedure(s) and dates for consideration an	d proceed to 5.			
Inva	sive pro	cedure 1:	Date:			
Inva	sive pro	cedure 2:	Date:			
Doc	ument aı	ny evidence of infection during invasive procedures above:				
5.	Addition	nal /Post-Discharge Infection Surveillance				
•	• Was there any documentation of surgical infection within the surveillance window, including while hospitalized or post-discharge (e.g. communication from patient or other hospital, visits to the ED or clinic)? (NOTE: Reporting of SSI to the surgical facility IP is required when SSI is detected at a different facility).					
	No	If No, proceed to 7.				
	If Yes, abstract information regarding infection status in the space below, and proceed to 6.					
6. D	ocumen	t SSI Definition Criteria				
(dat supe oper	e when terficial to erficial to n space i	ASN SSI Definitions criteria (See following), document which depth of inferthe last element used to meet NHSN infection criterion occurred). Note: An odeep); review the entire infection event and record the DEEPEST level of an 5 above, and the checklist that follows to document information and deform SSIs, continue to part 7B for attribution assignment.	Available criteria for SSI may progress (e.g. f SSI during the surveillance window. Use the			

6 (continued): NHSN SSI Definitions: Use checklist to establish elements met:						
Superficial Incisional HYST SSI	Deep incisional HYST SSI	Organ/Space HYST SSI				
 Occurs within 30 days or end of surveillance window (whichever comes first) 	Occurs within 30 days (HYST) or end of surveillance window (whichever comes first)	Occurs within 30 days (HYST) or end of surveillance window (whichever comes first)				
AND	AND	AND				
Involves only skin and subcutaneous tissue of the incision	Involves <u>deep soft tissues</u> (e.g., fascia and muscle layers) of the incision	Involves <u>any body part opened or manipulated</u> during surgery except skin incision, fascia or muscle.				
AND	AND	AND				
at least one of the boxes:	at least one of the boxes:	at least one of the boxes:				
 Purulent drainage from superficial incision 	Purulence from deep incision	 Purulence from a drain placed into the organ/space 				

	Time accaminant to co	aldated. Reference offig
 Organisms isolated from Aseptically-obtained culture of fluid or tissue superficial incision 		 Organisms isolated from Aseptically- obtained culture of fluid or tissue organ/space
 Attending physician* deliberately opened superficial incision AND Culture-positive or not-cultured AND At least one of: pain or tenderness localized swelling redness heat 	Spontaneous dehiscence or surgeon deliberately opened deep incision AND Deep incision is culture-positive or not-cultured AND At least one of: fever (>38.0°C) localized pain or tenderness	
 Diagnosis of superficial incisional SSI by attending physician* 	 Abscess or other evidence of infection involving the deep incision that is found on (at least one of) Direct examination During invasive procedure Histopathologic examination Imaging test 	Abscess or other evidence of infection involving the organ/space that is found on (at least one of) Direct examination During invasive procedure Histopathologic examination Imaging test
• , , , , , ,	ourposes of application of the NHSN SSI criteria e, emergency physician or physician's designee	, , , , , , , , , , , , , , , , , , , ,
injectious discuse, other physician on the cus	e, emergency physician or physician's designed	AND
		Meets at least one criterion for a specific organ/space infection site; particularly (for HYST) IAB, OREP, or VCUF. Document using Tennessee Checklist.
	Reporting Notes	
 Do not report stitch abscess, localized stab wound, pin site infection, or cellulitis alone 	The depth of SSI (SI, DI, or O/S) reported should reflect the deepest tissue layer involved during the surveillance window.	If a patient has O/S infection during the primary operative procedure subsequent continuation meeting NHSN SSI criteria is considered an O/S SSI.

	7. Outcome of 2014 HYST SSI audit 7(A): Select (a), (b), or (c); If (b) is selected, define depth and date of infection, then proceed to 7(B):						
	(a) Not a candidat	te HY	ST SSI: Did not meet NHSN Operative Procedure definition				
	(b) SSI:		(b1) Superficial incisional SSI	Date of SSI (date SSI was first met			
	(select deepest level during		(b2) Deep incisional SSI	at any depth):			
	surveillance window)		(b3) Organ/Space SSI (Specify site)				
	(c) No SSI						
7(B): Attribution of H	IYST S	SI to Procedure				
-	Was the SSI attrib	outabl	e to the HYST, or was the SSI attributable to another invasive concur	rent NHSN Operative Procedure or			
	to invasive manip	ulatio	on of the HYST operative site after the HYST procedure? (Select one):				
	HYST SSI		Note to validator: In the context of serial invasive manipulations (inc				
			operative site, an SSI is attributed to the most recent intervention. In				
			NHSN Operative Procedures through the same incision, superficial and deep incisional infections are				
	SSI not attributab	le	attributable to the procedure highest on the surgical hierarchy*, because there is no way to distinguish				
	to HYST; SSI		which of the NHSN Operative Procedures led to the infection. For organ/space SSIs, the specific location				
	attributable to		of infection should be examined for attribution; e.g., in the event of concurrent COLO and HYST, a vaginal				
	(specify):		cuff infection should be attributed to the HYST. E.g.; in the event of a				
			of the bed of the spleen should be attributed to the SPLE. E.g.; in the				
			COLO, deep pelvic abscess would be attributed to the HYST, whereas	the hierarchy would assign			
			peritonitis to the COLO. (*See hierarchy below)				

*2014 Sur	*2014 Surgical Hierarchy for Abdominal Operations, from NHSN Manual Ch 9, Table 5.				
Priority	Code	Abdominal Operations			
1	LTP	Liver transplant			
2	COLO	Colon surgery			
3	BILI	Bile duct, liver, or pancreatic surgery			
4	SB	Small bowel surgery			
5	REC	Rectal surgery			
6	KTP	Kidney transplant			
7	GAST	Gastric surgery			
8	AAA	Abdominal aortic aneurysm repair			
9	HYST	Abdominal hysterectomy			
10	CSEC	Cesarean section			
11	XLAP	Exploratory laparotomy			
12	APPY	Appendix surgery			
13	HER	Herniorrhaphy			
14	NEPH	Kidney surgery			
15	VHYS	Vaginal hysterectomy			

16	SPLE	Spleen surgery
17	CHOL	Gall bladder surgery
18	OVRY	Ovarian surgery

8.	(For health department use) If SSI was missed by facility, what was the reason?							
	☐ Surveillance missed	Provide detail:						
	Misinterpreted criteria							
	☐ Incorrect use of infection at another site							
	☐ MD ruled out an SSI							
	□ Other							

Don't forget to record the abstraction end time on page 1

2015 MRSA Bacteremia LabID Event (FacWideIN) Validation Tool

For validation of MRSA bacteremia (only) LabID Event reporting in acute care hospitals conducting LabID Event surveillance for either MRSA bacteremia or all MRSA clinical specimens (EXCLUDES SCREENING CULTURES for colonization). Note: Based on CDC sampling guidance, this tool will be used in two ways; [Sample A] to validate reportability of the FIRST inpatient MRSA culture for a patient and episode of care, and [Sample B] to validate reportability of a subsequent SELECTED (non-first) MRSA culture for a patient and episode of care. Sample A evaluates the facility's ability to link early inpatient MRSA cultures to recent episodes of care and affiliated ED/outpatient specimens on the date of admission. Sample B evaluates the facility's ability to correctly classify duplicate vs. reportable events

							_		_			_		
Facility	LabID Event S	Surveillance M	lethod (NH	SN monthly rep	orting pla	n)				mia only (note: <i>on</i> mens (note: <i>inclu</i> e				
Sample	e 🗆 Sa	ample A: valid	ating "first	" inpatient MRS	A culture			Date of "first" in	npatient I	MRSA culture				
Select of				CTED (non-first)		MRSA culti	ıre	Date of SELECTE	ED (non-f	first) inpatient lab	culture			
Instruct		1	0 -							,		<u> </u>		
For Sam other for ED or for calenda Through collecte For Sam to this s	nple A: Begin wan cility-affiliated acility-affiliated ar day of admis hadditional in dwithin that the begin was elected specin was acility.	I outpatient lou I outpatient sp sion, reporters vestigation, de imeframe for t ith the SELECT nen. If prior sp	cation on the ecimen is identification in the callowes termine if the patient of the callowes from the patient of the patient of the callowes from the callo	ne NHSN inpatien the NHSN inpatien the species of the same location of the same pation the same pation of the	t admissione "first" in the "first" in the pecimen to prior inpa n. Working A culture. tent were fi	n date. If su patient MR o the locatio tient stay w g across the Using the So ound, enter	ch a specime SA Culture in n of inpatien ithin the 14 c row, determ ample B sorte theses specin	determine whether a part is identified, enter in Table 2, row M1. No that admission. If the chosen ("firsted list, identify the momens starting in Table in the SELECTED (")	t in Table te that If n collectio st") inpati ost recent e 2, row S2	2, row S1. Then en a specimen was co an, and whether an ient MRSA Culture specimen collected 1. If no prior speci	iter the "first" ollected in ED y MRSA Cultu on the labora d from the sai imen was colle	'inpatient sp or affiliated o re specimens tory line list o me patient in ected from th	ecimen in ro outpatient lo were report was reportal the same lo ue same loca	w S2. If no cation on ted or ole* to NHSN. cation prior tion, enter
Patient	atient and Medical Record IDENTIFIERS													
NHSN c	orgID#:			Date of Audit:					Revie	ewer Initials:				
Review	Start Time:			End Time:				Time spent rev	iewing th	his record (minute	es):			
Patient	ID	Patient D	ООВ	NHSN Inpatient location as "ob		•	•	ced in inpatient	d in inpatient Facility Location 1 (Specific first inpatient bedded location name; not ED):					nme; not
HICNO		Gender	F M											
Table 1 validati		ocations and t	ransfer dat	tes: If validating	for Sampi	le B, enter d	ates and loca	ntions from ADT data	up to the	date of SELECTED	specimen . (N	lote: SKIP ste	o below for S	Sample A
Date tra	ansfer to Locat	ion 2		Facility Lo	cation 2			Date transfer to Locat	tion 4		Facility I	Location 4		
Date tra	ansfer to Locat	ion 3		Facility Lo	cation 3			Date transfer to Locat	tion 5		Facility I	Location 5		
Table 2	: Positive MRS	A Blood Cultu	res											
Α	В	С	D		E			F			G (Complete specimen su		Н	
Lab Date of specimen collection Location of specimen collection Location of specimen collection* Location of specimen collection* Location of specimen last positive MRSA cult from same NHSN location?				days since last pos patient in same lo previous episode	Was this a "duplicate specimen", i.e.; ≤14 days since last positive MRSA Culture AND patient in same location (could include a previous episode of care) Was this MRSA blood culture the first MRSA culture of any type fo this month?		first MRSA ny type for	Reportable to NHSN**?						
S1 /_/_ days □ no prior □ No □ Yes □ no prior			□ No		Yes	□ No	☐ Yes	□No	☐ Yes					
S2	_/_/_		days	S	□No	☐ Yes		□No		Yes	□No	☐ Yes	□ No	☐ Yes
S3			days	S	□No	☐ Yes		□ No		Yes	□No	☐ Yes	□No	□Yes
*Repor	table to NHSN	if:					· ·							

- Facility conducting MRSA all specimen surveillance, AND MRSA blood culture was first positive MRSA culture of any type for the month (Table 2, Column G = yes).
- No prior MRSA bacteremia from the patient in the same location
- More than 14 days since last MRSA bacteremia from the patient in the same location

2015 CDI LabID Event (FacWideIN) Validation Tool

For use in acute care hospitals (ACH). Note: This tool will be used in two ways; [Sample A] to validate reportability of the FIRST inpatient CDI toxin-positive specimen for a patient and episode of care, and [Sample B] to validate reportability of a subsequent SELECTED (non-first) CDI toxin-positive specimen for a patient and episode of care. Sample A evaluates the facility's ability to link inpatient specimens to recent episodes of care and affiliated ED/outpatient specimens on the date of admission, which could render the first inpatient specimen non-reportable; Sample B evaluates the facility's ability to correctly classify duplicate vs. reportable events.

eric racinty 5	ability to col	rectly classify	auplicate	vs. reportabl	C CVCIICS.									
Patient and	d Medical Re	ecord IDENTIF	IERS											
NHSN org	ID#:			Date of Au	dit:			Re	viewer Ir	nitials:				
Review St	art Time:			End Time:		Т	ime spent re	viewing t	his recor	d (minutes):				
Patient DO	OB Patie	nt ID	HICNO		I '		mission Date ation or admi	•		ed in inpatient	-	ation 1 (Specific first i ime; not ED):	npatient l	edded
Select	☐ Sam	ple A: validat	ing chose	en "first" in	oatient CDI to	oxin-posit	ive specime	n	Date of	chosen "first" inpatio	ent CDI toxin	-positive specimen:		
one:		ple B: validat				•	•					I toxin-positive specir	nen:	
Note: SKIP				· ·					-	a up to the date of SELE	<u> </u>			
	er to Location	r'			Location 2	,	-	Date trans			,	Facility Location 6		
Date transf	er to Location	on 3			Location 3			Date trans	fer to Loca	ation 7		Facility Location 7		
Date transfer to Location 4 Facility Location 4 Date transfer to Location 8 Facility Location 8														
Date transf	er to Location	on 5		Facility	Location 5			Date trans	fer to Loca	ation 9		Facility Location 9		
facility-affi facility-affi location on Through ac timeframe For Sample location pri specimen v SELECTED (liated outpa liated outpa calendar da Iditional inve for the patie B: Begin wit ior to this se vill then be i	tient location of tient speciment of admission estigation, detent and same let the SELECTE lected speciment C2. If no price to took to took to took took took took	on the NH in is identifing, reporter ermine if the occation. V ED (non-firen. If a prior specime kin-positiv	SN inpatient ied, enter the rs are allowed this patient hooking acrost inpatient ior specimen was collect	admission dat chosen ("first d to assign the ad a prior inpo s the row, det CDI toxin-posi from the same ted from the s	e. If such a ") inpatier specimen atient stay rermine if to tive specin e the patien	a specimen is id nt CDI toxin-po to the location within the prid he chosen ("fil men. Using the nt and location on, enter the S	dentified, e positive spec n of inpatie or 14 days, rst") inpati e Sample B n is found, SELECTED (enter it in ictimen in rictimen in rictimen in rictiment admission and when icent CDI to sorted listing enter this inon-first)	row C1. Then enter the cow C1. Note that If a spision. ther any CDI toxin-positioxin-positive specimen cost, identify the most receive specimen in row C1; the inpatient CDI toxin associated.	chosen ("first", pecimen was co ive specimens on the laborato nt specimen co e SELECTED (no	ras collected from this poor inpatient specimen in recollected in ED or affiliate were reported or collect ory line list was reportab collected from the same p on-first) inpatient CDI to looking across the row, of	ow C2. If n d outpatien ed within t le† to NHS atient in th xin-positive letermine i	o ED or nt hat N. ne same
A B		С	D			E				F			G	
list spe	e of cimen ection	Location of specimen collection*		umber of day OI toxin-posit			positive CDI to n from same N			Was this a "duplicate s positive CDI toxin-posit location (could include	tive specimen .	AND patient in same	Reporta NHSN†	ble to
C1	/			days	☐ no prior	□ No	☐ Yes	☐ no pr	ior	□ No		☐ Yes	□No	☐ Yes
C2	 //_		_	days		□No	☐ Yes	•		□ No		☐ Yes	□ No	☐ Yes
C3	 //_			days		□No	☐ Yes			□ No		☐ Yes	□No	☐ Yes
	n collected		ted outpa	tient locatior	on calendar o	day of adm	ission, reporte	ers are allo	wed to as	ssign specimen entered	in NHSN to the	e location of inpatient ac	lmission, to)

†Reportable to NHSN if:

- No prior positive CDI toxin-positive specimen from the patient in the same location –OR--
- More than 14 days since last CDI toxin-positive specimen from the patient in the same location

Appendix 4: Documentation of External Validation Results

Appendix 4.1: (Optional) Templates for Audit Discrepancies Discussion with Facilities

Please feel free to adapt these templates to meet your state's needs to discuss discordant outcomes and request changes

(Instructions: For each HAI Event with discordant outcome between reporters and validators, record the following [first row-enter hospital report; second row-enter recommended changes]. Use the Comment area to document reasons for error, e.g.: overlooked candidate culture; confusion re common commensals; did not meet alternative primary definition, not an uropathogen, etc. Many states have examined this type of data to identify common errors and direct future education and training. Keep a copy for your records and leave a copy with the facility). H=hospital; V=validator

Central line-associated Bloodstream Infection (CLABSI) Discrepancies

Pt. ID		Positive blood culture	Select One	:		If LCBI,	If LCBI,				
		event: first culture				Event	MBI*				
		date				date	LCBI?				
			Not	Alternative	LCBI1,		MBI Yes	POA,	Central line >2d?	Location of	ICU CLABSI Y/N
			candidate	primary	LCBI2,		or	HAI or		attribution	
			CLABSI	(specify)	LCBI3*		MBI No	neither			
	Н										
	V										
Comme	nt:		•	•		•	•	•	•		•
	Н										
	V										
Comme	nt:										
	Н										
	V										
Comme	nt:										
	Н										
	V										
Comme	nt:										
	Н										
	V										
Comme	nt:	•	•	•	•	•	•	•	•	•	•
*LCBI 1,	2, 3	(NHSN): types of laborate	ory- confirme	ed bloodstrear	n infection	ı. MBI-LCB	I (NHSN) mu	cosal barrie	er injury LCBI. See de	efinitions in NHSN Mar	nual Chapter 4.
LCDI 1,	2, 3	(INTISIN). LYPES OF IUDOTUL	ory-conjirine	a biooustieur	ii iiijeciioi	i. WIDI-LCD	i (iviisiv) illud	Josui Dullie	i injury LCDI. See ut	zjiiii.iioiis iii ivnsiv iviul	iuui Ciiuj

Catheter-asssociated Urinary Tract Infection (CAUTI) Discrepancies

vent: first culture ate	Not candidate CAUTI	SUTI 1a, SUTI 2a, SUTI 3, SUTI 4, ABUTI*	Did not meet UTI criteria (specify below)	If UTI, Event date	POA, HAI or neither	Urethral catheter >2d?	Location of attribution	ICU CAUTI Y/N
ate		SUTI 3, SUTI 4,	criteria (specify				attribution	
	CAUTI	SUTI 3, SUTI 4,	(specify			>2d?		
		SUTI 4,	(specify					
			1					
					-			
	_							
	I (NHSN): types of sym	I (NHSN): types of symptomatic urin	3 (NHSN): types of symptomatic urinary tract infe	I (NHSN): types of symptomatic urinary tract infection. ABUT	I (NHSN): types of symptomatic urinary tract infection. ABUTI (NHSN): asymptomatic	A (NHSN): types of symptomatic urinary tract infection. ABUTI (NHSN): asymptomatic urinary tr	L(NHSN): types of symptomatic uringry tract infection. ABUTI (NHSN): asymptomatic uringry tract infection.	I (NHSN): types of symptomatic urinary tract infection. ABUTI (NHSN): asymptomatic urinary tract infection. See definitions NHSN

Surgical Site Infection (SSI) Following Colon Procedure (COLO) Discrepancies

	Date:	window closed Date:	NHSN	No SSI	SI SSI	date	Attributable	ASA [†]	Age	SW	Duration of
			procedure Y/N		DI SSI O/S SSI* (specify)		to COLO? Y/N		J	class [‡]	procedure
Н											
V	'										
Comment:											
Н											
V	'										
Comment:											
Н											
V											
Comment:											
Н											
V											
Comment:											
Н											
V											
Comment:											
Н											
V											
Comment:											
Н											
V											
Comment:											

[‡] SW class: Surgical wound class. See definitions NHSN Manual Chapter 9.

Surgical Site Infection (SSI) Following Abdominal Hysterectomy Procedure (HYST) Discrepancies

Pt. ID		Procedure Date:	Surveillance	Select One:			If SSI, Event		Optional \	Validation o	of SSI Risk Fac	tors
			window closed Date:	NHSN procedure Y/N	No SSI	SI SSI DI SSI O/S SSI*	date	Attributable to HYST? Y/N	ASA [†]	Age	SW class [‡]	Duration of procedure
						(specify)						
	Н											
C	٧											
Comme	nt:											
	Η											
	٧											
Comme	nt:											
	Н											
	٧											
Comme	nt:											
	Н											
	٧											
Comme	nt:											
	Н											
	٧											
Comme	nt:											
	Н											
	٧											
Comme	nt:											
	Н											
	٧											
Comme	nt:											
		SSI (NHSN): depth (superficial inc American Society of Anesthesiolo		ncisional, orga	n/space)	of surgical s	ite infections.					

 $^{^{\}it t}$ SW class: Surgical wound class. See definitions NHSN Manual Chapter 9.

Methicillin-resistant Staphylococcus aureus (MRSA) Bacteremia LabID Event Discrepancies

Pt. ID		Admission Date	Date of first reportable LabID Event during this inpatient stay	NHSN location of LabID Event	Positive MRSA blood culture on date of admission? Y/N	Prior MRSA blood from same location within prior 14 days? Y/N	Other reason for error
	Н						
Comme	V						
Comme							
	Н						
	V						
Comme	nt:						
	Н						
	٧						
Comme	nt:						
	Н						
	٧						
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	Н						
	٧						
Comme	nt:						

Clostridium difficile Infection (CDI) LabID Event Discrepancies

Pt. ID		Admission Date	Date of first reportable LabID Event during this inpatient stay	NHSN location of LabID Event	CDI toxin-positive result from date of admission specimen? Y/N	Prior CDI toxin-positive result from same location within prior 14 days? Y/N	Other reason for error
	Н						
Comme	V nt:						
Comme							
	Н						
	٧						
Comme	nt:						
	Н						
	٧						
Comme	nt:						
	Н						
	٧						
Comme	nt:						
	Н						
	٧						
Comme	nt:						
	Н						
	٧						
Comme	nt:						
	Н						
	٧						
Comme	nt:						
	Н						
	٧						
Comme	nt:		,	•			

This document is outdated. Reference only. Appendix 4.2: Example Validation Follow-up Letters, With and Without Identified Problems

(Courtesy of New York State Department of Health)

Please feel free to adapt these templates to meet your state's needs

Version One: Problems identified. Letter should be adapted to circumstances.

Dear CEO Name,

The [Department of Health] Healthcare Associated Infection (HAI) Reporting Program completed an audit site visit at your facility for [year] at your facility. We wish to thank you and your staff, particularly the Infection Control, Microbiology, and Medical Records staff for their cooperation and the effort they contributed during our review and audit process.

The purposes of this audit were initially presented to you in the letter of notification. Based upon our review of X medical records during the audit, there were [e.g.: X missed and unreported central line-associated bloodstream infections (CLABSIs), and X missed and unreported surgical site infections (SSIs), including (X types), and X CLABSIs and X SSIs that need to be deleted from the NHSN database].

We observed the following trends that may contribute to surveillance inaccuracies: [e.g.: Of the X colon procedure records reviewed as entered in the NHSN database, X were not NHSN colon procedures. The reporting of non-colon procedures is an infection control program surveillance system issue. In addition, infection control was not made aware of X bloodstream infections identified by the microbiology laboratory, which may have resulted in omissions.] We reviewed the reporting requirements with [Name of IP] and [she] will be reporting the missing SSIs and deleting the non-NHSN colon and HYST procedures. Each record requiring corrections was reviewed with [Name of IP] and a list of a data entry edits to be made in NHSN was provided to [her]. All data errors and missed data entry must be edited in NHSN data base within 30 days of this notice.

The infection preventionist/infection prevention manager continues to enter surgical procedure data into NHSN manually, which is a labor-intensive method for larger hospitals. Data entry could be done by a clerical person with Infection Control oversight or by electronic submission after editing of the source data for accuracy by infection control staff. Additional IT support would be required to make this possible.

We investigated your facility's notification of other hospitals when patients who underwent procedures there were admitted to your hospital with surgical site infections during the post-operative period, and we found it to be lacking. [Stipulate state requirements if they exist]. Please note that such notifications are necessary for complete surveillance of SSIs statewide, and permitted under HIPAA for the purpose of healthcare operations. We also reviewed the timeliness of your reporting and found it acceptable.

Given the issue identified with colon procedure reporting, we request your hospital review all 2015 inpatient colon procedures entered in NHSN to validate they are NHSN colon procedures. A follow- up communication as to your findings and action plans to eliminate reporting non-NHSN colon procedures should be sent to my attention no later than [Date]. Your response can be faxed or electronically sent to me. If you need any additional information or have any further questions regarding this site visit please contact me directly at [phone, fax, email].

This document is outdated. Reference only. Version two: No problems identified. Letter should be adapted to circumstances.

Dear CEO name,

The [Department of Health] Hospital Acquired Infection (HAI) Reporting Program completed an audit site visit for [year] at your facility. We wish to thank you and your staff, particularly the Infection Control, Microbiology, and Medical Records staff for their cooperation and the effort they contributed during the review and audit process.

The purposes of this audit were initially presented to you in the letter of notification. Based upon our review of X medical records, no significant compliance issues were detected. During our [date] audit, we identified [one colon surgical site infection (SSI) and two colon procedures that need to be deleted from the NHSN database]. There were no unreported infections identified in the medical records reviewed during this audit visit. We also reviewed the timeliness of reporting and have found it to be acceptable.

There continues to be only one individual, [Name], with access to manage and report in the NHSN data system. In our [specify past years] post-audit letters, we recommended to select another NHSN user to receive administrative access, to serve as a backup to the infection preventionist (IP). We continue to strongly recommend your facility add another NHSN administrative user as soon as possible. The NHSN administrative user role should be reviewed with this individual periodically during the year to ensure that your facility will be able to meet the regulatory requirements for data submission should your IP be unable to work for any reason.

We also investigated your facility's notification of other hospitals when patients who underwent procedures there were admitted to your hospital with surgical site infections during the post-operative recovery period and found it to be adequate. [Stipulate requirements if they exist]. Please note that such notifications are necessary for complete surveillance of SSIs statewide, and permitted under HIPAA for the purpose of healthcare operations.

The infection prevention manager continues to manually enter surgical procedure data into NHSN. Data entry could be done by a clerical person with Infection Control oversight. NHSN does provide for electronic submission of denominator procedure data into their reporting database and may be an option when your OR documentation becomes electronic.

We have discussed infection definitions, reporting, and data entry issues or concerns that [Name of IP] may have had, in an ongoing effort to support the [state] HAI mandatory reporting. There are some data entry corrections to be made by your staff in the NHSN reporting system. A list of each record requiring data edits was reviewed with [Name of IP]. The data entry corrections should be completed within 30 days of the audit visit.

[Name of IP] is also a member of our State HAI public reporting Technical Advisory Workgroup. I would like to take this opportunity to thank you for supporting her membership and attendance at the semiannual workshop meetings. Her contributions to this workgroup are valued by the HAI public reporting program.

If you need any additional information or a have any further questions regarding this site visit please contact me directly at [phone, fax, email.

Appendix 4.3: External Validation Data Form

State Health Department Validation Record

Page 1 of 6		<u>"</u>	requirea	""conditionally required
Facility Validation Overview				
*Facility ID:				
*Facility Type: ☐ Acute care hospital	☐ Long term acute o	are hospital (LTAC/LTCH)		
☐ Oncology hospital	☐ Inpatient rehabilita	tion facility (IRF)		
*Sampling version: ☐ CDC Version 1 (Ta	rgeted Sampling)			
*Data for year: ☐ 2015				
*HAI validated at this facility, and reason:				
☐ CLABSI (ICU, includes NICUs if a	oplicable)			
☐ CAUTI (ICU, excludes NICUs)				
□ COLO (DI/OS SSI)				
☐ HYST (DI/OS SSI)				
☐ MRSA bacteremia LabID event				
☐ CDI LabID event				
Reason:				
	□ Torrested fosility		:1:4	
☐ All facilities are validated	☐ Targeted facility	☐ 5% random sample fac	шцу	
Numerator Validation				

*Sampling information for numerator audit at this facility

Front		Sampling Frame (# elements eligible	Total # events from facility reported to NHSN for year
Event	Sampling frame elements	for review for year)	(before validation)
**ICU (including NICU) CLABSI	Medical records with positive ICU blood culture(s)		
**ICU (excluding NICU) CAUTI	Medical records with positive ICU urine culture(s)		
**DI/OS ^a COLO SSI	COLO procedures		
**DI/OS ^a HYST SSI	HYST procedures		
**MRSA bacteremia labID event	Inpatient ^b blood cultures positive for MRSA		
**CDI labID event	Inpatient ^b stools toxin-positive for C. difficile, excluding those from "baby locations"		

^aDI/OS - deep incisional or organ/space SSI

Assurance of Confidentiality: The voluntarily provided information obtained in this surveillance system that would permit identification of any individual or institution is collected with a guarantee that it will be held in strict confidence, will be used only for the purposes stated, and will not otherwise be disclosed or released without the consent of the individual, or the institution in accordance with Sections 304, 306 and 308(d) of the Public Health Service Act (42 USC 242b, 242k, and 242m(d)).

Public reporting burden of this collection of information is estimated to average 15 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).

CDC 57.600 (Front) v8.3

Distribution Inpatient includes specimens collected on day of admission from ED or other outpatient location

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rage 2 t	01 6		
	rator Validation (continued)		
	y audit results, numerators		
**Cl	LABSI in ICU (including NICU):		
	Facility determination	Audit-CLABSI Yes	Audit-CLABSI No
	Date-matched CLABSI reported	a	b
	Date-matched CLABSI NOT reported	C	d
**C/	AUTI in ICU (excluding NICU):		
	Facility determination	Audit-CAUTI Yes	Audit-CAUTI No
	Date-matched CAUTI reported	a	b
	Date-matched CAUTI NOT reported	c	d
**DI	/OS COLO SSI:		
	Facility determination	Audit-DI/OS SSI Yes	Audit-DI/OS SSI No
	Date-matched DI/OS SSI reported	a	b
	Date-matched DI/OS SSI NOT reported	c	d
**DI	/OS HYST SSI:		
	Facility determination	Audit-DI/OS SSI Yes	Audit-DI/OS SSI No
	Date-matched DI/OS SSI reported	a	b
	Date-matched DI/OS SSI NOT reported	c	d
**M	RSA bacteremia LabID event:		
	Facility determination	Audit-MRSA bacteremia culture reportable LabID event	Audit-MRSA bacteremia culture NOT reportable LabID event
	Date-matched MRSA blood culture reported as LabID event	a	b
	Date-matched MRSA blood culture NOT reported as LabID event	C	d
**CI	DI LabID event:		
	Facility determination	Audit-CDI test reportable LabID event	Audit-CDI test NOT reportable LabID event
	Date-matched CDI test reported as LabID event	a	b
	Date-matched CDI test NOT reported as LabID event	c	d

Daniel				
Denominator Valida	ation: CLABSI			
**Which method was this year?	used by this facility for ICU C	LABSI denominator (patient d	ays and centra	l line days) counting for
☐ Manual counti	ing ☐ Electronic counting	☐ Both manual and electro	nic counting	
**Has this facility con	npleted an internal validation of	of ICU CLABSI denominator d	ata for this yea	r? □ Yes □ No
Note: Validation	of manual denominator data	counting requires either:	-	
		(with more experienced coun		
		data (reference) and standard	counting for $\geq t$	hree months
	ectronic denominator data coul	nting requires: minator counting (reference) v	rs electronic da	ota for > three months
• Metrioa	C – Concurrent manual denoi	minator counting (reference) v	3. electroriic de	ita for 2 tillee months
**If yes, provide t	the following information for al	I locations and months validate	ed:	
Location of valida	ation Month of validation	Validation method	Count 1	Count 2
		A, B, or C		
		A, B, or C		
		A, B, or C		
		A, B, or C		
		A, B, or C		
If Method B is c	hosen, Count 1 should be "Us	tual Count" and Count 2 should tual Count" and Count 2 should anual Count" and Count 2 shou	d be "Patient-le	vel (Referent) Count";
If Method B is c	hosen, Count 1 should be "Us hosen, Count 1 should be "Ma	rual Count" and Count 2 should	d be "Patient-le	vel (Referent) Count";
If Method B is could be a life of the second be a life	hosen, Count 1 should be "Us hosen, Count 1 should be "Ma ation: CAUTI	rual Count" and Count 2 should	d be "Patient-le uld be "Electror	vel (Referent) Count"; nic Count."
If Method B is confirmed in the Information of the	thosen, Count 1 should be "Us thosen, Count 1 should be "Ma ation: CAUTI s used by this facility for ICU C	ual Count" and Count 2 should anual Count" and Count 2 should be s	d be "Patient-le uld be "Electron ys and cathete	vel (Referent) Count"; nic Count."
If Method B is counting the second of the se	thosen, Count 1 should be "Us thosen, Count 1 should be "Mattion: CAUTI sused by this facility for ICU Cing Electronic counting	eual Count" and Count 2 should anual Count" and Count 2 should be	d be "Patient-le uld be "Electron ys and cathete nic counting	vel (Referent) Count"; nic Count." r days) counting for this
If Method B is could like the life of the	thosen, Count 1 should be "Us thosen, Count 1 should be "Mattion: CAUTI sused by this facility for ICU Coing Electronic counting impleted an internal validation of manual denominator data	anual Count" and Count 2 should anual Count" and Count 2 should anual and electron of ICU CAUTI denominator day counting requires either:	d be "Patient-le uld be "Electron ys and cathete nic counting ta for this year?	r days) counting for this
If Method B is confirmed by the second of t	thosen, Count 1 should be "Us thosen, Count 1 should be "Mattion: CAUTI sused by this facility for ICU Coing Electronic counting impleted an internal validation of manual denominator data of A – Concurrent dual counting	anual Count" and Count 2 should anual Count" and Count 2 should anual Count 2 should anual Count 2 should anual Count 2 should anual Count denominator (patient day counting requires either:	d be "Patient-le uld be "Electron ys and cathete nic counting ta for this year?	vel (Referent) Count"; nic Count." r days) counting for this D
If Method B is countries of the Method C is considered by the Meth	thosen, Count 1 should be "Us thosen, Count 1 should be "Mattion: CAUTI sused by this facility for ICU Coing Electronic counting impleted an internal validation of manual denominator data of A – Concurrent dual counting IB – Concurrent patient level of the should be a concurrent patient level of the should be "Us and be "Us an	anual Count" and Count 2 should anual Count" and Count 2 should anual Count 2 should anual Count 2 should anual Count 2 should anual Count and Electron of ICU CAUTI denominator data counting requires either: I (with more experienced count at a (reference) and standard	d be "Patient-le uld be "Electron ys and cathete nic counting ta for this year?	vel (Referent) Count"; nic Count." r days) counting for this D
If Method B is coll Method C is coll Method C is coll Method C is coll Method Was year? Manual counting the Method Method Validation of electric Method Collidation of electric Method C is collidati	thosen, Count 1 should be "Us thosen, Count 1 should be "Mattion: CAUTI sused by this facility for ICU Counting In Electronic counting In the property of the	anual Count" and Count 2 should anual Count" and Count 2 should anual Count 2 should anual Count 2 should anual Count" and Count denominator (patient data counting requires either: I (with more experienced count data (reference) and standard nting requires:	d be "Patient-le uld be "Electron ys and cathete nic counting ta for this year? ter as reference counting for ≥ t	vel (Referent) Count"; nic Count." r days) counting for this P □ Yes □ No e) for ≥ three months OF three months
If Method B is coll Method C is coll Method C is coll Method C is coll Method Was year? Manual counting **Has this facility consistent Method • Method Validation of election Method • Method Method • Method	thosen, Count 1 should be "Use thosen, Count 1 should be "Mattion: CAUTI is used by this facility for ICU Counting in Electronic counting in Electronic counting in the Electronic denominator data in the Electronic denominator data counting in Ele	anual Count" and Count 2 should anual Count" and Count 2 should anual Count 2 should anual Count" and Count 2 should anual Counting and electron of ICU CAUTI denominator data counting requires either: I (with more experienced counting and standard anting requires: minator counting (reference) with anual counting and standard anting requires:	d be "Patient-le uld be "Electron ys and cathete nic counting ta for this year? ter as reference counting for ≥ to rs. electronic da	vel (Referent) Count"; nic Count." r days) counting for this P □ Yes □ No e) for ≥ three months OF three months
If Method B is confirmed by the second by t	thosen, Count 1 should be "Us thosen, Count 1 should be "Mathemation: CAUTI sused by this facility for ICU Coing Electronic counting impleted an internal validation of manual denominator data of A – Concurrent dual counting IB – Concurrent patient level of the concurrent manual denominator data counting IC – Concurrent manual d	anual Count" and Count 2 should anual Count" and Count 2 should anual Count 2 should anual Count" and Count 2 should anual Count and electron of ICU CAUTI denominator data counting requires either: If (with more experienced count and the counting requires and standard anting requires: If (mith more experienced count and the counting requires) with the counting requires and the counting (reference) with the counting (reference) with the counting (reference) with the counting and months validated.	d be "Patient-le uld be "Electron ys and cathete nic counting ta for this year? ter as reference counting for ≥ to ss. electronic dated:	vel (Referent) Count"; nic Count." r days) counting for this P □ Yes □ No e) for ≥ three months OF three months ata for ≥ three months
If Method B is coll Method C is coll Method C is coll Method C is coll Method Was year? Manual counting **Has this facility consistent Method • Method Validation of election Method • Method Method • Method	thosen, Count 1 should be "Us thosen, Count 1 should be "Mathemation: CAUTI sused by this facility for ICU Coing Electronic counting impleted an internal validation of manual denominator data of A – Concurrent dual counting IB – Concurrent patient level of the concurrent manual denominator data counting IC – Concurrent manual d	anual Count" and Count 2 should anual Count" and Count 2 should anual Count 2 should anual Count" and Count 2 should anual Counting requires either: a (with more experienced counting requires: minator counting (reference) with more counting (refe	d be "Patient-le uld be "Electron ys and cathete nic counting ta for this year? ter as reference counting for ≥ to rs. electronic da	vel (Referent) Count"; nic Count." r days) counting for this P □ Yes □ No e) for ≥ three months OF three months
If Method B is confirmed by the second by t	thosen, Count 1 should be "Us thosen, Count 1 should be "Mathemation: CAUTI sused by this facility for ICU Coing Electronic counting impleted an internal validation of manual denominator data of A – Concurrent dual counting IB – Concurrent patient level of the concurrent manual denominator data counting IC – Concurrent manual d	EAUTI denominator (patient da Dount 2 should anual Count" and Count 2 should anual Count" and Count 2 should be accounted an and electron of ICU CAUTI denominator data counting requires either: If (with more experienced counting requires: minator counting (reference) with the counting (reference) with the counting and months validated Validation method A, B, or C	d be "Patient-le uld be "Electron ys and cathete nic counting ta for this year? ter as reference counting for ≥ to ss. electronic dated:	vel (Referent) Count"; nic Count." r days) counting for this P □ Yes □ No e) for ≥ three months OF three months ata for ≥ three months
If Method B is could Method C is considered by the Method C is considered by the Method was year? □ Manual counting the Method •	thosen, Count 1 should be "Us thosen, Count 1 should be "Mathemation: CAUTI sused by this facility for ICU Coing Electronic counting impleted an internal validation of manual denominator data of A – Concurrent dual counting IB – Concurrent patient level of the concurrent manual denominator data counting IC – Concurrent manual d	anual Count" and Count 2 should anual Count" and Count 2 should anual Count and Count 2 should anual Count" and Count 2 should anual Counting requires either: a (with more experienced counting requires: minator counting (reference) with the cou	d be "Patient-le uld be "Electron ys and cathete nic counting ta for this year? ter as reference counting for ≥ to ss. electronic dated:	vel (Referent) Count"; nic Count." r days) counting for this P □ Yes □ No e) for ≥ three months OF three months ata for ≥ three months
If Method B is co. If Method C is co. Denominator Valida **Which method was year? Manual counti **Has this facility con. Note: Validation. Method Validation of ele. Method **If yes, provide to.	thosen, Count 1 should be "Us thosen, Count 1 should be "Mathemation: CAUTI sused by this facility for ICU Coing Electronic counting impleted an internal validation of manual denominator data of A – Concurrent dual counting IB – Concurrent patient level of the concurrent manual denominator data counting IC – Concurrent manual d	anual Count" and Count 2 should anual Count" and Count 2 should anual Count and Count 2 should anual Count" and Count 2 should anual Counting requires either: a (with more experienced counting requires: minator counting (reference) with the cou	d be "Patient-le uld be "Electron ys and cathete nic counting ta for this year? ter as reference counting for ≥ to ss. electronic dated:	vel (Referent) Count"; nic Count." r days) counting for this P □ Yes □ No e) for ≥ three months OF three months ata for ≥ three months
If Method B is co. If Method C is co. Denominator Valida **Which method was year? Manual counti **Has this facility con. Note: Validation. Method Validation of ele. Method **If yes, provide to.	thosen, Count 1 should be "Us thosen, Count 1 should be "Mathemation: CAUTI sused by this facility for ICU Coing Electronic counting impleted an internal validation of manual denominator data of A – Concurrent dual counting IB – Concurrent patient level of the concurrent manual denominator data counting IC – Concurrent manual d	anual Count" and Count 2 should anual Count" and Count 2 should anual Count and Count 2 should anual Count" and Count 2 should anual Counting requires either: a (with more experienced counting requires: minator counting (reference) with the cou	d be "Patient-le uld be "Electron ys and cathete nic counting ta for this year? ter as reference counting for ≥ to ss. electronic dated:	vel (Referent) Count"; nic Count." r days) counting for this P □ Yes □ No e) for ≥ three months OF three months ata for ≥ three months

If Method A is chosen, Count 1 should be "Usual Count" and Count 2 should be "Expert (Referent) Count"; If Method B is chosen, Count 1 should be "Usual Count" and Count 2 should be "Patient-level (Referent) Count"; If Method C is chosen, Count 1 should be "Manual Count" and Count 2 should be "Electronic Count."

Page 4 or 6 Denominator Validation: COLO					
**Document number of COLO procedures from two systems by month:					
	Month	Number of COLO procedures entered into NHSN by facility before validation Number of ICD-9 procedure codes for COLO identified from hospital discharge billing			
Denomir	nator Validat	on: HYST			
**Docum	ent number o	f HYST procedures from two	o systems by mo	onth:	
	Month Number of HYST procedures entered into NHSN by facility before validation Number of ICD-9 procedure codes for HYST identified from hospital discharge billing				
		on: MRSA bacteremia Lab	oID event & CDI	LabID event	
NHSN inpatient location validation **Do any inpatient locations require mapping or re-mapping within NHSN? \[\triangle \tr					
**Do any inpatient locations require mapping or re-mapping within NHSN? \square Yes \square No **If yes, indicate which locations need to be mapped/re-mapped and recommendations:					
Loca		Current CDC location code designation	Current bed count	Recommended CDC location code designation	Recommended bed count
**How do	es this facility	obtain inpatient admissions	s data?		
□ Ele	☐ Electronic from billing ☐ Electronic from vendor system ☐ Electronic from ADT			ADT	
□ Other (specify):					
**How does this facility obtain inpatient patient days data?					
☐ Electronic from billing ☐ Electronic from vendor system ☐ Electronic from ADT					
□ Other (specify):					

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Denominator Validation: MRSA bacteremia LabID event & CDI LabID event (continued)						
**Has this facility completed any internal validation of LabID event denominator data counting?						
	□ Yes □ No Note: Validation of denominator data counting requires concurrent patient level denominator counting (reference) vs. standard electronic data for three specified location types [one ICU, one LDRP if available, and one or more wards where observation patients are frequently housed] for ≥1 month; validated data should fall within 5% of the reference standard (see validation Guidance and Toolkit Appendix 1).					
*	*If yes, provide the follo	wing information for a	Il months validate	ed:		
		MRS	A bacteremia La			. =
				ssions		t Days
-	Location of validation	Month of validation	Usual count	Manual count	Usual count	Manual count
-						
-						
-						
L					l	
			CDI LabID eve	ent ^c		
			Admi	ssions	Patien	t Days
	Location of validation	Month of validation	Usual count	Manual count	Usual count	Manual count
_						
-						
c_ L	al des feet des des de					
ΕX	cludes 'baby locations'					
Ris	k Adjustment Variable	• Validation				
	U mapping (ICU CLAB		CU CAUTI [exclu	des NICUs])		
	Number of ICU location	ns correctly mapped as	s ICUs in NHSN (includes NICUs):		
Number of locations incorrectly mapped as ICUs (includes NICUs):						
Number of ICUs (includes NICUs) omitted from ICU mapping: Number of ICU mapping errors (ICUs vs. non-ICUs):						
**T6	eaching hospital affiliation	on (ICU CLABSI, ICU	CAUTI, MRSA ba	acteremia LabID e	event, CDI LabID	event)
Facility teaching hospital affiliation reported on 2015 NHSN annual facility survey:						
	□ Non-teaching □ Major □ Graduate □ Undergraduate □ N/A (IRF & LTAC)					
	Is facility teaching hosp	ital affiliation correct?	□ Yes	□ No		
** A		-\				
A	SA score (COLO, HYS) Number (% of audited)					
	Number (% of audited)					

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Risk Adjustment Variable Validation (continued)					
**Patient age (COLO, HYST)					
Number (% of audited) correct for COLO:					
Number (% of audited) correct for HYST:					
**Facility bed size (all inpatient locations, including 'baby loca Facility bed size reported on 2015 NHSN annual facility s Validated bed size:					
Custom Fields					
Label	Label				
					
Comments					

Appendix 5: Facility/Provider to Facility/Provider Communications under HIPAA: Questions and Answers

Note: The following document was developed by CDC scientists and lawyers in collaboration with HHS Office of Civil Rights (OCR) program and legal staff, who oversee administration of the Health Insurance Portability and Accountability Act of 1996 (HIPAA). This information may not be modified without express permission of OCR.

Facility/Provider to Facility/Provider Communications under HIPAA: Questions and Answers

Health care providers [i.e., individual clinicians and facilities (including hospitals and other health care facilities such as nursing homes and rehabilitation facilities)] are increasingly active in addressing concerns about patient safety and minimizing patients' risks of adverse healthcare events. In an era when the public, policymakers, and many health care providers seek greater transparency and accountability in healthcare, these efforts include but are not limited to new or renewed emphasis on information sharing among providers themselves about adverse events that are a consequence of a care process, care process omission, or some other risk exposure during a health care episode, such as exposure to an infectious agent.

Health care providers have raised questions as to whether the HIPAA Privacy Rule permits information sharing between individual providers and/or facilities for patient safety-related purposes. This guidance assumes that the provider seeking to share such patient information is a HIPAA covered entity. While any health care provider may be faced with these questions, they tend to arise more frequently at the facility level. The term "patient" is also used here to encompass persons residing in nursing homes or other facilities, where they are often referred to as "residents." "Source facility" or "source provider" refers to the health care facility or individual provider that first cared for the patient. Protected health information ("PHI") is individually identifiable health information, such as information that identifies (or can be used to identify) a patient.

Question One

Does HIPAA permit a health care facility to share PHI with the source facility where a patient was previously treated or where a patient previously resided, without the patient's authorization, for purposes of providing notification of an infection with potential infection control implications at the source facility?

In these scenarios a resident of a nursing home is admitted into a hospital, certain medical conditions are diagnosed, and the hospital wants to disclose this health information back to the nursing home.

- A practitioner at the hospital diagnoses a patient's tuberculosis and wants to inform the nursing home so that the staff there can quarantine the coughing roommate of the index case.
- The patient is admitted with sepsis and later dies in the hospital. Blood cultures drawn at admission grow group A streptococcus. The hospital seeks to disclose that this patient was diagnosed with invasive group A streptococcal infection (which causes serious outbreaks in nursing homes) to the nursing home for infection control purposes, even though the patient will not be returning.
- The hospital diagnoses the patient with influenza early in the flu season and wants to disclose this diagnosis to the nursing home for infection control purposes.

In each scenario the hospital will want to disclose the name of the patient so the nursing home can verify that this patient had been a resident in their home and the date and location of service.

Answer One

The HIPAA Privacy Rule permits a covered health care provider to <u>use or disclose</u> PHI for treatment purposes without the authorization of the patient. (Generally, disclosures of psychotherapy notes require written patient authorization, but these notes do not appear relevant here.) 45 CFR 164.506(c) and 164.508(a)(2). "Treatment" is defined to include the provision, coordination, or management of "health care" and related services. 45 CFR 164.501. "Health care" is defined to include preventive care. 45 CFR 160.103. Treatment refers to activities undertaken on behalf of individual patients. While in most cases, the information regarding an individual is needed for the treatment of that individual, the HIPAA Privacy Rule also allows the information regarding one individual (e.g., a patient) to be used or disclosed for the treatment or preventive care (e.g., vaccinations or quarantine) of other persons (e.g., patients at risk).

In these scenarios, the patient (and former nursing home resident) has or had a medical condition while at the nursing home that may directly impact the health of certain or all residents at that facility. In some cases, the nursing home did not know of this condition, or the condition had not manifested itself at the time the patient was at the nursing home. The hospital may disclose PHI of the patient (and former nursing home resident) to the nursing home for treatment purposes involving other residents.

A distinction is made between <u>use</u> and <u>disclosure</u> of PHI for treatment purposes with regard to the "minimum necessary" requirement. The "minimum necessary" requirement does <u>not</u> apply to <u>disclosures</u> of PHI for treatment purposes, and the disclosures discussed above are treatment disclosures that are permitted under the HIPAA Privacy Rule.

After PHI is disclosed to the nursing home, the information may be <u>used</u> for the provision of treatment to the nursing home residents. For example, preventive measures, such as cohorting, isolation, or prophylaxis of specific patients who may be at risk at the nursing home, are considered treatment under the Privacy Rule. The <u>uses</u> of PHI by the nursing home for treatment purposes in the above scenarios <u>are</u> subject to the Privacy Rule's "minimum necessary" requirement, and the nursing home's minimum necessary policies. A nursing home, as a covered entity, must identify those persons or classes of persons in its workforce who need access to PHI, and for each such person or classes of person, the category or categories of PHI to which access is needed, and any conditions appropriate to such access. 45 CFR 164.514(d)(2). For more information on the "minimum necessary" requirement,

see: http://www.hhs.gov/ocr/privacy/hipaa/fag/minimum necessary/207.html.

Question Two

Under HIPAA, is a health care facility permitted to share PHI with another health care facility that previously treated or housed a patient, without that patient's authorization, for purposes of notifying this source facility of a potential complication of care related to the health care provided at the source facility so as to monitor and improve care and prevent future complications?

- A hospital identifies a surgical site infection (SSI) that is probably attributable to an ambulatory surgical
 care facility and/or surgeon that performed the surgery within the past 12 months. The hospital seeks
 to notify the ambulatory surgical care facility about the SSI, or in a given situation, notify the surgeon
 directly.
- A patient is admitted to Hospital B with a surgical site infection (SSI) after an operation at another hospital (Hospital A), where the patient had been operated on and then discharged without signs or symptoms of infection. Because of federal requirements (e.g., the Centers for Medicare and Medicaid

Services' Inpatient Quality Reporting program requirements) or state law or policy, both hospitals are committed to reporting all SSIs following the type of operation performed on the patient. Hospital B seeks to report the SSI to Hospital A, where the SSI is presumed to have originated, so that Hospital A can fully account for SSIs attributable to its care.

Answer Two

The HIPAA Privacy Rule permits a covered entity to use or disclose PHI for certain "health care operations" purposes without the authorization of the patient. 45 CFR 164.506(c). This includes a covered entity disclosing PHI to another covered entity for certain purposes if each entity either has or had a relationship with the individual who is the subject of the information, and the PHI being disclosed pertains to the relationship. 45 CFR 164.506(c)(4). Of relevance here, disclosures are permitted for the purpose of the covered entity receiving the information "conducting quality assessment and improvement activities; . . . population-based activities relating to improving health [and] protocol development." 45 CFR 164.501 (definition of "health care operations"). Only the minimum amount of PHI necessary for the particular health care operations purpose may be disclosed.

The disclosures discussed above are health care operations disclosures that are permitted under the HIPAA Privacy Rule. In these scenarios we assume that the hospitals sharing the PHI, the ambulatory surgical care facility, and the surgeon are all HIPAA covered entities. The hospitals disclosing the PHI would be sharing information regarding a patient who the surgical facilities (either the ambulatory care facility or the hospital) and/or surgeon had treated, and the communication is in regard to the treatment that had been provided. The disclosures are so that the surgical facilities and/or surgeon can monitor and improve the quality of care provided. This falls under "conducting quality assessment and improvement activities," and perhaps "population-based activities relating to improving health," and/or "protocol development." In these scenarios, information regarding the patient with an SSI can be shared with the surgical facilities and/or surgeon. While only the minimum amount of information regarding the patient may be disclosed, in these scenarios the identity of the patient may be shared because it is needed to investigate the cause of the infections (e.g., the dates and locations of care, and the staff involved.) There is likely to be no need to share health information regarding these patients that is unrelated to investigating the SSI.

For additional information regarding disclosures for treatment and healthcare operations purposes, see: http://www.hhs.gov/ocr/privacy/hipaa/understanding/coveredentities/usesanddisclosuresfortpo.html.