
How To Assess Risk to Patients When There Is a Disinfection and Sterilization Failure

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Disclosure

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Risk Analysis of Disinfection and Sterilization Failures

- Overview of disinfection and sterilization principles
- Failure scenarios
- Recommended protocol for exposure evaluation

ORIGINAL ARTICLE

How to Assess Risk of Disease Transmission to Patients When There Is a Failure to Follow Recommended Disinfection and Sterilization Guidelines

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BACKGROUND. Disinfection and sterilization are critical components of infection control. Unfortunately, breaches of disinfection and sterilization guidelines are not uncommon.

OBJECTIVE. To describe a method for evaluating a potential breach of guidelines for high-level disinfection and sterilization of medical devices.

METHODS. The appropriate scientific literature was reviewed to determine the frequency of failures of compliance. A risk assessment model was constructed.

RESULTS. A 14-step protocol was constructed to aid infection control professionals in the evaluation of potential disinfection and sterilization failures. In addition, a model is presented for aiding in determining how patients should be notified of the potential adverse event. Sample statements and letters are provided for communicating with the public and individual patients.

CONCLUSION. Use of a protocol can guide an institution in managing potential disinfection and sterilization failures.

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In the United States in 1996, there were approximately 46,500,000 surgical procedures and a much larger number of infection failure on record involved the distribution of an inactive lot of glutaraldehyde disinfectant solution that had



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Failure to Follow Disinfection and Sterilization Principles

● Overview

- Achieving disinfection and sterilization through the use of disinfection and sterilization practices is essential for ensuring that medical and surgical instruments do not transmit pathogens to patients
- Deficiencies leading to infection have occurred when there has been failure to follow disinfection and sterilization principles
- These failures resulted from human error, equipment failures or system problems
- Discuss a 14 step method for managing a failure incident

Disinfection and Sterilization Principles

Disinfection and Sterilization

EH Spaulding believed that how an object will be disinfected depended on the object's intended use.

CRITICAL - objects which enter normally sterile tissue or the vascular system or through which blood flows should be **sterile**.

SEMICRITICAL - objects that touch mucous membranes or skin that is not intact require a disinfection process (**high-level disinfection[HLD]**) that kills all microorganisms but high numbers of bacterial spores.

NONCRITICAL -objects that touch only intact skin require **low-level disinfection**.

Efficacy of Disinfection/Sterilization

Influencing Factors

Cleaning of the object

Organic and inorganic load present

Type and level of microbial contamination

Concentration of and exposure time to disinfectant/sterilant

Nature of the object

Temperature and relative humidity

Critical Patient Care Objects

Processing “Critical” Patient Care Objects

Classification:	Critical objects enter normally sterile tissue or vascular system, or through which blood flows.
Object:	Sterility.
Level germicidal action:	Kill all microorganisms, including bacterial spores.
Examples:	Surgical instruments and devices; cardiac catheters; implants; etc.
Method:	Steam, gas, hydrogen peroxide plasma, hydrogen peroxide vapor, ozone or chemical sterilization.

Critical Objects

- Surgical instruments
- Cardiac catheters
- Implants

Recommendations

Methods of Sterilization

- Steam is preferred for critical items not damaged by heat
- Follow the operating parameters recommended by the manufacturer
- Use low temperature sterilization technologies for reprocessing critical items damaged by heat

Chemical Sterilization of “Critical Objects”

Glutaraldehyde ($\geq 2.0\%$)

Hydrogen peroxide-HP (7.5%)

Peracetic acid-PA (0.2%)

HP (1.0%) and PA (0.08%)

HP (7.5%) and PA (0.23%)

Glut (1.12%) and Phenol/phenate (1.93%)

Glut (3.4%) and Isopropanol (26%)

Exposure time per manufacturers' recommendations

Semicritical Patient Care Objects

Processing “Semicritical” Patient Care Objects

Classification:	Semicritical objects come in contact with mucous membranes or skin that is not intact.
Object:	Free of all microorganisms except high numbers of bacterial spores.
Level germicidal action:	Kills all microorganisms except high numbers of bacterial spores.
Examples:	Respiratory therapy and anesthesia equipment, GI endoscopes, thermometer, etc.
Method:	High-level disinfection

Semicritical Items

- Endoscopes
- Respiratory therapy equipment
- Anesthesia equipment
- Endocavitary probes
- Tonometers
- Diaphragm fitting rings

High Level Disinfection of “Semicritical Objects”

Exposure Time \geq 8 m-30m (US), 20°C

<u>Germicide</u>	<u>Concentration</u>
Glutaraldehyde	\geq 2.0%
Ortho-phthalaldehyde (12 m)	0.55%
Hydrogen peroxide*	7.5%
Hydrogen peroxide and peracetic acid*	1.0%/0.08%
Hydrogen peroxide and peracetic acid*	7.5%/0.23%
Hypochlorite (free chlorine)*	650-675 ppm
Accelerated hydrogen peroxide	2.0%
Glut and isopropanol	3.4%/26%
<u>Glut and phenol/phenate**</u>	<u>1.21%/1.93%</u>

*May cause cosmetic and functional damage; **efficacy not verified

Noncritical Patient Care Objects

Processing “Noncritical” Patient Care Objects

Classification:	Noncritical objects will not come in contact with mucous membranes or skin that is not intact.
Object:	Can be expected to be contaminated with some microorganisms.
Level germicidal action:	Kill vegetative bacteria, fungi and lipid viruses.
Examples:	Bedpans; crutches; bed rails; EKG leads; bedside tables; walls, floors and furniture.
Method:	Low-level disinfection

Low-Level Disinfection for “Noncritical” Objects

Exposure time \geq 1 min

Germicide

Use Concentration

Ethyl or isopropyl alcohol

70-90%

Chlorine

100ppm (1:500 dilution)

Phenolic

UD

Iodophor

UD

Quaternary ammonium

UD

Accelerated hydrogen peroxide

0.5%

UD=Manufacturer’s recommended use dilution

Disinfection and Sterilization of Emerging Pathogens

Disinfection and Sterilization of Emerging Pathogens

- Hepatitis C virus
- *Clostridium difficile*
- *Cryptosporidium*
- *Helicobacter pylori*
- *E.coli* 0157:H7
- Antibiotic-resistant microbes (MDR-TB, VRE, MRSA)
- SARS Coronavirus, avian influenza, norovirus
- Bioterrorism agents (anthrax, plague, smallpox)

Disinfection and Sterilization of Emerging Pathogens

Standard disinfection and sterilization procedures for patient care equipment are adequate to sterilize or disinfect instruments or devices contaminated with blood and other body fluids from persons infected with emerging pathogens

Endoscopes/AERS

Murphy Was an ICP!

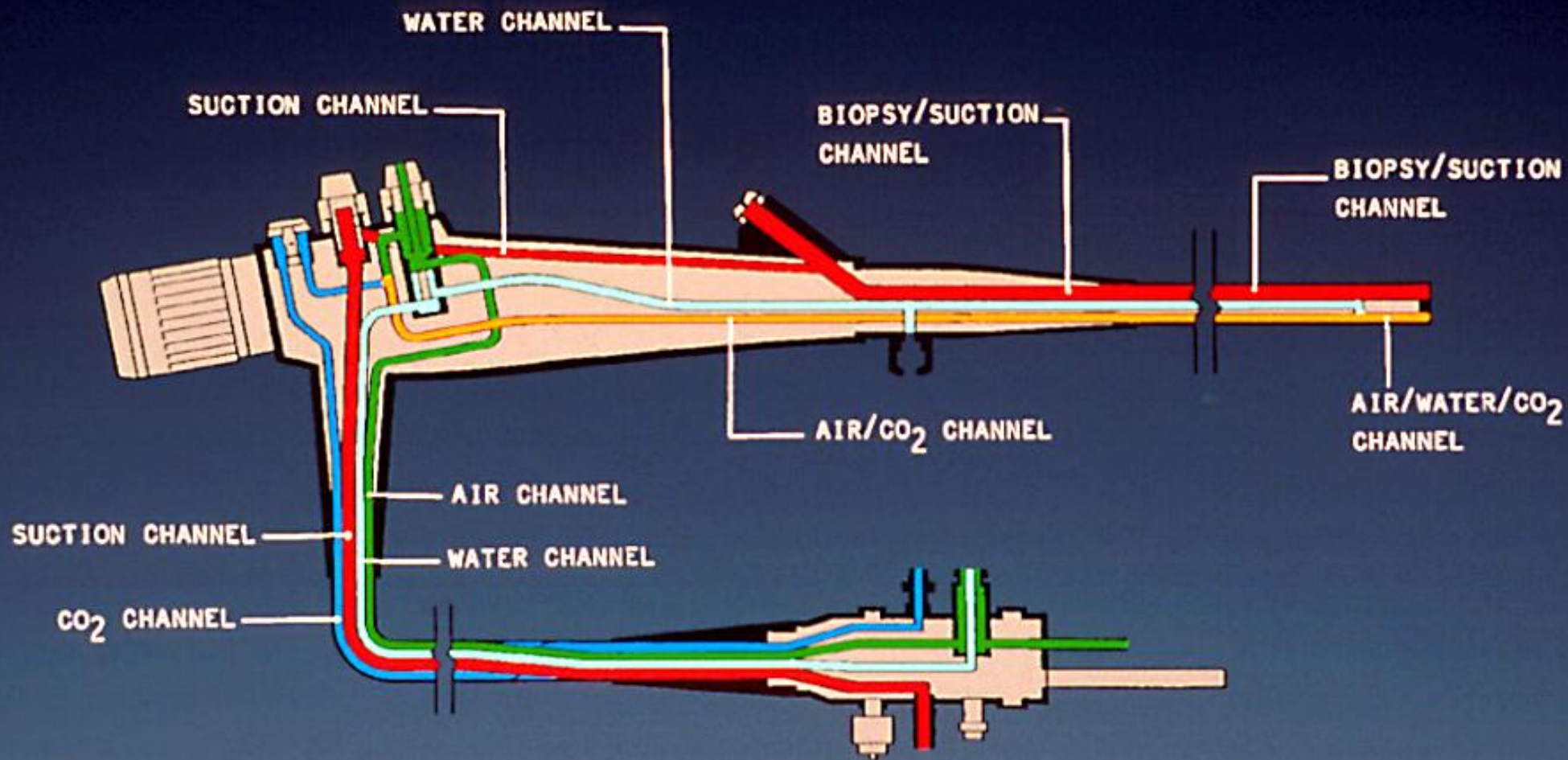
Murphy's Law

“Whatever can go wrong will go wrong”

Corollary

“...in the worst possible way at the worst possible time”

ENDOSCOPE CHANNELS



GI ENDOSCOPES AND BRONCHOSCOPES

- Widely used diagnostic and therapeutic procedure
- Endoscope contamination during use (GI 10^9 in/ 10^5 out)
- Semicritical items require high-level disinfection minimally
- Inappropriate cleaning and disinfection has lead to cross-transmission
- In the inanimate environment, although the incidence remains very low, endoscopes represent a risk of disease transmission

TRANSMISSION OF INFECTION

- Gastrointestinal endoscopy
 - >300 infections transmitted
 - 70% agents *Salmonella sp.* and *P. aeruginosa*
 - Clinical spectrum ranged from colonization to death (~4%)
- Bronchoscopy
 - 90 infections transmitted
 - *M. tuberculosis*, atypical *Mycobacteria*, *P. aeruginosa*

Spach DH et al Ann Intern Med 1993; 118:117-128 and Weber DJ, Rutala WA Gastroint Dis 2002;87

ENDOSCOPE INFECTIONS

- Infections traced to deficient practices
 - Inadequate cleaning (clean all channels)
 - Inappropriate/ineffective disinfection (time exposure, perfuse channels, test concentration)
 - Failure to follow recommended disinfection practices (tapwater rinse)
 - Flaws in design of endoscopes or AERs

ENDOSCOPE DISINFECTION

- CLEAN-mechanically cleaned with water and enzymatic cleaner
- HLD/STERILIZE-immerse scope and perfuse HLD/sterilant through all channels for at least 12 min
- RINSE-scope and channels rinsed with sterile water, filtered water, or tap water followed by alcohol
- DRY-use forced air to dry insertion tube and channels
- STORE-prevent recontamination

Disinfection and Sterilization

Conclusions

- When properly used, disinfection and sterilization can ensure the safe use of invasive and non-invasive medical devices.
- Method of disinfection and sterilization depends on the intended use of the medical device
- Cleaning should always precede high-level disinfection and sterilization
- Current disinfection and sterilization guidelines must be strictly followed.

Failure to Follow Disinfection and Sterilization Principles

Failure to Follow Disinfection and Sterilization Principles

- These events are relatively frequent; however, not commonly appreciated
- Human errors
 - Time setting of 132°C steam sterilizer at 0 min rather than 4 min
 - Failure to sterilize items after cleaning
 - Exposure time on AER set at 5 min rather than 20 min for glut
- Equipment failures-biopsy port caps not secure
- System problems-unwrapped specula

TABLE 1. Reprocessing Failures of Semicritical or Critical Medical Instruments Resulting in Patient Notification

Location or institution, year	Instrument involved	No. of persons exposed
Sacramento, CA, 2002	Endoscope	750
Toronto, ON, 2003	Endoscope	146
Seattle, WA, 2004	Endoscope	600
Sacramento, CA, 2004	Endoscope	1,331
San Francisco, CA, 2004	Endoscope	2,000
Long Island, NY, 2004	Endoscope	177
Charleston, NC, 2004	Endoscope	1,383
Toronto, ON, 2003	Prostate biopsy probe	900
Pittsburgh, PA, 2005	Endoscope	200
Leesburg, VA 2005	Endoscope	144
San Diego, CA, 2006	Endoscope	300
Augusta, ME, 2006	Prostate biopsy needle	481
Dept Veterans Affairs, 2006	Prostate biopsy equipment	2,075
San Diego, CA, 2006	Surgical instrument	82

NOTE. Modified from a presentation by Douglas Nelson, MD, at the 33rd Annual Conference and International Meeting of the Association for Professionals in Infection Control and Epidemiology; Tampa, Florida, 2006.

Failure to Follow Disinfection and Sterilization Principles

- Method for assessing patient risk for adverse events
- Although exposure events are often unique, can approach the evaluation of potential failure using a standardized approach
- Propose a sequence of 14 steps that form a general approach to a possible failure of disinfection/sterilization (D/S)
- D/S failure could result in patient exposure to an infectious agent

Failure to Follow Disinfection and Sterilization Principles

Scenario:

Hospital A has been purchased an AER for GI endoscope reprocessing. The AER has been in use for 9 months. The hospital was using >2% glutaraldehyde with an intended exposure time of 20 minutes. It was discovered that the exposure time was incorrectly set at 10 minutes. Endoscopes for 9 months were processed at 10 minutes rather than the recommended 20 minutes.

What Do You Do?



Failure to Follow Disinfection and Sterilization Principles

Scenario:

Hospital B discovered that for the past 3 days all surgical instruments were exposed to steam sterilization at 132°C for 0 minutes rather than the intended 4 minutes. A central processing technician turned the timer to 0 minutes in error.

What Do You Do?

1. Confirm disinfection or sterilization reprocessing failure
2. Impound any improperly disinfected/sterilized items
3. Do not use the questionable disinfection/sterilization unit (e.g., sterilizer, automated endoscope reprocessor) until proper functioning can be assured
4. Inform key stakeholders
5. Conduct a complete and thorough evaluation of the cause of the disinfection/sterilization failure
6. Prepare a line listing of potentially exposed patients
7. Assess whether disinfection/sterilization failure increases patient risk for infection
8. Inform expanded list of stakeholders of the reprocessing issue
9. Develop a hypothesis for the disinfection/sterilization failure and initiate corrective action
10. Develop a method to assess potential adverse patient events
11. Consider notification of state and federal authorities
12. Consider patient notification
13. Develop long-term follow-up plan
14. Perform after-action report

FIGURE 1. Protocol for exposure investigation after a failure of disinfection and sterilization procedures

Failure to Follow Disinfection and Sterilization Principles

- Step 1-confirm failure
 - Confirm that the suspected failure did, in fact, occur.
 - ICP must review the circumstances of the reported failure including: the time and date of the possible failure; type of D/S method; and evidence of process parameters (printout) and results of physical, chemical and/or biological indicators

Failure to Follow Disinfection and Sterilization Principles

- Step 1-confirm failure

- If the initial evaluation reveals that no medical items that were potentially inadequately processed were used in patient care, there is no patient safety issue involved
- Then one can limit the evaluation to determining if the disinfection/sterilization **process failed and correcting the processing error**
- All potentially inadequately processed items must, of course, be reprocessed
- If a disinfection/sterilization failure is not confirmed, the investigation may be concluded

Failure to Follow Disinfection and Sterilization Principles

- Step 2-embargo improperly D/S items
 - If a D/S failure has occurred, one should immediately embargo any medical items that may not have been appropriately D/S
 - All items since the last successful processing (as demonstrated by process measures and/or physical, chemical, or biological indicators) should be embargoed.
 - Retrieving all items may require visiting all areas where the medical/surgical items may be stored or used including CP, ORs, community-based practices, storerooms, etc



Orth Carolina Hospitals
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STERILIZED S1
INDEFINITE SHELF LIFE
UNLESS
DAMAGED OR OPENED

Failure to Follow Disinfection and Sterilization Principles

- Step 3-do not use questionable D/S item
 - The incriminated D/S item should be immediately placed off line and not used for D/S of medical or surgical devices until its proper function can be assured
 - This may involve several runs with assessment of process parameters and physical, chemical and/or biological indicators
 - Medical engineering or the manufacturer's representative usually performs repairs and evaluation of the unit

Failure to Follow Disinfection and Sterilization Principles

- Step 4-inform key stakeholders
 - All key stakeholders should be informed of the problem
 - ◆ Risk management
 - ◆ Medical/nursing director of the involved units (e.g., OB, GI)
 - ◆ Personnel involved in disinfection/sterilization
 - It is often easier to arrange a face-to-face conference to assure complete transmission of the facts with feedback than to use email or telephone consultation

Failure to Follow Disinfection and Sterilization Principles

- Step 5-investigate the cause of the D/S problem
 - A complete and thorough evaluation of the possible D/S failure should be rapidly completed.
 - ICP should review the exact circumstances of the possible D/S failure including dates and results of all process measures (e.g., temperature, time, sterilant/HLD concentration) and physical, chemical and biological indicators obtained in the recent past going back far enough to assess the time/date of the first possible malfunction

Failure to Follow Disinfection and Sterilization Principles

- Step 6-line listing of exposed patients
 - Once a failure of D/S has been documented, it is important to initiate the evaluation of potential patient exposures
 - First step is to create a line listing of all possible patients who may have been exposed to possibly contaminated medical/surgical devices
 - ◆ Patient name, identification number, date(s) of exposure, contaminated device used, underlying risk factors for infection, development of HAIs (pathogen, body site), and other potentially adverse events

Failure to Follow Disinfection and Sterilization Principles

- Step 7-does D/S failure increase patient risk for infection
 - Once a failure of D/S process has been documented with possible exposure to a contaminated item, it is crucial to determine whether in fact the failure could result in an adverse patient event.
 - For example, 3 min for flash sterilization rather than 4 min. Would not consider 3 min flash sterilization cycle as representing a patient hazard.
 - Assessing risk should always include on a review of the scientific literature and national guidelines

Failure to Follow Disinfection and Sterilization Principles

- Step 8-inform expanded list of stakeholders
 - All stakeholders should be informed of the progress of the investigation, especially if an increased risk to patients is possible or documented
 - ◆ Risk management
 - ◆ Medical/nursing director of the involved units (e.g., OB, GI)
 - ◆ Personnel involved in disinfection/sterilization
 - ◆ Public relations, healthcare administration, and legal
 - A press release should be prepared in case of need and a spokesperson appointed

The ABC Hospital announced today that it is contacting patients because of a sterilizer malfunction. The Department of Hospital Epidemiology (Infection Control) was notified on July 7 that sterilizers runs conducted on July 5 and July 6 were performed at a reduced temperature (250°F versus 270°F). The sterilizer was used to process surgical instruments.

ABC Hospital has taken the following actions:

- Upon notification of the malfunction, Hospital Epidemiology immediately took the sterilizer out of use.
- As chemical indicators (used to check temperature) are included in each sterilizer run, Hospital Epidemiology has reviewed the results of these indicators.
- All surgical instruments sterilized in the malfunctioning unit were impounded and will be resterilized.
- All patients who were operated on using instruments processed in this unit are being notified by phone and registered mail.
- A "hot line" has been set up to answer any questions by patients. The number is -----.

Our staff has fully evaluated the impact of the reduced temperature on the sterilization of surgical instruments. We are committed to providing the highest quality care to our patients and to notifying them of any mishap even if we believe it poses no risk. Based on our review of the processing procedure and the scientific literature, ABC Hospital believes that this mishap will not lead to any increased risk of infection by patients operated on using instruments processed in the malfunctioning unit.

- All surgical instruments were mechanically cleaned in a washer-disinfector that is highly effective in removing bacteria (ie, removes >99.9%).
- Unpublished studies have shown that the time and temperature (250°F for 5 minutes in a gravity steam sterilizer) used at ABC Hospital inactivates high numbers of bacterial pathogens (ie, inactivates >99.999%).
- Studies have shown that the microbial load associated with cleaned surgical instruments is low (ie, 85% of surgical instruments contain less than 100 organisms).

We have stressed to our patients that it is unlikely that the surgical instruments processed at reduced temperature represent any increased risk. However, in any procedure, there is always a risk of infection. Patients who develop any symptoms of infection (fever, redness, drainage, warmth) should immediately contact their surgeon.

ABC Hospitals regrets that this event occurred and is taking steps to assure that this problem does not occur again.

FIGURE 4. Sample press release regarding the potential exposure event caused by the sterilization failure

Failure to Follow Disinfection and Sterilization Principles

- Step 9-develop hypothesis for D/S failure and initiate corrective action
 - Corrective actions (e.g., reset timer, monitor concentration of HLD) should be initiated to correct the deficiencies in reprocessing
 - Reprocessing of any item that may not have been appropriately disinfected/sterilized must be done

Failure to Follow Disinfection and Sterilization Principles

- Step 10-assess adverse patient events
 - Initiate a more detailed study, if necessary, of possible adverse outcomes in patients
 - This may entail designing a prospective cohort study
 - This may require reviewing medical records and/or examining patients for infections, chemical reactions, or other adverse events
 - Specific laboratory tests may be necessary such as testing source patients and exposed persons for bloodborne pathogens such as HIV, HBV, and HCV

Failure to Follow Disinfection and Sterilization Principles

- Step 11

- In conjunction with the legal department, notify state and federal authorities if required by regulation or law

Failure to Follow Disinfection and Sterilization Principles

- Step 12-consider patient notification
 - Consider whether patients should be notified of the disinfection/sterilization failure
 - If it is determined the failure could result in adverse patient events, then patients should be notified
 - Determine who will notify the patients
 - ◆ Patient's local medical provider, risk management, attending physician at the time of failure, ICP
 - One should develop a script to be used in notification to ensure all patients receive the same information

I am Dr....., may I please speak to..... I am calling to tell you that during your recent Obstetrics-Gynecology examination at, you may have been examined using a speculum that was cleaned and disinfected but not high-level disinfected. We believe that the chance you could get any infection from this infection is extremely unlikely, almost certainly less than 1 in a billion. However, if you are concerned, we would be happy to see you at and provide free testing for viruses that could theoretically have been transmitted during your examination. We would offer testing now and again in 3 and 6 months

The viruses we are talking about include HIV, and the viruses that cause hepatitis B and hepatitis C. We can also give you the hepatitis B vaccine if you desire. Again, all tests and the vaccine will be provided free.

We regret this occurred and have taken steps to assure that in the future all medical devices are both cleaned and disinfected before use.

FIGURE 2. Sample script to be used when discussing potential exposure with a patient by telephone

Failure to Follow Disinfection and Sterilization Principles

- Step 12 (continued)
 - Notification may be accomplished by a face-to-face meeting, phone or registered mail
 - More than one method may be used to ensure complete notification
 - Notification should include: an assessment of risk, possible adverse events that may occur, symptoms and signs of the adverse event, time period for the adverse event, risk to other contacts, possible prophylactic therapy (risks and benefits) and recommended medical follow-up

Dear (Patient's Name):

ABC Hospital routinely monitors the effectiveness of its sterilizers through the use of physical, chemical and biological indicators. Biological indicators are used to validate sterilization efficacy.

This letter is intended to notify you that on July 5 and 6 sterilizer runs were conducted at a reduced temperature (250°F versus 270°F). Surgical packs processed at this reduced temperature may have been used in surgical services (if only certain services specify). Chemical indicators are packaged in each tray and in all cases these indicators demonstrated processing. Two biological indicators were included in runs at the reduced temperature and both were positive demonstrating that all spores were not inactivated.

We believe that the reduced temperature used in the sterilizer is unlikely to have an impact upon patients operated upon using equipment processed at the reduced temperature for the following reasons.

- All chemical indicators used demonstrated an acceptable result.
- Unpublished studies have shown that the time and temperature (250°F for 5 minutes in gravity steam sterilizer) used at ABC Hospital inactivates high numbers of bacterial pathogens (ie, 3,000,000 *Staphylococcus aureus*)(Unpublished data, Rutala and Weber, May 2006).
- The surgical instruments were mechanically cleaned in a washer-disinfector that is effective in inactivating/removing bacteria.
- Studies have shown that the microbial load associated with decontaminated surgical instruments is low (ie, 85% of surgical instruments contain less than 100 organisms)^{a, b}. The microbial load associated with surgical instruments is at least 1000 less than the microbial load contained in the biological indicator.
- The organism contained in the biological indicator, *Geobacillus stearothermophilus*, is far more resistant to steam sterilization than the bacteria present on surgical instruments before processing.
- The scientific literature supports the statement that our current standard operating time-temperature is conservatively chosen.

The sterilizer in question has been repaired and is now operating at the standard temperature of 270°F. All remaining surgical packs, which were processed at the reduced temperature, have been recalled and reprocessed.

Again, we believe it is unlikely that the surgical packs processed at reduced temperature represent any increased risk of infection to you. However, in any procedure, there is always a risk of infection. Should you have any symptoms of infection, including fever or increased redness, swelling or pain near the operative site, you should notify....

We regret this occurred and have taken steps to assure that this problem does not occur again. If you have any questions or concerns, please contact...

Failure to Follow Disinfection and Sterilization Principles

- Step 12 (continued)
 - The healthcare facility must decide who will provide these services and whether the facility will cover the cost of care.
 - In general, we believe that if the facility was responsible for the failure then it should provide these services at no patient charge
 - However, if the exposure resulted from failures outside the institution (receipt by the facility of inadequately sterilized devices), then the facility may want to offer the services but at patient expense or causative party's expense (e.g., manufacturer)

Failure to Follow Disinfection and Sterilization Principles

Scenario:

Endoscopes were processed in an AER using OPA for 2m rather than the recommended 12m. The HLD exposure time in the AER was incorrectly reset by the biomedical engineer to 2m 10 months ago. During that period of time there were 2,134 patients that received GI endoscopy procedures. All the endoscopes were reprocessed in the same AER.

Failure to Follow Disinfection and Sterilization Principles

Can estimate the per patient risk for HIV as follows:

- HIV prevalence in the US population: 0.37%, ~4:1000, $\sim 4 \times 10^{-3}$
- Risk of transmission (via mm): 0.09%, 1:1000, 1×10^{-3}
- Efficacy of AER without HLD: 99.999%, 1:100,000, 1×10^{-3}
- Efficacy of OPA against HIV in 2m: 99.999%, 1:100,000, 1×10^{-5}
- Effect of HIV drying: 99%, 1:100, 1×10^{-2}
- Individual risk = $\sim 4 \times 10^{-16}$ (4 in 10 quadrillion)

Failure to Follow Disinfection and Sterilization Principles

- How about if you were able to conduct a risk assessment and the risk for infection was 4 in 10 quadrillion
 - There is no fixed or accepted frequency that necessitates risk disclosure.
 - ◆ Hospital could conclude that the risk frequency of 4 in 10 quadrillion is so small that they are effectively, legally, of no weight or less than the risk of many other daily life exposures we all endure
 - ◆ Hospital could conclude that all exposures should be communicated to the patient regardless of the 4 in 10 quadrillion risk for an adverse event
 - Decision to inform patients is made by the hospital stakeholders

TABLE 2. Lifetime Odds of Death Due to Selected Types of Injury, United States, 2002

Type of injury or event	Lifetime odds of death
Transportation accident	1 in 77
Pedestrian	1 in 612
Car occupant	1 in 228
Drowning	1 in 1,081
Fall	1 in 229
Exposure to smoke, fire, flames	1 in 1,179
Venomous snake or lizard bite	1 in 1,241,661
Accidental poisoning	1 in 212
Lightning	1 in 56,439
Flood	1 in 413,887
Intentional self-harm	1 in 118

NOTE. National Safety Council estimates based on data from the National Center for Health Statistics and the U.S. Census Bureau. To determine the odds per year, multiply by 77.3 years (eg, the annual odds of dying as a result of injury caused by lightning are 1 in 4,362,735).

Failure to Follow Disinfection and Sterilization Principles

- Step 13-develop long term follow-up plan
 - Once the problem leading to the D/S failure has been identified and corrective action initiated, it is essential to assess whether these interventions have eliminated the problem over the long-term
 - This may require long-term surveillance, changes in current policies or procedures, development of new policies or procedures, evaluation of current equipment, etc

Failure to Follow Disinfection and Sterilization Principles

- Step 14-perform after-action report
 - A report of the event should be prepared for presentation to the appropriate healthcare system committees
 - Consideration should be given to publishing the evaluation if it provides a contribution to the scientific literature

Failure to Follow Disinfection and Sterilization Principles

Summary

- Follow the 14 steps-they provide a general outline, but each event is unique and you must be flexible and adaptable
- Steps are delineated in a linear fashion but the evaluation is often done simultaneously
- Communication among key stakeholders is very important
- Ethical to notify patients if there is a risk-should be upfront and factual
- Train staff and access processes/practices to minimize recurrence
- These are stressful events (patients and staff) but the goal is to assess failure and protect patients rather than assessing blame

Risk Analysis of Disinfection and Sterilization Failures

- Overview of disinfection and sterilization principles
- Failure Scenarios
- Recommended Protocol for Exposure Evaluation

Thank you



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