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INTRODUCTION

Of the patients with end-stage renal disease (ESRD) treated by maintenance dialysis in the United States, approximately 90% are on maintenance hemodialysis and 10% are on peritoneal dialysis. Maintenance hemodialysis patients are at higher risk for infection, because uremia is known to make patients with ESRD more susceptible to infectious agents through defects in cellular immunity, neutrophil function, and complement activation. In addition, because the process requires vascular access for long periods in an environment where multiple patients receive hemodialysis concurrently, repeated opportunities exist for transmission of infectious agents. Patient-to-patient transmission of infectious agents, directly or indirectly through contaminated devices, equipment, supplies, injectable medications, environmental surfaces, or hands of healthcare personnel have all been demonstrated. Furthermore, hemodialysis patients require frequent hospitalizations and surgery, which increases their opportunities for exposure and risk for developing healthcare-associated infections. This chapter describes (1) the major infectious diseases that can be acquired in the dialysis center setting, (2) important epidemiological and environmental microbiological considerations, and (3) infection control strategies.

FACTORS CONTRIBUTING TO INFECTIONS AMONG HEMODIALYSIS PATIENTS

Technical development and clinical use of hemodialysis delivery systems improved dramatically in the late 1960s and early 1970s. However, a number of microbiological parameters were not accounted for in the design of many hemodialysis machines and their respective water supply systems. There are many situations where certain types of gram-negative water bacteria can persist and actively multiply in hemodialysis water supplies and aqueous environments associated with hemodialysis equipment. This can result in massive numbers of gram-negative bacteria, which can directly or indirectly lead to septicemia or endotoxemia. These bacteria can adhere to surfaces and form biofilms (glycocalyces), which are virtually impossible to eradicate. Control strategies are designed not to eradicate bacteria but to reduce their concentration to relatively low levels and to prevent their regrowth.

Although certain genera of gram-negative water bacteria (e.g., Burkholderia, Flavobacterium, Pseudomonas, Ralstonia, Serratia, Stenotrophomonas maltophilia, and Sphingomonas) are most commonly encountered, virtually any bacterium...
that can grow in water can be a problem in a hemodialysis unit. Several species of nontuberculous mycobacteria may also contaminate water treatment systems, including *Mycobacterium chelonae*, *M. abscessus*, *M. fortuitum*, *M. gordonae*, *M. mucogenicum*, *M. scrofulaceum*, *M. kansasii*, *M. avium*, and *M. intracellulare*; these microorganisms do not contain bacterial endotoxin but are comparatively resistant to chemical germicides.

Gram-negative water bacteria can multiply even in water containing relatively small amounts of organic matter, such as water treated by distillation, softening, deionization, or reverse osmosis, reaching levels of $10^5$ to $10^7$ microorganisms/mL; these levels are not associated with visible turbidity. When treated water is mixed with dialysis concentrate, the resulting dialysis fluid is a balanced salt solution and growth medium almost as rich in nutrients as conventional nutrient broth. Gram-negative water bacteria growing in dialysis fluids can reach levels of $10^8$ to $10^9$ microorganisms/mL, producing visible turbidity.

Bacterial growth in water used for hemodialysis depends on the types of water treatment system used, dialysate distribution systems, dialysis machine type, and method of disinfection (Table 25.1).

### TABLE 25.1 Factors Influencing Microbial Contamination in Hemodialysis Systems

| Factors                                      | Comments                                                                 |
|----------------------------------------------|--------------------------------------------------------------------------|
| **Water Supply (Water Source)**              |                                                                          |
| Groundwater                                  | Contains endotoxin and bacteria                                          |
| Surface water                                | Contains high levels of endotoxin, bacteria, and other organisms        |
| **Water Treatment at the Dialysis Center**   |                                                                          |
| None                                         | Not Recommended                                                          |
| Filtration                                   |                                                                          |
| Prefilter                                    | Particulate filter to protect equipment; does not remove microorganisms  |
| Absolute filter (depth or membrane)          | Removes bacteria but unless changed frequently or disinfected, bacteria will accumulate and grow through the filter; acts as a significant reservoir of bacteria and endotoxin |
| Granular activated carbon (GAC)              | Removes organics and available chlorine or chloramine; significant reservoir of water bacteria and endotoxin |
| **Water Treatment Devices**                  |                                                                          |
| Ion exchange (softener, deionization)        | Softeners and deionizers remove cations and anions, contaminants from source water; significant reservoir for bacteria and endotoxin |
| Reverse osmosis (RO)                         | Removes bacteria, endotoxin, chemicals, and must be cleaned and disinfected; most systems employed for dialysis applications operate under high pressure |
| Ultraviolet (UV) germicidal irradiator       | Kills most bacteria, but there is no residual; some UV-resistant bacteria can develop |
| Ultrafilter                                   | Removes bacteria and endotoxin; operates on normal line pressure; can be positioned distal to storage tank and deionizer; must be disinfected or changed |
| **Water and Dialysate Distribution System**  |                                                                          |
| Distribution pipes                           | Oversized diameters and length decrease fluid flow and increases bacterial reservoir in the form of biofilms for both treated water and central delivery systems (bicarbonate concentrate or bicarbonate dialysate) |
| Materials                                    | Pipe materials influence bacterial colonization and biofilm formation, as well as what types of chemical disinfectants can be used |
| Construction                                 | Rough joints, dead ends, and unused branches can act as bacterial reservoirs |
| Elevation                                    | Outlet taps should be located at highest elevation to prevent loss of disinfectant |
| Storage tanks                                | Generally undesirable because of large surface area and can act as a reservoir for water bacteria; a properly designed tank can minimize this risk |
| **Dialysis Machines**                        |                                                                          |
| Single pass                                  | Disinfectant should have contact time with all parts of the machine that are in contact with treated water or dialysate |
| Recirculating single pass, or recirculating batch | Recirculating pumps and machine design allow for massive contamination levels if not properly disinfected; overnight disinfection recommended |

Bacterial contamination of water used for hemodialysis may also contaminate water treatment systems, including *Mycobacterium chelonae*, *M. abscessus*, *M. fortuitum*, *M. gordonae*, *M. mucogenicum*, *M. scrofulaceum*, *M. kansasii*, *M. avium*, and *M. intracellulare*; these microorganisms do not contain bacterial endotoxin but are comparatively resistant to chemical germicides. Each component is discussed separately next.

## Microbial Contamination of Water

Water used for the production of dialysis fluid must be treated to remove chemical and microbial contaminants. The Association for the Advancement of Medical Instrumentation (AAMI) published guidelines and recommended practices for the chemical and microbial quality of water used to prepare dialysis fluid and reprocess hemodialyzers (Table 25.2). The Centers for Medicare and Medicaid Services (CMS) has incorporated into their ESRD facility conditions for coverage infection control requirements that dialysis facilities need to follow, including water quality standards. Some components of the water treatment system may allow for amplification of water bacteria. For example, ion exchangers such as...
TABLE 25.2 AAMI Microbial Quality Standards for Dialysis Fluids

| Type of Fluid               | MICROBIAL BIOBURDEN | ENDOTOXIN |
|----------------------------|---------------------|-----------|
|                            | Maximum Contaminant  | Action Level | Maximum Contaminant  | Action Level |
|                            | Level (CFU/mL)      |            | Level (EU/mL)        |             |
| Water for all purposes     | 100                   | 50 CFU/mL | 0.25 EU/mL           |             |
| Conventional dialysate     | 100                   | 50 CFU/mL | 0.50 EU/mL           |             |
| Ultrapure dialysate        | 1 CFU/10 mL          | 0.03 EU/mL | 0.25 EU/mL           |             |
| Dialysate for infusion*    | This online process shall be validated by the manufacturer to produce fluid that is sterile and nonpyrogenic. | |

*Compliance with a maximum bacterial level of 10⁻⁶ CFU/mL cannot be demonstrated by culturing, but by processes developed by the machine manufacturers.

as water softeners and deionizers do not remove endotoxin or microorganisms and provide many sites for significant bacterial multiplication.³⁵ Granular activated carbon adsorption media (i.e., carbon filters) are used primarily to remove certain organic compounds and available chlorine (free and combined) from water, but they also significantly increase the level of water bacteria, yeast, fungi, and endotoxins.

A variety of filters are marketed to control bacterial contamination of water and dialysis fluids. Most are inadequate, especially if they are not routinely disinfected or frequently changed. Particulate filters, commonly called prefilters, operate by depth filtration and do not remove bacteria or endotoxin. These filters can become colonized with gram-negative water bacteria, resulting in higher levels of bacteria and endotoxin in the filter effluent. Absolute filters, including membrane types, temporarily remove bacteria from passing water. However, some of these filters tend to clog, and gram-negative water bacteria can “grow through” the filter matrix and colonize downstream surfaces of the filters within a few days. Further, absolute filters do not reduce levels of endotoxin in the effluent water. These filters should be changed regularly in accordance with the manufacturer’s directions and disinfected in the same manner and at the same time as the rest of the water distribution system.

Ultraviolet germicidal irradiation (UVGI) is sometimes used to reduce microbial contamination in water, but the use of UVGI has some special considerations. The lamp should be appropriately sized for the flow rate of water passing through the device, and the energy output should be monitored to ensure effectiveness of the lamp. Manufacturers of the lamp may require routine replacement schedule. Some bacterial populations may develop resistance to UVGI. In recirculating dialysis distribution systems, repeated exposure to UVGI are used to ensure adequate disinfection; however, this approach allows for progressive removal of sensitive microorganisms and selection of UVGI-resistant organisms. In addition, bacterial endotoxins are not affected.

Reverse osmosis (RO) is an effective water treatment modality that is used in more than 97% of US hemodialysis centers. RO possesses the singular advantage of being able to remove a variety of substances, including microorganisms and endotoxins, from supply water based primarily on particle size and adsorption to the membrane. However, low numbers of gram-negative and acid-fast organisms may penetrate the membrane or by other means (leaks around seals), and colonize downstream portions of the water distribution system. Consequently, the RO unit must be disinfected routinely.

We recommend a water treatment system that produces chemically adequate water while avoiding high levels of microbial contamination. The components in a typical water system should include (1) prefilters, (2) a water softener, (3) carbon adsorption tanks (at least two in series), (4) a particulate filter (to protect the reverse osmosis membrane), and (5) an RO unit. If one includes a deionization unit as a polisher (post–reverse osmosis unit) and a storage tank, the final component should be an ultrafilter to remove microorganisms and endotoxin. As the incoming tap water passes through the system components, it becomes more chemically pure, but the level of microbial contamination increases, which is why ultrafiltration and RO are important. Additional components or processes may be included in the pretreatment chain (see Table 25.1) depending on the pH, potable water disinfectant, and chemical quality of the incoming municipal water. If the system is adequately disinfected and properly maintained, the microbial content of water should be well within the recommended limits.

Distribution Systems

Water that has passed through the water distribution system (product water) is then distributed to individual dialysis machines where it is combined with dialysate concentrates and to a reprocessing area if a facility reprocesses hemodialyzers. It may also be combined with concentrates at a central location where the resulting dialysis fluid is supplied to the individual machines. Plastic pipe (most often polyvinyl chloride) is then used to distribute water, or dialysis fluids to the dialysis machines. Distribution systems should include the use of a loop-based system and no dead-ended pipes. Outlets to dialysis machines should have a relatively short path with the least amount of fittings and the use of valves with minimal dead space. Voids, dead ends, and large surface areas serve as sites for microbial colonization. Also large diameter pipes decrease fluid velocity and increase the wetted surface area available for microbial colonization. In addition, long pipe runs also increase the available surface area for colonization.
Gram-negative water bacteria in fluids remaining in pipes overnight can rapidly multiply and colonize wetted surfaces of the distribution system, producing microbial populations and endotoxin in quantities proportional to the total volume of the surface area. Such colonization results in the formation of protective biofilm, which is difficult to remove and protects the bacteria and other organisms from disinfection. Continuous circulation of water slows down this process.

Disinfection of the water or dialysate distribution system should be performed on a regular basis so that the microbial quality of the fluids is within the acceptable standards range. The frequency of disinfection should be validated by each facility and should be performed after any changes or modifications to the system. AAMI standards and recommended practices are community consensus standards, and do not specify a schedule for disinfection other than to suggest that routine disinfection be conducted. In many instances, microbiological monitoring can be used to determine the frequency of testing of disinfection of the distribution system. In some circumstances, repeat disinfection of the system cannot adequately control microbial growth because of established biofilm and replacement of the system is the only option.

To prevent disinfectant from draining from pipes by gravity before adequate contact time, distribution systems should be designed with all taps at equal elevation and at the highest point of the system. Furthermore, the system should be free of rough joints and dead-end pipes. Fluid trapped in such stagnant areas can serve as reservoirs for bacteria and fungi that later contaminate the rest of the distribution system.

Storage tanks greatly increase the volume of fluid and surface area of the distribution system. If used, these should be designed with a conical bottom so that water exits the storage tank at its lowest point (and allows the tank to be drained), be fitted with a tight-sealing lid, be equipped with a spray head, and possess an air vent containing a bacteriological filter. If used, the storage tanks should be routinely cleaned, disinfected, and drained. To remove biofilm, use of strong oxidizers may aid in stripping biofilm from surfaces; however, physical scrubbing of the inner surfaces of the tank may be necessary. When using a storage tank, an ultrafilter should be incorporated before water is pumped into the distribution system.

**Hemodialysis Machines, Effluent, and Environmental Surfaces**

In the 1970s, most dialysis machines were of the recirculating or recirculating single-pass type; their design contributed to relatively high levels of gram-negative bacterial contamination in dialysis fluid. Virtually all dialysis machines in the United States now are single-pass machines (i.e., the dialysate flows through the machine once). Single-pass machines tend to respond to adequate cleaning and disinfection procedures and, in general, have lower levels of bacterial contamination than do recirculating machines. Levels of contamination in single-pass machines depend primarily on the microbiological quality of the incoming water and the method of machine disinfection. Earlier dialysis machines had a port (waste-handling option) that allowed disposal of the extracorporeal circuit priming fluids. If one-way check valves in the waste-handling option are not maintained, checked for competency, or disinfected as recommended, it allows backflow from the effluent dialysate path into and contamination of the port and the attached bloodline. This led to outbreaks of infections among hemodialysis patients. The waste-handling option is much less commonly used now.

The external surfaces of dialysis machines and components are also likely sources for contamination. These include frequently touched surfaces (e.g., the control panel, dialysis chairs, keyboard, shared charting computers), attached priming buckets used during the priming of the dialyzers, blood tubing draped or clipped to waste containers, or other equipment brought into the station. For example, among nine outbreaks of bacteremia, fungemia, and pyrogenic reactions not related to dialyzer reuse investigated by the Centers for Disease Control and Prevention (CDC), inadequate disinfection of the water distribution system or dialysis machines was implicated in seven (Table 25.3). Surface contamination has been described as a potential contributor to transmission of bloodborne pathogens in the context of other poor practices. A novel source of transmission has been identified: dialysis wall boxes, which contain several connections that allow the dialysis machine to hook up to the water supply and drain effluent. A large outbreak of bloodstream infections caused by *Serratia marcescens*, *Pseudomonas aerugiosa*, *Enterobacter cloacae*, and other gram-negative bacteria was identified and wall boxes were determined to be the source. More work is needed to understand the role of wall boxes and other surfaces and related infection control aspects in transmission of pathogens.

**Hemodialyzer Reuse**

Reuse of disposable hollow-fiber dialyzers in the United States increased between 1976 and 1982, from 18% to 43% of facilities reporting reuse; the highest percentage was 82% in 1997. By 2002 the percentage of facilities reporting reusing dialyzers had declined to 63%. Recent data from the CDC’s National Healthcare Safety Network indicated that only 1.8% of facilities reported reuse in 2017 (CDC, unpublished data). This decline coincides with decisions made by several large dialysis organizations to discontinue the practice of reuse and to only use single-use dialyzers. Although dialyzer reuse is still common in developing countries, it has become less popular in developed countries and some have plans to phase out this practice. After a series of outbreaks of bacterial infections associated with reuse and reprocessing of dialyzers, CDC recommended that single use of dialyzers be the preferred practice and stated that it should be used whenever possible.

In 1986, AAMI standards for reprocessing hemodialyzers were adopted by the United States Public Health Service and was incorporated into regulation by CMS. In the United States, dialyzer reuse has not been associated with the transmission of bloodborne pathogens such as hepatitis B (HBV), hepatitis C (HCV), or human immunodeficiency...
| Description | Cause(s) of Outbreak | Corrective Measure(s) Recommended | Reference |
|-------------|---------------------|----------------------------------|-----------|
| Bacteremia, Fungemia, or Pyrogenic Reactions Not Related to Dialyzer Reuse | | | |
| Pyrogenic reactions in 49 patients | Untreated city water contained high levels of endotoxin | Install a reverse osmosis system | 4 |
| Pyrogenic reactions in 45 patients | Inadequate disinfection of the fluid distribution system | Increase disinfection frequency and contact time | 40 |
| Pyrogenic reactions in 14 patients; 2 cases of bacteremia; 1 death | Reverse osmosis water storage tank contaminated with bacteria | Remove or properly maintain and disinfect the storage tank | 28 |
| Pyrogenic reactions in 6 patients; 7 cases of bacteremia | Inadequate disinfection of water distribution system and dialysis machines; improper microbial assay procedure | Use correct microbial assay procedures; disinfect water treatment system and dialysis machines following manufacturer’s recommended procedures | 302 |
| Bacteremia in 35 patients with central venous catheters (CVCs) | CVCs used as facilities’ primary vascular access; median duration of infected catheters was 311 days; improper aseptic techniques | Uses CVCs when only absolutely necessary for vascular access; use appropriate aseptic technique when inserting and performing routine catheter care | 303 |
| Three pyrogenic reactions and 10 cases of bacteremia in patients treated on machines with a port for disposal of dialyzer priming fluid (waste handling option [WHO] port) | Incompetent check valves allowing backflow of fluid from the waste side of the machine into attached blood tubing; bacterial contamination of the WHO | Routine disinfection and maintenance of the dialysis machine including the WHO; check competency of WHO before patient treatment | 41 |
| Bacteremia in 10 patients treated on machines with WHO port | Incompetent backflow to allow backflow from dialysate effluent side of the machine in the WHO port and attached bloodlines | Routine maintenance, disinfection, and check for check valve competence of the WHO port | 42 |
| Outbreak of pyrogenic reactions and gram-negative bacteremia in 11 patients | Water distribution system and machines were not routinely disinfected according to manufacturer’s recommendations | Disinfect machines according to manufacturer’s recommendations; include reverse osmosis water distribution system in the weekly disinfection schedule; microbiological assay should be performed via membrane filtration or spread plate using Trypticase soy agar | 9 |
| *Phialemonium curvatum* access infections in four dialysis patients; two of these patients died of systemic disease | Observations at the facility noted some irregularities in site preparation for needle insertion | Review infection control practices clean and disinfect HVAC system where water accumulated; perform surveillance on all patients | 304 |
| *P. curvatum* bloodstream infection (BSI) in two patients | All affected patients had synthetic grafts | Conduct routine maintenance and disinfection of machines and WHO ports; redesign water system to eliminate dead legs; have a routine schedule for disinfection of the water system | 43 |
| | One environmental sample was positive for *P. curvatum* (condensate pan of HVAC serving the unit) | | |
| | Water and dialysate samples were cultured using a calibrated loop and blood agar plates—results always indicated no growth | | |
| | Water system and dialysis machines with WHO ports not routinely maintained; water system contained dead legs and laboratory used wrong assays | Improve infection control practices, cleaning and disinfection of wall boxes | 47 |
| | | | |
| Outbreak of gram-negative BSI in 58 patients | Poor infection control practices, contamination from dialysis wall boxes | | |
| | | | |
| Bacteremia/Pyrogenic Reactions Related to Dialyzer Reprocessing | | | |
| Mycobacterial infections in 27 patients | Inadequate concentration of dialyzer disinfectant | Increase formaldehyde concentration used to disinfect dialyzers to 4% | 22 |
| Mycobacterial infections in five high-flux dialysis patients; two deaths | Inadequate concentration of dialyzer disinfectant and inadequate disinfection of water treatment system | Use higher concentration of peracetic acid for reprocessing dialyzers and follow manufacturer’s labeled recommendations; increase frequency of disinfecting the water treatment system | 26 |
| Bacteremia in six patients | Inadequate concentration of dialyzer disinfectant; water used to reprocess dialyzers did not meet AAMI standards | Use AAMI-quality water; ensure proper germicide concentration in the dialyzer | CDC unpublished data |

Continued
**TABLE 25.3 Outbreaks of Dialysis-Associated Illnesses Investigated by the Centers for Disease Control and Prevention, 1975–2016—cont’d**

| Description                                                                                                                                  | Cause(s) of Outbreak                                                                 | Corrective Measure(s) Recommended                                                                 | Reference |
|--------------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------|-----------|
| Bacteremia and pyrogenic reactions in six patients                                                                                           | Dialyzer disinfectant diluted to improper concentration                              | Use disinfectant at the manufacturer’s recommended dilution and verify concentration             | 60        |
| Bacteremia and pyrogenic reactions in six patients                                                                                           | Inadequate mixing of dialyzer disinfectant                                            | Thoroughly mix disinfectant and verify proper concentration                                     | 10        |
| Bacteremia in 33 patients at 2 dialysis centers                                                                                              | Dialyzer disinfectant created holes in the dialyzer membrane                         | Change disinfectant (product was withdrawn from the market by the manufacturer)                | 305, 306  |
| Bacteremia in six patients; all blood isolates had similar plasmid profiles                                                                 | Dialyzers were contaminated during removal and cleaning of headers with gauze; staff not routinely changing gloves; dialyzers not reprocessed for several hours after disassembly and cleaning | Do not use gauze or similar material to remove clots from header; change gloves frequently; process dialyzers after rinsing and cleaning | 59        |
| Pyrogenic reactions in three high-flux dialysis patients                                                                                     | Dialyzer reprocessed with two disinfectants; water for reuse did not meet AAMI standards | Do not disinfect dialyzers with multiple germicides; more frequent disinfection of water treatment system and conduct routine environmental monitoring of water for reuse | 307       |
| Pyrogenic reactions in 14 high-flux dialysis patients; 1 death                                                                            | Dialyzers rinsed with city (tap) water containing high levels of endotoxin; water used to reprocess dialyzers did not meet AAMI standards | Do not rinse or reprocess dialyzers with tap water; use AAMI-quality water for rinsing and preparing dialyzer disinfectant | 308       |
| Pyrogenic reactions in 18 patients                                                                                                           | Dialyzers rinsed with city (tap) water containing high levels of endotoxin; water used to reprocess dialyzers did not meet AAMI standards | Do not rinse or reprocess dialyzers with tap water; use AAMI-quality water for rinsing and preparing dialyzer disinfectant | 11        |
| Pyrogenic reactions in 22 patients                                                                                                           | Water for reuse did not meet AAMI standards; improper microbiological technique was used on samples collected for monthly monitoring | Use the recommended assay procedure for water analysis of water and dialysate; disinfect water distribution system | 8         |
| Bacteremia and candidemia among patients in seven dialysis units (in Minnesota and California)                                              | Dialyzers were not reprocessed in a timely manner; some dialyzer refrigerated for extended periods before reprocessing; company made changes to header cleaning protocol | Reprocess dialyzers as soon as possible; follow joint CDC and dialyzer reprocessing equipment and disinfectant manufacturer guidance for cleaning and disinfecting headers of dialyzer | CDC unpublised Data |
| Outbreak of gram-negative BSI, including *Burkholderia cepacia* and *Stenotrophomonas maltophilia* in 17 patients                          | *B. cepacia* was isolated from header cleaning machine matched patient isolates Contamination likely was due to incomplete disinfection during reprocessing | Reuse was stopped                                                                               | 52        |
| **Transmission of Viral Agents**                                                                                                            | **Leakage of coil dialyzer membranes and use of recirculating bath dialysis machines** | **Separation of HBsAg+ patients and equipment from all other patients**                          | 262       |
| 26 patients seroconverted to HBsAg+ during a 10-month period                                                                            | No specific cause determined; false-positive HBsAg results caused some susceptible patients to be dialyzed with infected patients | Laboratory confirmation of HBsAg+ results; strict adherence to glove use and use of separate equipment for HBsAg+ patients | 309       |
| 19 patients and 1 staff member seroconverted to HBsAg+ during a 14-month period                                                            | Staff not wearing gloves; surfaces not properly disinfected; improper handling of needles/sharps resulting in many staff needlestick injuries | Separation of HBsAg+ patients and equipment from susceptible patients; proper precautions by staff (e.g., gloves; handling of needles and sharps) | 262       |
| 24 patients and 6 staff seroconverted to HBsAg+ during a 10-month period                                                                  | Extrinsic contamination of intravenous medication being prepared adjacent to an area where blood samples were handled | Separate medication preparation area from area where blood processing for diagnostic tests is performed | 267       |
| 13 patients and 1 staff member seroconverted to HBsAg+ during a 1-month period                                                             | Extrinsic contamination of multidose medication vial shared by HBsAg+ and HBsAg-susceptible patients | No sharing of supplies, equipment, and medications between patients (CDC, unpublished data) |           |
### TABLE 25.3 Outbreaks of Dialysis-Associated Illnesses Investigated by the Centers for Disease Control and Prevention, 1975–2016—cont’d

| Description | Cause(s) of Outbreak | Corrective Measure(s) Recommended | Reference |
|-------------|----------------------|-----------------------------------|-----------|
| Seven patients seroconverted to HBsAg+ during a 3-month period | Same staff caring for HBsAg+ and HBsAg-susceptible patients | Separation of HBsAg+ patients from other patients; same staff should not care for HBsAg+ and HBsAg– patients | 264 |
| Eight patients seroconverted to HBsAg+ during a 1-month period | Not consistently using external pressure transducer protectors; same staff members cared for both HBsAg+ patients and susceptible patients | Use external pressure transducer protectors and replace after each use; same staff members should not care for HBV-infected and -susceptible patients on the same shift | 310 |
| 14 patients seroconverted to HBsAg+ during a 6-week period | Failure to review results of admission and monthly HBsAg testing; inconsistent handwashing and use of gloves; adjacent clean and contaminated areas; <20% of patients vaccinated | Proper infection control precautions for dialysis facilities; routine review of serological testing; hepatitis B vaccination of all patients | 265 |
| Seven patients on the same shift seroconverted to HBsAg+ during a 2-month period | Same staff members cared for HBsAg+ and HBsAg– patients on the same shift; common medication and supply carts were moved between stations, and multidose vials were shared | Dedicated staff for HBsAg+ patients; no sharing of equipment or supplies between any patients; centralized medication and supply areas; hepatitis B vaccination of all patients | 265 |
| Four patients seroconverted HBsAg+ during a 3-month period | Transmission appeared to occur during hospitalization at an acute care facility | Hepatitis B vaccination of all patients | 265 |
| 11 patients seroconverted to HBsAg+ during a 3-month period | Staff, equipment, and supplies were shared between HBsAg+ and HBs– patients; no patients were vaccinated | Dedicated staff for HBsAg+; no sharing of medication or supplies between any patients; hepatitis B vaccination of all patients | 265 |
| Two patients converted to HBsAg+ during a 4-month period | Transmission appeared to occur during hospitalization at an acute care facility; Same staff cared for HBsAg+ and HBs– patients; no patients vaccinated | Hepatitis B vaccination of all patients; dedicate staff for the care of HBsAg+ patients; no sharing of supplies or medication between patients | 268 |
| One patient converted to HBsAg+ | Transmission from a patient with history of resolved HBV infection, but the infection reactivated as a result of immunosuppression; multiple infection control breaches observed | Awareness of the reactivation/reserve seroconversion situation | 271 |
| 36 patients with liver enzyme elevations consistent with non-A, non-B hepatitis | Environmental contamination with blood | Use proper precautions (e.g., gloving of staff; environmental cleaning); monthly liver function tests (e.g., ALT) | 311 |
| 35 patients developed elevated liver enzymes consistent with non-A, non-B hepatitis during a 22-month period; 82% of probable cases were anti-HCV | Inconsistent use of infection control precautions, especially hand washing | Strict compliance to aseptic technique and dialysis center precautions | 312 |
| HCV infection developed in 7 out of 40 (17.5%) HCV-susceptible patients; shift specific attack rates of 29%–36% | Multidose vials left on top of machine and used by multiple patients; routine cleaning and disinfection of surfaces and equipment between patients not routinely done; arterial line for draining prime draped into a bucket that was not routinely cleaned or disinfected between patients | Strict compliance with infection control precautions for all dialysis patients; routine HCV testing | 239, 241 |
| HCV infection developed in 5 out of 61 (8%) HCV-susceptible patients | Sharing of equipment and supplies between chronically infected and susceptible patients; gloves not routinely used; clean and contaminated areas not separated | Strict compliance with infection control precautions for all dialysis patients; CDC does not recommend separation of equipment/supplies between HCV-infected and -susceptible patients | 239, 241 |
**TABLE 25.3 Outbreaks of Dialysis-Associated Illnesses Investigated by the Centers for Disease Control and Prevention, 1975–2016—cont’d**

| Description | Cause(s) of Outbreak | Corrective Measure(s) Recommended | Reference |
|-------------|----------------------|-----------------------------------|-----------|
| HCV infection developed in 3 out of 23 (13%) HCV-susceptible patients | Supply carts moved between stations and contained both clean and blood-contaminated items; medications prepared in the same area used for disposal of used injection equipment | Strict compliance with infection control precautions for all dialysis patients | 241 |
| HCV infection developed in 7 out of 52 (13%) HCV-susceptible patients; shift-specific attack rates 4%–21% | Medication cart moved between stations and contained both clean and blood-contaminated items; single-dose medication vials used for multiple patients; cleaning and disinfection of surfaces and equipment between patients not routinely done | Strict compliance with infection control precautions for all dialysis patients | 241 |
| HCV infection developed in 9 out of 119 (7.6%) patients; attack rate 10% | Cleaning and disinfection of surfaces and equipment between patients not routinely done; gloves not routinely used; medications not stored in separate clean area | Strict compliance with infection control precautions for all dialysis patients; perform routine HCV testing | 227 |
| HCV infections developed in 6 out of 66 (9%) patients | Clean and contaminated areas not well delineated; clean supplies accessed with contaminated gloves; medication preparation in proximity to blood specimen processing; reuse of single-dose vials | Strict compliance with infection control precautions for all dialysis patients | CDC unpublished data |
| HCV infections developed in 8 out of 149 (5.4%) patients; attack rate 8.6% | Multidose heparin vials taken to individual dialysis stations; poor hand hygiene and glove use; poor cleaning and disinfection practices | Strict compliance with infection control precautions for all dialysis patients | 244 |
| HCV infections developed in 18 patients; attack rate 16.7% | Poor hand hygiene and glove use; poor cleaning and disinfection practices; blood stains found on machine surfaces after cleaning | Strict compliance with infection control precautions for all dialysis patients | 46 |
| HCV infections developed in 16 patients at 9 facilities between 2013 and 2015 | Multiple infection control breaches identified | Strict compliance with infection control precautions for all dialysis patients | 313 |

ALT, Alanine aminotransferase; HBsAg, hepatitis B surface antigen; HVAC, heating, ventilation, and air conditioning.

Updates on HBV and HCV outbreaks among dialysis patients reported to CDC are available at https://www.cdc.gov/hepatitis/outbreaks/healthcarehepoutbreaktable.htm.

However, the reprocessing of dialyzers has been associated with pyrogenic reactions and bacterial infections. These adverse events may be the result of the use of incorrect concentrations of chemical germicides, the failure to maintain appropriate water quality, breaks in reprocessing procedures, or practical challenges to achieving complete disinfection of reused dialyzers. Manual reprocessing of dialyzers, which is allowed in the United States, does not include testing for membrane integrity, such as a pressure-leak test, may fail to detect membrane defects, and relies on disinfection processes that are particularly difficult to standardize.

Dialyzer reprocessing can be performed in myriad ways with few quality control checks. Procedures used to reprocess hemodialyzers generally constitute high-level disinfection rather than sterilization. Several liquid chemical germicides have been used for high-level disinfection of dialyzers. There are commercially available chemical germicides specifically formulated for this purpose (e.g., peroxyacetic acid, chlorine-based, and glutaraldehyde-based products that are approved by the US Food and Drug Administration [FDA] as sterilants or high-level disinfectants for reprocessing hemodialyzers). During the period between 1983 and 2002, the percentage of centers using formaldehyde for reprocessing dialyzers decreased from 94% to 20%, whereas the percentage using peroxyacetic acid increased from 5% to 72%. Only a minority of facilities (4%) reported used either glutaraldehyde or heat disinfection.

Using a suboptimal disinfectant may lead to outbreaks of infection, such as nontuberculous mycobacteria. An outbreak of systemic mycobacterial infections in five hemodialysis patients, resulting in two deaths, occurred when high-flux dialyzers were contaminated with *Mycobacterium abscessus* during manual reprocessing and disinfected with a commercial disinfectant prepared at a concentration that did not ensure complete inactivation of mycobacteria. These and other outbreaks of infections in dialysis patients emphasize the need to reconsider the safety and necessity of dialyzer reuse.

Outbreaks of pyrogenic reactions (defined as fever or chills in a patient who was afebrile and had no signs or symptoms of an infection before the start of the dialysis treatment session)
have often resulted from reprocessing hemodialyzers with water that did not meet AAMI standards (see Table 25.3). In most instances the water used to rinse dialyzers or to prepare the dialyzer disinfectants exceeded the allowable AAMI microbial or endotoxin standards, because the water distribution system was not disinfected frequently, the disinfectant was improperly prepared, or routine microbial assays were improperly performed. Several outbreaks associated with dialyzer reuse have been reported. Breaches in disinfection of dialyzer components (such as an O-ring) and contamination caused by poor infection control practices during reprocessing steps have been identified as major contributors to those outbreaks. In at least one outbreak, no major breaches in reprocessing were identified. Rather, it was determined that dialyzers are difficult to reprocess safely and completely under typical conditions. This is due to poorly trained staff (often in low-paying jobs), variability in procedures, and few quality control standards.

As described in the most recent investigation of a reuse associated outbreak that resulted in 17 cases, “In practice, reuse and reprocessing of dialyzers poses an increased risk for infection to patients.” In this investigation, each additional use of a dialyzer was associated with higher odds of bloodstream infection. In the era of affordable single-use dialyzers, dialysis providers have discontinued reuse in the interest of patient safety. For facilities or regions where reuse and reprocessing continues to be performed, improved standardization of processes and rigorous quality assurance programs are needed.

High-Flux Dialysis and Bicarbonate Dialysate

High-flux dialysis uses dialyzer membranes and hydraulic permeability that are 5 to 10 times greater than conventional dialyzer membranes. There has been concern that bacteria or more likely endotoxin in the dialysate may penetrate these highly permeable membranes.

High-flux membranes require the use of bicarbonate rather than acetate dialysate. Bicarbonate dialysate must be prepared from two concentrates, an acid concentrate (acetic acid or citric acid) with a pH of 2.8 that is not conducive to microbial growth and a bicarbonate concentrate with a relatively neutral pH and a salt molarity of 1.2 M. Because the bicarbonate concentrate will support rapid growth, its use can increase microbial and endotoxin concentrations in the dialysate and theoretically may contribute to an increase in pyrogenic reactions, especially when used during high-flux dialysis.

Some of the concern appeared justified by results of surveillance data during the 1990s showing a significant association between use of high-flux dialysis and reporting of pyrogenic reactions among patients during dialysis. However, a prospective study of pyrogenic reactions in patients receiving more than 27,000 conventional, high-efficiency, or high-flux dialysis with bicarbonate dialysate containing high concentrations of bacteria and endotoxin found no association between pyrogenic reactions and the type of dialysis treatment. Although there seems to be conflicting data on the relationship between high-flux dialysis and pyrogenic reactions, centers providing high-flux dialysis should ensure that dialysate meets AAMI microbial standards (see Table 25.2).

Disinfection of Hemodialysis Systems

Routine disinfection of isolated components of the dialysis system often produces inadequate results. Consequently, the total dialysis system (water treatment system, distribution system, and dialysis machine) should be included in the disinfection procedure.

Disinfection of dialysis systems usually employs sodium hypochlorite solutions, hydrogen peroxide solutions, commercially available peracetic acid disinfectants, ozone, and, in some systems, hot water pasteurization. Sodium hypochlorite solutions are convenient and effective in most parts of the dialysis system when used at the manufacturer’s recommended concentrations. Also, the test for residual available chlorine to confirm adequate rinsing is simple and sensitive. However, because chlorine is corrosive, it is usually rinsed from the system after a relatively short dwell time of 20 to 30 minutes. The rinse water invariably contains organisms that can multiply to significant levels if the system is permitted to stand overnight. Therefore disinfection with chlorine-based disinfectants are best used before the start of the first patient treatment session rather than at the end of the day. However, for models of machines that most dialysis facilities are using, options for disinfection include heat at the end of the day and use of other disinfectants with longer contact time that also require overnight dwell. There is no need to disinfect the fluid pathway between patients.

Aqueous formaldehyde, peroxyacetic acid, hydrogen peroxide, or glutaraldehyde solutions can produce good disinfection results. These products are not as corrosive as hypochlorite solutions and can be allowed to dwell in the system for long periods of time when the system is not in operation. However, formaldehyde, which has good penetrating power, is considered an environmental hazard and potential carcinogen and has irritating qualities that may be objectionable to staff. The US Environmental Protection Agency has also limited the amount of formaldehyde that can be discharged into the wastewater stream, which has drastically reduced the use of this chemical in the dialysis community as a disinfectant. Peroxyacetic acid and glutaraldehyde are commercially available and are designed for use with dialysis machines when used according to the manufacturers labeled instructions. Glutaraldehyde use is also limited because it is considered a sensitizer and may pose a risk to healthcare workers.

Some dialysis systems (both water treatment and distribution systems, some hemodialysis machines) use hot-water disinfection (pasteurization) for control of microbial contamination. In this type of system water heated to >80°C (176°F) is passed through the water distribution system and hemodialysis machine or just the hemodialysis machine at the end of the day. These systems are excellent for controlling microbial contamination. However, it should be noted that heat disinfection of the hemodialysis machine would not
control microbial contamination of the waste lines and effluent drains. Additional processes may be needed to disinfect waste lines, drains, and wall boxes.

**Monitoring of Water and Dialysis Fluid**

Microbiological and endotoxin standards for water and dialysis fluids (see Table 25.2) were originally based on the results of culture assays performed during outbreak investigations. There is increasing evidence that the microbial quality of hemodialysis fluids plays a role in the chronic inflammatory response syndrome, affects anemia management, accelerates loss of residual renal function, and affects serum albumin levels in dialysis patients. Increasing data suggest that use of ultrapure water and dialysate would benefit maintenance hemodialysis patients and potentially save costs. A large cohort study from Japan found a lower all-cause mortality in facilities using ultrapure water. However, there have been no randomized controlled studies to evaluate and confirm these studies, so regulatory agencies have not yet mandated these higher water standards.

Water samples for routine testing should be collected from a source as close as possible to where water enters the dialysate proportioning unit. In most cases this is the tap (not from that hose connecting the tap to the dialysis machine) at the dialysis station (Fig. 25.1). Water samples should be collected at least monthly from several locations within the dialysis unit, including samples at different dialysis stations. Samples should also be collected using a similar approach after any modifications or maintenance have been made to the water treatment system water distribution system. Dialysate samples should be collected during or at the end of the dialysis treatment from a source close to where the dialysis fluid either enters or leaves the dialyzer (Fig. 25.2). Dialysate samples should be collected at least monthly from a representative number of dialysis machines. Samples of water and dialysate should also be collected when a pyrogenic reaction is suspected. If centers reprocess hemodialyzers for reuse, water used to prepare disinfectant and rinse dialyzers should also be assayed monthly.

The maximum contaminant levels for water are 100 CFU/mL and 0.25 EU/mL. Methods for microbiological and endotoxin testing are available elsewhere.

In an outbreak investigation, the assay methods may need to be both qualitative and quantitative; also detection of non-tuberculous mycobacteria and in some cases fungi in water or dialysate may be desirable. In such instances, plates should be incubated for 5 to 14 days at both 36°C and 28°C to 30°C. Laboratories should be notified of special testing requests outside of routine water testing, such as if the facilities would like to look for specific pathogens.

**DIALYSIS-ASSOCIATED PYROGENIC REACTIONS**

Gram-negative bacterial contamination of dialysis water or components of the dialysis system (water, dialysate, water used for reprocessing) can cause pyrogenic reactions. A pyrogenic reaction is defined as objective chills (visible rigors) or fever (oral temperature ≥ 37.8°C [100°F]) or both in a patient who was afebrile (oral temperature up to 37°C [98.6°F]) and
had no signs or symptoms of an infection before the start of the dialysis treatment session.\textsuperscript{87,88} Depending on the type of dialysis system and the level of contamination, fever and chills may start 1 to 5 hours after dialysis has been initiated. Other symptoms may include hypotension, headache, myalgia, nausea, and vomiting. Pyrogenic reactions can occur without bacteria; because presenting signs and symptoms cannot differentiate bacteremia from pyrogenic reactions, blood cultures are necessary.

During 1990–2002 an annual average of 20% to 24% of the hemodialysis centers in the United States reported at least one pyrogenic reaction in the absence of septicemia in their patients undergoing maintenance hemodialysis.\textsuperscript{48,49,89–97} Pyrogenic reactions can result from passage of bacterial endotoxin (lipopolysaccharide) or other substances in the dialysate across the dialyzer membrane\textsuperscript{98–102} or from the transmembrane stimulation of cytokine production in the patient’s blood by endotoxin in the dialysate.\textsuperscript{99,103–105} In other instances, endotoxin can enter directly into the bloodstream with fluids that are contaminated with gram-negative bacteria.\textsuperscript{106} The signs and symptoms of pyrogenic reactions without bacteremia generally abate within a few hours after the dialysis has been stopped. If gram-negative sepsis is associated, fever and chills may persist and hypotension is more refractory to therapy.\textsuperscript{4,106}

When a patient develops a pyrogenic reaction (i.e., onset of fever or chills) while being dialyzed, the following steps are recommended: (1) careful physical examination of the patient to identify signs and symptoms and evaluate other possible causes of chills and fever (e.g., pneumonia, vascular access site infection); (2) blood cultures, other diagnostic tests (e.g., chest radiograph), and other cultures as clinically indicated; (3) collection of dialysate from the dialyzer (i.e., postdialyzer effluent sample) for quantitative and qualitative microbiological culture; and (4) recording of the incident in a log or other permanent record. In addition, empiric antibiotic treatment should be administered to the patient. Determining the cause of such episodes is important because they may be the first indication of a remediable problem that can affect a potentially large number of patients.

The higher the level of bacteria and endotoxin in dialysis fluid, the higher the probability that the bacteria or their products will pass through the dialyzer membrane to produce bacteremia or stimulate cytokine production. In an outbreak of febrile reactions among patients undergoing hemodialysis, the attack rates were directly proportional to the level of microbial contamination in the dialysis fluid.\textsuperscript{6} Prospective studies also reported a lower pyrogenic reaction rate among patients when they underwent dialysis with dialysis fluid from which most bacteria had been removed by filtration, compared with patients who underwent dialysis fluid that was highly contaminated (mean 19,000 CFU/mL).\textsuperscript{5,87,107}

**DISINFECTION, STERILIZATION, AND ENVIRONMENTAL CLEANING IN DIALYSIS FACILITIES**

Good cleaning, disinfection, and sterilization procedures are important components of the infection control program in the hemodialysis center. The procedures do not differ from those recommended for other healthcare settings,\textsuperscript{108,109} but the
high potential for blood contamination makes the hemodialysis setting unique. In addition, the need for routine aseptic access of the patient's vascular system makes the hemodialysis unit more akin to a surgical suite than to a standard hospital room. Medical items are categorized as critical (e.g., needles and catheters), which are introduced directly into the bloodstream or normally sterile areas of the body; semicritical (e.g., fiberoptic endoscopes), which come in contact with intact mucous membranes; and noncritical (e.g., blood pressure cuffs), which touch only intact skin.\textsuperscript{109,110}

Cleaning and housekeeping in the dialysis center have two goals: to remove soil and waste on a regular basis, thereby preventing the accumulation of potentially infectious material, and to maintain an environment that is conducive to good patient care.\textsuperscript{110} Crowding of patients and patient stations, as well as overtaxing of staff members, may increase the likelihood of microbial transmission. Adequate cleaning may be difficult if there are multiple wires, tubes, and hoses in a small area. There should be enough space to move completely around each patient's dialysis station without interfering with the neighboring stations. According to the Facility Guidelines Institute, each dialysis station should be at least 80 square feet and allow at least 4 feet distance between stations to avoid contamination.\textsuperscript{111} However, most of dialysis facilities do not have space to meet that guideline. To avoid contamination, cleaning should only start when patients have left their stations and staff should not allow new patients into chairs until cleaning and disinfection is complete. Creating unit-wide patient-free intervals between treatment shifts is likely to improve the adequacy of station cleaning and disinfection between patients.

Where space is limited, elimination of unneeded items, orderly arrangement of required items and removal of excess lengths of tubing, hoses, and wires from the floor can improve accessibility for cleaning. Because of the special requirements for cleaning in the dialysis center, staff should be specially trained in this task.

After each patient treatment, frequently touched environmental surfaces, including external surfaces of the dialysis machine, should be properly disinfected; some surfaces may also require precleaning (with a detergent) before disinfection. A study in the Netherlands and an investigation of a large HCV outbreak in the United States where the investigators used chemiluminescent agents to detect nonvisible blood contamination have demonstrated the importance of environmental cleaning.\textsuperscript{46,112} Antiseptics, such as formulations with povidone-iodine, hexachlorophene, or chlorhexidine, should not be used for surface disinfection because these are formulated for use on skin and are not designed for use on hard surfaces. Given the role of environmental surfaces of components adjacent to the machine (e.g., wall boxes) in transmission of pathogens, as illustrated in recent outbreaks,\textsuperscript{47} attention should be paid to cleaning and disinfection of those surfaces as well.

**BLOODSTREAM INFECTIONS AND OTHER INFECTIONS**

The annual adjusted mortality rate among hemodialysis patients is 169 per thousand patient-years at risk. Infection is the second leading cause of mortality in this patient population, accounting for 8% of all deaths.\textsuperscript{1} In a number of published studies that have evaluated bacterial infections in outpatient hemodialysis, bacteremia occurred in 0.6% to 1.7% of patients per month and vascular access infections (VAIs; with or without bacteremia) in 1.3% to 7.2% of patients per month.\textsuperscript{113-123} A review of four studies published during 2002 estimated that 1.8% of hemodialysis patients have vascular access associated bacteremia each month, amounting to 50,000 episodes nationally per year.\textsuperscript{124} In a study of 27 French hemodialysis centers, 28% of 230 infections in hemodialysis patients involved the vascular access, whereas 25% involved the lung, 23% the urinary tract, 9% the skin and soft tissues, and 15% other or unknown sites.\textsuperscript{119}

Because of the importance of bacterial infections in hemodialysis patients, the CDC initiated a voluntary ongoing surveillance system in the United States called the Dialysis Surveillance Network (DSN) in 1999.\textsuperscript{122} At the time, only bacterial infections associated with hospital admission or intravenous antimicrobial receipt were counted; as a result, this system likely only detected more severe infections. During 1999–2001, 109 dialysis centers reported data. Rates per 100 patient-months were 3.2 for all VAIs (including access infections both with and without bacteremia), 1.8 for vascular access associated bacteremia, 1.3 for wound infection not related to the vascular access, 0.8 for pneumonia, and 0.3 for urinary tract infection. Among patients with fistulas or grafts, wounds were the most common site for infection. Among patients with hemodialysis catheters, infections of the vascular access site were the most common site for infection.\textsuperscript{122} The surveillance project expanded and evolved into National Healthcare Safety Network (NHSN), of which Dialysis Event Surveillance is a component (https://www.cdc.gov/nhsn/dialysis/index.html). The NHSN is an Internet-based surveillance system that enables facilities to report healthcare-associated infection data to the CDC. VAIs in dialysis patients and related events are reported to NHSN's Dialysis Event Surveillance. Outpatient hemodialysis facilities in the United States eligible to participate in the surveillance are instructed to follow a standard protocol,\textsuperscript{125} by which all outpatients who receive hemodialysis at the facility are monitored for three NHSN-defined dialysis events. The three types of dialysis events (positive blood culture; intravenous antimicrobial start; and pus, redness, or increased swelling at the vascular access site) are reported using a standard data collection form. During 2007–2011, 193 facilities reported to NHSN; the rate of bloodstream infection (BSI) and access-related BSI was 1.27 and 0.88 per 100 patient-months, respectively.\textsuperscript{126} Data reported to NHSN have been used by CMS as part of the ESRD Quality Incentive Program since 2012; as a result, almost all outpatient hemodialysis facilities now report to NHSN. In 2014 more than 6000 facilities now reported 160,971 events, including 29,516 BSIs and 22,576 access-related BSIs; the rate of BSI was 0.64 per 100 patient-months. The rate of BSI was much higher among patients with a central venous catheter (2.16 per 100 patient-months) compared with other vascular access types.\textsuperscript{127}
Vascular Access Infections

Access site infections are particularly important because they can cause disseminated bacteremia or loss of the vascular access. Local signs of VAI include erythema, warmth, induration, swelling, tenderness, breakdown of skin, loculated fluid, or purulent exudates. Based on data from DSN collected during 1995–2005, the overall VAI rate was 3.1 per 100 patient-months and varied from 0.6 for fistulas to 10.1 for temporary catheters. In the 2014 NHSN surveillance data report, the VAI rate was 1.21 per 100 patient-months. The access-related BSI rate was 0.49 per 100 patient-months, which varied by access type: 0.16 for fistulas, 0.27 for grafts, and 1.83 for central venous catheters (tunneled and nontunneled).

VAIs are caused (in descending order of frequency) by Staphylococcus aureus (32% to 53% of cases), coagulate-negative staphylococci (20% to 32% of cases), gram-negative bacilli (10% to 18%), other gram-positive cocci (including enterococci; 10% to 12%), and fungi (<1%). Among BSIs, S. aureus remained the most commonly reported pathogen in 2014 NHSN data (31% of BSI and 32% of access-related BSI), and 40% of cases of S. aureus were resistant to methicillin.

The primary risk factor for vascular access–related infection is access type, with catheters having highest risk for infection; grafts intermediate; and native arteriovenous (AV) fistulas the lowest. Other potential risk factors for VAI include (1) location of the access in the lower extremity; (2) recent vascular access surgery; (3) trauma, hematoma, dermatis, or scratching over the access site; (4) poor patient hygiene; (5) poor needle insertion technique; (6) older age; (7) diabetes; (8) immunosuppression; (9) iron overload; (10) intravenous drug use; and (11) chronic inflammatory state.

Based on the relative risk for both infectious and noninfectious complications, native AV fistulas are considered the preferred vascular access type; a goal of no more than 10% of patients maintained with permanent catheter–based hemodialysis treatment is recommended. To minimize infectious complications, patients should be referred early for creation of an arteriovenous access, thereby decreasing the time dialedyzed through a temporary catheter. During the period between 1995 and 2002, the percentage of patients dialedyzed through fistulas increased from 22% to 33%, with most of the increase occurring after 1999. Data from Dialysis Outcomes and Practice Patterns Study indicated that from August 2010 to August 2013, AV fistula use increased from 63% to 68%, whereas catheter use declined from 19% to 15%. However, the majority of incident patients still initiated dialysis with a catheter. The US Renal Data System (USRDS) annual data report for 2016 indicated that whereas 18.8% of prevalent hemodialysis patients used a catheter, 80.3% of incident patients started dialysis with a catheter.

Etiology and Prevention of Bloodstream Infection

Bacterial pathogens causing infection can either be exogenous (i.e., acquired from contaminated dialysis fluids or equipment) or endogenous (i.e., caused by invasion of bacteria present in or on the patient). Catheter-related infections are most often caused by bacteria from the patient’s skin colonizing the outside of the catheter or from direct contact (e.g., touch contamination by healthcare personnel) with the catheter hub, leading to contamination of the inner surface of the catheter. Surveillance data indicate that S. aureus and other coagulate-negative staphylococci were the most common pathogens for BSI and access-related BSI. Endogenous sources may also be more likely causes of VAI among fistula and graft patients. Contaminated infusates and hematogenous spread are thought to be less common causes of BSI in this patient population, regardless of vascular access type.

Exogenous pathogens have caused numerous outbreaks, most of which resulted from inadequate dialyzer reprocessing procedures (e.g., contaminated water or inadequate disinfectant concentration) or inadequate disinfection and maintenance of the water treatment and distribution system. During 1995–2006, five outbreaks were traced to contamination of the waste handling option on one type of dialysis machine. Recommendations to prevent such outbreaks have been published elsewhere. Contaminated medication vials are also a source of bacterial infection for patients. In 1999, an outbreak of Serratia liquefaciens bloodstream infections and pyrogenic reactions among hemodialysis patients was traced to contamination of vials of erythropoietin. These vials, which were intended for single use, were contaminated by repeated puncture to obtain additional doses and by pooling of residual medication into a common vial.

Recommendations for preventing VAIs have been developed by the CDC and the Healthcare Infection Control Practices Advisory Committee and the National Kidney Foundation. The CDC has developed a recommended “Approach to BSI Prevention in Dialysis Facilities” that includes core interventions to prevent BSI among hemodialysis patients (Table 25.4). Facilities that implemented this set of interventions were able to reduce their access-related BSI rates and sustained these lowered rates for at least 4 years. The core interventions include (1) BSI surveillance using NHSN and feedback to clinical staff; (2) hand hygiene observations with feedback to staff; (3) catheter vascular access care observations to ensure clinical staff adherence to aseptic technique and good infection control practices (with staff feedback); (4) development of staff infection prevention skills, demonstrated through competency assessments; (5) patient education and engagement in infection control processes; (6) decrease catheter prevalence; (7) catheter hub disinfection; and (8) bacitracin zinc/polymyxin B sulfate (Polysporin) triple ointment or povidone-iodine ointment applied to catheter exit sites. The CDC has also developed tools, protocol, and guidance to assist in the implementation of the interventions (https://www.cdc.gov/dialysis/prevention-tools/index.html).

Other strategies that might assist in implementation of recommended interventions include staff engagement and safety culture. Use of a behavioral change strategy (“Positive Deviance”), in which positive BSI prevention practices by certain staff were encouraged among all staff, was found to contribute to the reduction of BSI in one dialysis facility.
Incorporate efforts (e.g., through patient education modalities) in conjunction with other preventive strategies such as (1) antimicrobial catheter lock solutions in patients with multiple BSIs despite optimal adherence to aseptic technique; (149) Routine prophylactic use of antimicrobial lock solutions for hemodialysis catheter-related BSI is not recommended at this time. (148,155)

In hemodialysis patients, the Infectious Diseases Society of America has recommended treatment with nasal mupirocin in documented S. aureus carriers who have catheter-related BSI with S. aureus and continue to need a hemodialysis catheter. (156,157) Otherwise, the routine use of nasal mupirocin in patients with hemodialysis catheters is not recommended by either the CDC or the National Kidney Foundation. (137,138,148) The CDC also updated the Guidelines for the Prevention of Intravascular Catheter-Related Infections and included the recommendation of using chlorhexidine-impregnated dressings to protect the insertion site of short-term, nontunneled central venous catheters in patients aged 18 years and older (https://www.cdc.gov/infectioncontrol/guidelines/bsi/c-i-dressings/recommendations.html). However, no recommendations were made for patients with long-term, tunneled catheters, and the effect of chlorhexidine-impregnated dressings on reducing catheter-related bloodstream infections among hemodialysis patients remains unclear. (158,159)

A recently developed chlorhexidine-impregnated catheter cap (ClearGuard®) has been reported in a cluster randomized trial to reduce catheter-related BSIs and hospital admissions for BSI. (160) A needle-free connector (TEGO® needle-free hemodialysis connector [ICU Medical, Inc., San Clemente, Calif.]) was found to be significantly associated with less intravenous antibiotic use among hemodialysis patients; however, the risk for catheter-related BSI among patients who used the connector was not statistically significantly decreased. (161)

Poor injection safety practices have led to BSI among dialysis patients, and thus improving injection practices should be considered as a strategy to reduce the spread of both bloodborne viruses (e.g., hepatitis B and C) and BSIs. (147) To reduce the risk for infection, the CDC recommends (a) preparing medications in a clean room or, if a clean room is not available, in an area separated from the patient treatment area and designated for medications; (b) performing hand hygiene and using aseptic technique when preparing medication; (c) disinfecting the rubber septum of vials with alcohol and using a new needle and a new syringe to withdraw medication; (d) discarding single-dose vials and storing multidose vials appropriately; (e) not handling or storing used supplies, equipment, blood samples, or biohazard containers in or adjacent to areas where medications and clean (i.e., unused) equipment and supplies are handled; (f) delivering medications separately to each patient and not using common carts within the patient treatment area to prepare or distribute medications; and (g) performing hand hygiene, putting on new, clean gloves, scrubbing the injection port with antiseptic, and using aseptic technique when administering medications. (162) Intravenous medication vials labeled for single use, including erythropoietin, should not be punctured more than once. Multidose medication vials should be assigned to a single patient whenever possible. (163)

### TABLE 25.4 Core Interventions for Dialysis Bloodstream Infection Prevention

| Interventions                               | Description                                                                 |
|---------------------------------------------|-----------------------------------------------------------------------------|
| Surveillance and feedback using NHSN       | Conduct monthly surveillance for BSIs and other dialysis events using NHSN, and actively share results with frontline clinical staff. |
| Hand hygiene observations                  | Perform observations of hand hygiene opportunities monthly and share results with clinical staff. |
| Catheter/vascular access care observations | Perform observations of vascular access care and catheter accessing quarterly. Assess staff adherence to aseptic technique when connecting and disconnecting catheters and during dressing changes. Share results with clinical staff. |
| Staff education and competency            | Train staff on infection control topics, including access care and aseptic technique. Perform competency evaluation for skills such as catheter care and accessing every 6–12 months and on hire. |
| Patient education/engagement              | Provide standardized education to all patients on infection prevention topics. |
| Catheter reduction                         | Incorporate efforts (e.g., through patient education, vascular access coordinator) to reduce catheters by identifying and addressing barriers to permanent vascular access placement and catheter removal. |
| Chlorhexidine for skin antisepsis          | Use an alcohol-based chlorhexidine (>0.5%) solution as the first-line skin antiseptic agent for central line insertion and during dressing changes. |
| Catheter hub disinfection                  | Scrub catheter hubs with an appropriate antiseptic after cap is removed and before accessing. |
| Antimicrobial ointment                     | Apply triple antibiotic ointment or povidone-iodine ointment to catheter exit sites during dressing change. |

Additional recommendations for preventing hemodialysis-catheter–associated infections include (1) using sterile technique and maximal sterile barrier precautions (cap, mask, sterile gown, large sterile drapes, and gloves) during catheter insertion; (2) limiting use of noncuffed catheters to 3 to 4 weeks; (3) restricting catheter manipulation and dressing changes to trained personnel; (4) replacing catheter site dressing if damp, loosened, or soiled. (148,152)

A number of studies have looked at the use of various antimicrobial locks to prevent catheter-related BSI among hemodialysis patients. Two meta-analyses of these studies concluded that (1) antimicrobial catheter lock solutions reduce catheter-related bloodstream infections and the (2) use of these lock solutions should be considered in routine clinical practice in conjunction with other prevention modalities. (153,154) However, the long-term consequence of using antibiotics routinely in catheter locking solutions is unknown. CDC and the Healthcare Infection Control Practices Advisory Committee guidelines recommend lock solutions in patients with multiple BSIs despite optimal adherence to aseptic technique. (149) Routine prophylactic use of antimicrobial lock solutions for hemodialysis catheter-related BSI is not recommended at this time. (148,155)

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Poor injection safety practices have led to BSI among dialysis patients, and thus improving injection practices should be considered as a strategy to reduce the spread of both bloodborne viruses (e.g., hepatitis B and C) and BSIs. (147) To reduce the risk for infection, the CDC recommends (a) preparing medications in a clean room or, if a clean room is not available, in an area separated from the patient treatment area and designated for medications; (b) performing hand hygiene and using aseptic technique when preparing medication; (c) disinfecting the rubber septum of vials with alcohol and using a new needle and a new syringe to withdraw medication; (d) discarding single-dose vials and storing multidose vials appropriately; (e) not handling or storing used supplies, equipment, blood samples, or biohazard containers in or adjacent to areas where medications and clean (i.e., unused) equipment and supplies are handled; (f) delivering medications separately to each patient and not using common carts within the patient treatment area to prepare or distribute medications; and (g) performing hand hygiene, putting on new, clean gloves, scrubbing the injection port with antiseptic, and using aseptic technique when administering medications. (162) Intravenous medication vials labeled for single use, including erythropoietin, should not be punctured more than once. Multidose medication vials should be assigned to a single patient whenever possible. (163)
Respiratory Infections

Hospital admissions for pneumonia have been declining overall for dialysis patients; however, pneumonia rates for hemodialysis patients are 1.8 to 2.0 times that of renal transplant recipients or peritoneal dialysis patients. Hospital admissions for pneumonia are also 102% higher among hemodialysis patients compared with the general population. In one study of a group of 433 dialysis patients over a 9-year period, pneumonia was the third most common cause of infection (after vascular access and infections below the knee) and accounted for 13% of all infections. One- and five-year survival probabilities are 0.55 and 0.17, respectively. Pneumonia is common among hemodialysis patients, carries a poor prognosis, and is often the antecedent to cardiovascular death. A recent analysis of incident hemodialysis patients found pneumonia to be associated with chronic obstructive pulmonary disease, inability to transfer or ambulate, hemodialysis as initial therapy, advanced age (≥75 years), and body mass index ≥ 30 kg/m². According to the Advisory Committee on Immunization Practices, patients with chronic renal failure should be vaccinated with the pneumococcal polysaccharide vaccine. Both 13-valent pneumococcal conjugate vaccine and 23-valent pneumococcal polysaccharide vaccine can be used; the schedule depends on the vaccination status of patients and is available on the CDC website (https://www.cdc.gov/dialysis/pdfs/vaccinating_dialysis_patients_and_patients_dec2012.pdf).

Patients with chronic kidney diseases, including hemodialysis patients, are at high risk for developing complications of seasonal influenza, which can be effectively prevented by vaccination. Influenza vaccination is associated with lower risk for hospitalization and death in ESRD patients. Although transmission of influenza, even during pandemics, in US dialysis clinics is not evident through the published literature, anecdotal reports of transmission exist. Because of their higher risk for complications, it is important to maintain influenza vaccination coverage among hemodialysis patients. In addition to influenza vaccination for patients, vaccination for dialysis providers (clinicians, nurses, technicians) is also important. Vaccination for healthcare personnel has been found to decrease absenteeism and healthcare facility acquired influenza. However, coverage among dialysis healthcare personnel was still suboptimal: data indicated that only 73% of healthcare personnel in dialysis clinics received influenza vaccination in the 2011–2012 season. Annual vaccination is therefore recommended for both dialysis patients and healthcare personnel. ESRD patients with latent tuberculosis (TB) infection are at higher risk for developing TB disease. For this reason, the CDC recommends that all dialysis patients be tested at least once on admission for latent TB infection or TB disease using a tuberculin skin test or TB blood test. Patients who test positive should be evaluated for treatment and for the presence of TB disease. TB transmission in US dialysis facilities has been very rare. The most recently reported instance occurred in 2003 when an infected healthcare worker transmitted the bacteria to patients and other healthcare workers at a dialysis facility. Another episode in 1998 involved a patient with smear-positive pulmonary TB, but no transmission to other patients at the dialysis facility was identified. Suspected or confirmed TB exposure occurring within a dialysis facility should be reported to the appropriate state or local public health authority.

A new emerging respiratory infection, Middle East respiratory syndrome (MERS) caused by a corona virus, was first reported in 2012. No MERS infections or transmissions have been reported in the United States, but because of their significance and transmissibility, healthcare facilities, including dialysis clinics, should remain vigilant for MERS and other respiratory pathogens.

To prevent the transmission of respiratory infections (e.g., influenza) in dialysis facilities, staff should have systems in place to detect patients with respiratory symptoms on presentation to the facilities and implement interventions to decrease transmission. Facilities should educate patients about respiratory hygiene and provide necessary supplies such as tissues, mask, and hand hygiene materials. Patients should be encouraged to notify facility staff of any respiratory symptoms when they arrive. Facilities should also have policies that encourage dialysis healthcare personnel to not work while sick with respiratory infection. For new and emerging pathogens, dialysis providers should maintain awareness of current issues and recommendations from state and local public health departments and the CDC. Any possible instances of transmission of one of these new and emerging pathogens in a dialysis facility should be reported to public health authorities.

Antimicrobial-Resistant Bacteria

Hemodialysis patients have been in the forefront of the epidemic of antimicrobial resistance, especially vancomycin resistance. One of the earliest reports of vancomycin-resistant enterococci (VRE) was from a renal unit in London in 1988. The prevalence of VRE stool colonization among dialysis patients has varied from 1.5% among pediatric dialysis patients in the United Kingdom and 2.4% of adult dialysis patients at three dialysis centers in Indianapolis, Indiana, to 9.5% at a university hospital in Baltimore, Maryland. In one center the prevalence of rectal carriage of VRE was 9%, and 2% of noncarriers developed VRE infections in 1 year. A meta-analysis of studies from 100 facilities and 4800 patients worldwide reported a pooled VRE colonization prevalence of 6.2%. It appears that hospital acquisition of VRE contributes substantially to the increasing prevalence of VRE in the maintenance hemodialysis patient population. Among enterococci causing bloodstream infections in hemodialysis patients, up to 11.4% have been reported to be resistant to vancomycin. Vancomycin resistance in S. aureus has also been reported in dialysis patients. Five of the first six US patients with infections associated with vancomycin–intermediate S. aureus were receiving either peritoneal dialysis or hemodialysis. In addition, the first US patient found to be infected with a vancomycin-resistant S. aureus (VRSA) strain was a maintenance hemodialysis patient; the VRSA was isolated from...
a diabetic foot wound and from a temporary central venous catheter exit site. In the period between 2002 and 2009 there were nine cases of VRSA in the United States; three of these patients had chronic renal failure and two were hemodialysis patients. Five of those VRSA cases occurred in southeastern Michigan and contained a plasmid carrying the vanA gene, which had been donated from a VRE donor. To date, 14 cases of VRSA have been reported to the CDC, of which the most recent case was in a dialysis patient. A guide to investigation and control of VRSA is available from the CDC and includes suggested strategies for VRSA control in dialysis centers.

The percentage of hemodialysis facilities reporting methicillin-resistant *S. aureus* (MRSA) infection or colonization has increased from 40% in 1995 to 76% in 2002. In a 2005 CDC study assessing the incidence of invasive MRSA infection among dialysis patients, the incidence of invasive MRSA infection was found to be 42.5 cases per 1000 dialysis patient population. This is approximately 100-fold higher than the general population, in which rates for invasive MRSA infection are 0.2 to 0.4 cases per 1000 population. The rate of invasive MRSA infections among hemodialysis patients appears to be decreasing. In 2015, invasive MRSA incidence had decreased to 14.8 cases per 1000 population, still much higher than the incidence among general population. A study in the United Kingdom found that MRSA was responsible for 30% of *S. aureus* catheter-related infections in hemodialysis patients. In the United States, 30.6% of BSIs in hemodialysis patients were caused by *S. aureus* and 39.5% of the *S. aureus* BSI isolates were methicillin-resistant strains.

Patients with chronic kidney disease, including end-stage renal disease, are at high risk for *Clostridium difficile* infection (CDI). Limited data are available on CDI among dialysis patients in the United States. In a review of USRDS (Medicare claims) data between 2005 and 2008, 4.25% of dialysis patients were diagnosed with first episode of CDI. In a cohort of dialysis patients followed from 1999 to 2007, 14.3% of hemodialysis patients developed CDI (a rate of 8.3 cases per 100 patient-years). The UK Renal Registry reported an incidence of 1.09 CDI per 100 patient-years among hemodialysis patients in 2013–2014. An outbreak of CDI in a hemodialysis facility has been reported. The outbreak investigation revealed several challenges in prevention and control of CDI among dialysis patients (e.g., shared patient environment and equipment, lack of physical barriers between patient treatment stations, and adequacy of typical cleaning and disinfection procedures). CDI control strategies that were employed during this outbreak included designation of select dialysis stations as CDI contact isolation stations, use of dedicated, disposable gown and gloves by staff while caring for a patient in contact isolation, handwashing with soap and water after caring for CDI patients, use of 1:10 dilution of bleach to disinfect environmental surfaces in stations after treatment of CDI patients, and heightened diligence to ensure adequate wet contact time of bleach on surfaces.

To combat emerging antimicrobial resistance in dialysis patients, one must understand the transmission kinetics involved with each organism. For certain patients, including those infected with MRSA or VRE, contact precautions are used in the hospital setting. The CDC has not recommended routine use of contact precautions in hemodialysis centers for patients infected or colonized with multidrug-resistant organisms (MDROs). Transmission of pathogenic bacteria is well documented in hospitals. At least one study has suggested that the majority of transmission and acquisition of resistant pathogens among dialysis patients occurs when these patients are admitted to the acute care setting. However, studies have demonstrated MDRO spread in dialysis centers. The CDC recommends additional precautions be used during treatment of patients who might be at higher risk for transmitting pathogenic bacteria (i.e., those with an infected skin wound with drainage that is not contained by dressings or fecal incontinence or uncontrolled diarrhea). These interventions include the following: (1) Staff members treating the patient should wear a separate gown over their usual clothing and remove the gown when finished caring for the patient; (2) patients should be dialyzed at a station away from the main flow of traffic and with as few adjacent stations as possible (e.g., at the end or corner of the unit). However, preventing transmission of resistant pathogens depends primarily on adherence to basic infection control practices and these additional practices. More work is needed to understand the transmission of targeted MDROs in dialysis settings and the effectiveness of interventions to reduce transmission.

One major contributor to the development of antimicrobial-resistant bacteria is inappropriate use of antimicrobial drugs. Antibiotics are commonly used in dialysis patients, especially vancomycin, cefazolin, and third- and fourth-generation cephalosporins. In a small study, as many as 30% of antibiotic indications were found to be inappropriate. Reasons for those inappropriate uses and possible strategies for improved antibiotic stewardship in dialysis facilities have been proposed. More data are needed to understand the relationship between antibiotic prescribing patterns in dialysis centers and antibiotic resistance to better target potential stewardship activities.

### HEPATITIS C VIRUS

HCV is a single-stranded RNA virus that belongs to the family *Flaviviridae*. HCV was first recognized as non-A, non-B hepatitis virus in 1974 until cloning of the etiological agent in 1989. HCV is a relatively efficiently transmitted bloodborne viral pathogen in the dialysis setting. It is not as efficiently transmitted as HBV in this setting, and generally, recommended infection control practices do prevent transmission among hemodialysis patients (without need for isolation). However, new acquisition of hepatitis C infection continues to occur among maintenance hemodialysis patients and outbreaks of hepatitis C are far more common than outbreaks of hepatitis B in the dialysis setting.
Epidemiology

In 2002, 63% of dialysis centers tested patients for antibodies against HCV (anti-HCV). In the facilities that performed screening, the incidence rate in 2002 was 0.34%, and among these centers, the prevalence of anti-HCV among patients was 7.8%, a decrease of 25.7% since 1995. Only 11.5% of dialysis facilities reported newly acquired HCV infection among their patients. Higher incidence rates have been reported from cohort studies of dialysis patients in the United States (<1% to 3%), Japan (<2%), and Europe (3% to 15%). Higher prevalence rates (10% to >85%) also have been reported in individual facilities and in other countries.

HCV is moderately stable in the environment and can survive drying and environmental exposure to room temperature for at least 16 hours. Longer survival, up to several weeks, has been reported. HCV is most efficiently transmitted by direct percutaneous exposure to blood, and like HBV, the chronically infected patient is central to the epidemiology of HCV transmission. Risk factors associated with HCV infection among hemodialysis patients include blood transfusions from unscreened donors, injection drug use, low staff-to-patient ratios, dialysis in a facility with high HCV prevalence, and number of years on dialysis. The number of years on dialysis is a risk factor that is independently associated with higher rates of HCV infection. Multiple studies found that as the time patients spent on dialysis increased, their prevalence of HCV infection increased.

These studies, as well as investigations of dialysis-associated outbreaks of hepatitis C infection, indicate that HCV transmission most likely occurs because of inadequate infection control practices. The practices that have been found to be associated with higher prevalence of HCV in dialysis facilities include handling blood specimens near medication preparation area or other clean areas, use of a mobile cart to distribute medications, poor disinfection of priming buckets, and inconsistent cleaning of dialysis machines. The CDC tracks HCV outbreaks in dialysis settings (https://www.cdc.gov/hepatitis/outbreaks/healthcarehepoutbreaktable.htm); during 1998–2008, the CDC helped investigate five outbreaks of HCV infection among patients in hemodialysis centers. From 2008 to 2015, 18 outbreaks involving at least 98 newly infected patients were reported to the CDC. In those outbreaks a common finding was that seroconversions were associated with receiving dialysis immediately after or at a machine adjacent to a chronically infected patient. Multiple opportunities for cross-contamination were observed in the involved facilities, including (a) equipment and supplies that were not disinfected between patient use; (b) use of common medication carts to prepare and distribute medications at patient stations; (c) sharing of multiple dose vials, which were used at patients’ stations; (d) contaminated priming buckets that were not routinely changed or cleaned and disinfected between patients; (e) machine surfaces that were not routinely cleaned and disinfected between patients; and (f) blood spills that were not cleaned up promptly. Investigation of an outbreak involving four different clusters found multiple lapses in infection control and blood contamination of environmental surfaces as a result of poor cleaning and disinfection practice. In these outbreaks, a single common exposure event is rarely identified, and many outbreaks involve separate chains of transmission occurring over time. Moreover, it has been noted that station turnover procedures are rushed and disinfection of machine surfaces is initiated before the patient has left the treatment station. These common practices are challenges to proper cleaning and disinfection and prevention of cross-transmission of bloodborne pathogens such as HCV.

Other traditional risk factors for acquiring HCV include injection drug use, exposure to an HCV-infected sexual partner or household contact, multiple sexual partners, and perinatal exposure. The efficiency of transmission in settings involving sexual or household exposure to infected contacts is low, and the magnitude of risk and the circumstances under which these exposures result in transmission are not well defined. When a new HCV infection (includes acute, symptomatic infection or HCV seroconversion) occurs in a dialysis facility, it should be assumed that the infection was healthcare related and investigated as such. State and local health departments to whom these infections should be reported have extensive expertise in evaluating traditional risk factors that the patient might have in addition to healthcare exposures.

Treatment for HCV infection has gained significant achievements in the past several years, and recent data have indicated that ESRD patients infected with HCV can be treated successfully. All dialysis patients with HCV infection should be referred to care and assessment. Because dialysis in a facility with high HCV prevalence is a risk factor for HCV infection, HCV treatment may reduce the number of infected patients and therefore help decrease the number of new infections. However, the effect of HCV treatment on transmission of HCV in dialysis facilities is unknown.

Screening and Diagnostic Tests

FDA-licensed or approved tests to screen for HCV antibodies (anti-HCV) in the United States comprise immunoassays, immunoblot assays, and immunochromatography-based rapid tests. None discriminate between active and resolved HCV infection, and confirmatory recombinant immunoblot tests have been discontinued. All individuals who test anti-HCV positive should be further tested for HCV RNA by an FDA-approved nucleic acid test to determine current infection status.

Routine testing of hemodialysis patients for anti-HCV on admission and every 6 months has been recommended since 2001. For routine HCV screening of hemodialysis patients, the anti-HCV screening immunoassay (either rapid test or laboratory-based assay) is recommended, and if positive, this should be confirmed with HCV RNA testing (Box 25.1).

Prevention of Hepatitis C Virus Transmission

Lessons from investigations of HCV outbreaks in dialysis indicate that breaches in infection control practices are the
of care and is recommended to all susceptible hemodialysis patients. The vaccine series should ideally be administered before starting dialysis for ESRD.\textsuperscript{162}

## Epidemiology

During the early 1970s, HBV infection was endemic in maintenance hemodialysis units and outbreaks were common. Subsequently, the incidence and prevalence of HBV infection among maintenance hemodialysis patients in the United States has declined dramatically and by 2002 was 0.12% and 1%, respectively.\textsuperscript{49} Data from 2002 indicated that newly acquired HBV infections were reported by 2.8% of US hemodialysis centers, and 27.3% of centers reported one or more chronically infected patients.\textsuperscript{49} New hepatitis B infections in hemodialysis patients are now rarely reported.

The chronically infected patient is central to the epidemiology of HBV transmission. HBV is transmitted by percutaneous (i.e., puncture through the skin) or mucosal (direct contact with mucous membranes) exposure to infectious blood or body fluids that contain blood. All hepatitis B surface antigen (HBsAg)–positive persons who are also positive for hepatitis Be antigen (HBeAg) have an extraordinary level of infectious virus in their blood, approximately $10^8$ to $10^9$ virions per milliliter.\textsuperscript{257,258} With virus titers this high in blood, HBV can transfer virus to susceptible patients through contaminated blood, blood or body fluids containing serum or blood may also contain high levels of HBV and are potentially infectious. Furthermore, HBV at titers of $10^2$ to $10^3$ virions/mL can be present on environmental surfaces in the absence of any visible blood and still cause infection.\textsuperscript{257,259-261}

HBV is relatively stable in the environment and has been found to remain viable for at least 7 days on environmental surfaces at room temperature.\textsuperscript{257,259,261} HBsAg has been detected in dialysis facilities on hemostats, scissors, dialysis machine control panels, and door knobs.\textsuperscript{261} Thus blood-contaminated surfaces that are not routinely cleaned and disinfected represent a reservoir for HBV transmission. Dialysis staff members can transfer virus to susceptible patients through contamination in the environment.\textsuperscript{257,259,261}

Most HBV outbreaks among hemodialysis patients (see Table 25.3) were caused by cross-contamination to patients via (1) environmental surfaces, supplies (e.g., hemostats, clamps, etc.), or equipment that were not routinely clean and disinfected after each use; (2) multiple-dose vials or intravenous solutions that were not used exclusively for one patient; (3) medications for injections that were prepared adjacent to areas where blood samples were handled; and (4) staff members who simultaneously provided care for both infected (HBsAg-positive) patients and susceptible patients.\textsuperscript{106,262-268} Once the factors that promote HBV transmission among hemodialysis patients were identified, recommendations for control were published.\textsuperscript{253}

The segregation of HBsAg-positive patients and their equipment from HBV-susceptible patients resulted in 70% to 80% reduction in the incidence of HBV infections among hemodialysis patients.\textsuperscript{255,269,270} The success of isolation practices in preventing transmission of HBV infection is linked to other infection control practices, including routine

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**HEPATITIS B VIRUS**

HBV is the most highly efficiently transmitted pathogen in the dialysis setting. Recommendations for control of hepatitis B in hemodialysis setting were first published in 1977,\textsuperscript{253} and by 1980 their widespread implementation was associated with a sharp decrease in the incidence of HBV infection among both patients and staff members.\textsuperscript{254,255} In 1982 the hepatitis B vaccine was recommended for all susceptible patients and staff members.\textsuperscript{256} Hepatitis B vaccination is currently the standard...
serological surveillance and routine cleaning and disinfection. Frequent serological testing for HBsAg detects patients recently infected with HBV so that isolation procedures can be implemented before cross-contamination can occur. Environmental control by routine cleaning and disinfection procedures reduces the opportunity for cross contamination, either directly from environmental surfaces or indirectly by hands of personnel.

In past studies, independent risk factors among maintenance hemodialysis patients for acquiring HBV infection included the presence of ≥1 HBV-infected patient in the hemodialysis facility who was not isolated, as well as a vaccination rate <50% among patients. However, transmission has been rarely reported in the United States in the past 20 years because of high rates of vaccination, screening, and isolation. The most recent documented transmission in a dialysis clinic in the United States was due to reactivation of hepatitis B infection that occurred in a patient with previous infection who became antigen positive as a result of immunosuppression. The CDC has received anecdotal reports of atypical hepatitis B serological test results among dialysis patients that may represent reactivation of HBV infection or HBV mutant strains; however, no further cases of dialysis-related transmission have been identified.

Other risk factors for acquiring HBV infection include injection drug use, sexual and household exposure to HBV-infected contacts, exposure to multiple sexual partners, male homosexual activity, and perinatal exposure. Dialysis patients should be educated about these and other risks and, for those patients with active HBV infection (HBsAg positive), informed that their sexual partners and household contacts should be vaccinated. HBV-infected patients should be evaluated for HBV treatment.

**Screening and Diagnostic Tests**

Several well-defined antigen-antibody systems are associated with HBV infection, including HBsAg and antibody to HBsAg (anti-HBs); hepatitis B core antigen (HBcAg) and antibody to HBcAg (anti-HBc); and HBeAg and antibody to HBeAg (anti-HBe). Serological assays are commercially available for all of these except for HBcAg because no free HBcAg circulates in the blood. One or more of these serological markers are present during different phases of HBV infection (Table 25.5). HBV infection can also be detected using qualitative or quantitative tests for HBV DNA. These tests are most commonly used for HBV-infected patients being managed with antiviral therapy.

In some individuals the only HBV serological marker detected is total anti-HBc (i.e., isolated anti-HBc). Among most asymptomatic persons in the United States tested for HBV infection, an average of 2% (range: <0.1% to 6%) test positive for anti-HBc; among injecting drug users, however, the rate is 24% to 28%. This pattern can occur after HBV infection among individuals who have recovered but whose anti-HBs have waned or among individuals who have low-level chronic HBV infection and failed to develop anti-HBs. It may also represent a false positive total anti-HBc result or someone in the window of infection. HBV DNA has been detected in <10% of individuals with isolated anti-HBc, and these individuals are unlikely to be infectious to others except under unusual circumstances involving direct percutaneous exposures to large quantities of blood (e.g., transfusion). In most persons with isolated anti-HBc, the result appears to be false positive. Data from several studies have indicated that a primary anti-HBs response develops in most of these individuals after a three-dose series of hepatitis B vaccinations. No data exist on response to vaccination among hemodialysis patients with this serological pattern. Testing and follow-up recommendations for hemodialysis patients with isolated anti-HBc are available.

**Prevention of Hepatitis B Virus Transmission**

The following recommendations can be applied to prevent transmission of HBV in hemodialysis facilities: (1) serological screening of patients (and staff members) for HBV infection, including monthly testing of all susceptible patients for HBsAg; (2) HBV vaccination of susceptible patients (and patient care staff); (3) isolation of all HBsAg-positive patients in a separate room; (4) assignment of staff members to HBsAg-positive patients and not to HBV-susceptible patients during the same or overlapping shifts; (5) assignment of dedicated dialysis equipment to HBsAg-positive patients; (6) cleaning and disinfection of nondisposable items (e.g., hemostats, clamps, scissors) before use on another patient; (7) glove use whenever patient or hemodialysis equipment is touched and glove changes and hand hygiene between each

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**TABLE 25.5 Interpretation of Serological Test Results for Hepatitis B Virus Infection**

| SEROLOGIC MARKERS | Total | Anti-HBc | Anti-HBc | Anti-HBs | Interpretation |
|-------------------|-------|----------|----------|----------|---------------|
| HBsAg – | –     | –        | –        | –        | Susceptible, never infected |
| +     | –     | –        | –        | –        | Acute infection, early incubation |
| +     | +     | +        | –        | –        | Acute infection |
| –     | +     | +        | –        | –        | Acute resolving infection |
| –     | –     | –        | –        | +        | Past infection, recovered and immune |
| +     | +     | –        | –        | –        | Chronic infection |
| –     | +     | –        | –        | –        | False positive (i.e., susceptible, past infection, or low-level chronic infection) |
| +     | –     | –        | +        | –        | Immune if titer ≥ 10 mIU/mL |

Anti-HBc, Antibody to hepatitis B core antigen; Anti-HBs, antibody to hepatitis B surface antigen; HBsAg, hepatitis B surface antigen; IgM, Immunoglobulin M.
patient (and station); and (8) routine cleaning and disinfection of equipment and environmental surfaces.298,299 Because dialysis patients can have waning immunity to hepatitis B vaccine, patients who require one or more booster doses of vaccine should not be cared for by the same staff as infected patients.

**Hepatitis Delta Virus**

Delta hepatitis is caused by the hepatitis delta (HDV), a relatively small defective virus that causes infection only in persons with active HBV infection. The prevalence of HDV infection is extremely low in the United States, with rates <1% among HBsAg-positive persons in the general population and >10% among HBsAg-positive persons with repeated percutaneous exposures (e.g., intravenous drug users, persons with hemophilia).291 Only one transmission of HDV among dialysis patients has been reported in the United States.290 In this episode, transmission occurred from a patient who was chronically infected with HBV and HDV to an HBsAg-positive patient after a massive bleeding incident; both patients received dialysis at the same station. Therefore, in dialysis settings, HDV-infected patients should be isolated from other HBV-infected patients.

**Human Immunodeficiency Virus Infection**

During 1985–2002, the percentage of US hemodialysis centers that reported providing maintenance hemodialysis for patients with HIV infection increased from 11% to 39% and the proportion of patients with known HIV infection increased from 0.3% to 1.5%.49 Although the proportion of patients with HIV infection has remained stable during the past decade, the number of infected patients has increased, as has the number of centers treating patients with HIV infection. HIV is transmitted by blood and other body fluids that contain blood. No patient-to-patient transmission of HIV has been reported in a US hemodialysis center. However, there have been reports of transmission of HIV among patients in other countries. All these outbreaks have been attributed to several breaks in infection control: (a) reuse of access needles and inadequately disinfected equipment, (b) sharing of syringes among patients, and (c) and sharing of dialyzers among different patients.293-297 The most recent reported outbreak involved three new HIV infections and was associated with sharing of multidose heparin vials, inadequately disinfected hemodialysis equipment, and dialysis staff who used blood-contaminated gloves to manipulate vascular access for multiple patients.298 Adherence to recommended infection control practices is adequate to prevent HIV transmission in dialysis facilities.162

**Other Emerging Infections**

In 2014 the largest outbreak of Ebola in history occurred in West Africa. Healthcare personnel caring for Ebola patients are at high risk for becoming infected, and during this outbreak, a significant number of healthcare personnel acquired the virus.299 Other high-consequence pathogens have continued to be identified, including Candida auris, a yeast that is resistant to multiple antifungals.300 Dialysis center staff and management should prepare for the possible introduction of highly virulent pathogens into their communities and dialysis centers by developing contingency plans, improving baseline adherence to recommended infection prevention practices, and strengthening communication channels with public health departments. The CDC has released on its website recommendations for infection control to prevent transmission of C. auris (https://www.cdc.gov/fungal/diseases/candidiasis/c-auris-infection-control.html), including dialysis-specific recommendations.

**Summary of Recommendations and Future Directions**

Preventing transmission of pathogens and reducing healthcare-associated infections among maintenance hemodialysis patients requires implementation of a comprehensive infection control program that can support consistent adherence to infection control recommendations (Table 25.6; Boxes 25.2 to Box 25.4) among all staff members. Adherence to core prevention practices (see Table 25.4) has been found to sustainably reduce highly morbid bloodstream infections among dialysis patients with central venous catheters. An active infection control program is the foundation of these efforts. The components of such a program include routine

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**Table 25.6 Schedule for Routine Testing for Hepatitis B Virus and Hepatitis C Virus Infections**

| Patient Status                          | On Admission*       | Monthly | Semi-Annual | Annual |
|----------------------------------------|---------------------|---------|-------------|--------|
| All patients                           | HBsAg, Anti-HBc (total), Anti-HBs, Anti-HCV, ALT | HBsAg   |             |        |
| HBV susceptible, including vaccine nonresponders | HBsAg, Anti-HBc (total), Anti-HBs, Anti-HCV, ALT | Anti-HBs |             |        |
| Anti-HBs positive (≥10 mIU/mL), anti-HBc negative | HBsAg |             |             |        |
| Anti-HBs and Anti-HBc positive         | No additional testing is needed | ALT     |             | Anti-HCV |
| Anti-HCV negative                      |                     |         |             |        |

ALT, Alanine aminotransferase.
*Results of HBV testing should be known before patient begins dialysis.
实施感染防控和控制措施旨在预防血液透析相关的感染：(1) 每个透析单位应至少有一名工作人员具备基础感染控制知识和经验；(2) 感染预防培训应包括人员和患者；(3) 定期实施感染预防和控制实践；(4) 建立安全文化，包括前线人员的积极参与；(5) 定期进行血清学检测；(6) 感染监测应定期进行，并已开展相关工具和评估工具。

**BOX 25.2 Recommended Infection Control Practices for Hemodialysis Units**

**Infection Control Precautions for All Patients**
- Hand hygiene should be performed.
- Before and after having direct contact with a patient’s intact skin.
- After contact with blood, body fluids or excretions, mucous membranes, non-intact skin, or wound dressings.
- After contact with inanimate objects (including medical equipment) in the immediate vicinity of the patient.
- If hands will be moving from a contaminated-body site to a clean-body site during patient care.
- After glove removal.
- Wear disposable gloves when caring for the patient or touching the patient’s equipment at the dialysis station; remove gloves and perform hand hygiene (if hands are visibly soiled wash with soap and water) between each patient or station.
- Items taken into the dialysis station should be disposed of or cleaned and disinfected before taken to a common clean area or used on another patient.
- Nondisposable items taken to the patient treatment station that cannot be cleaned or disinfected (e.g., adhesive tape) should be discarded after use.
- Unused medications (including multi-dose vials) or supplies (syringes, alcohol swabs, etc.) taken to the patient’s station should be used only for that patient and should not be returned to a common clean area or used on other patients.
- When multidose medication vials are used (including vials containing diluents), prepare individual patient doses in a clean (centralized) area away from dialysis stations and deliver separately to each patient. Do not carry multidose medication vials from station to station.
- Do not use common medication carts to deliver medications to patients. Do not carry medication vials, syringes, alcohol swabs, or supplies in pockets. If trays are used to deliver medication to individual patients, they must be cleaned between patients.
- Clean areas should be clearly designated for the preparation, handling, and storage of medications and unused supplies and equipment. Clean areas should be clearly separated from contaminated areas where used supplies and equipment are handled. Do not handle and store medications or clean supplies in the same or adjacent area to where used equipment or blood samples are handled.
- Use external transducer protectors (venous or arterial) for each patient treatment to prevent blood contamination of the dialysis machine’s pressure monitoring equipment. Change these external transducer protectors between each patient treatment and when they become wet, and do not reuse them. The redundant internal transducer protectors do not need to be changed routinely between patients. If the external transducer protectors are contaminated with blood the internal transducer protector should be assessed for contamination before dialyzing another patient with the same machine.
- Clean and disinfect the dialysis station (chairs, beds, tables, machines, etc.) between patients.
- Start cleaning only when patient has left the station and only admit new patient after cleaning and disinfection are complete.
- Give special attention to cleaning control panels on the dialysis machine and other surfaces that are frequently touched and potentially contaminated with patient’s blood.
- Discard all fluid, and clean and disinfect all surfaces and containers associated with the prime waste (including buckets attached to the machines).
- For dialyzers and blood tubing that will be reprocessed, cap dialyzer ports and clamp tubing. Place all used dialyzers and tubing in a leak-proof containers for transport from station to reprocessing or disposal area.

**BOX 25.3 Hepatitis B Vaccination**

- Vaccinate all susceptible patients against hepatitis B.
- Test for anti-HBs 1–2 months after the last dose.
- If anti-HBs is <10 mIU/mL, consider patient susceptible, revaccinate with an additional three doses, and retest for anti-HBs.
- If anti-HBs is >10 mIU/mL, consider immune and retest annually.
- Give booster dose of vaccine if anti-HBs declines to <10 mIU/mL and continue to retest annually.

**BOX 25.4 Management of HBsAg-Positive Patients**

- Follow infection control practices for hemodialysis units for all patients.
- Dialyze HBsAg-positive patients in a separate room using separate machines, equipment, instruments, and supplies.
- Staff members caring for HBsAg-positive patients should not care for HBV-susceptible patients at the same time (e.g., during same shift or during patient change over).
- FDA Safety Alert. Modified from Centers for Disease Control and Prevention, Recommendations for preventing transmission of infections among chronic hemodialysis patients. MMWR Recomm Rep, 2001. 50(RR-5): pp. 1-43.

https://www.cdc.gov/dialysis/prevention-tools/index.html; (4) a culture of safety should be developed, including engaged leadership and involvement of frontline staff in infection prevention efforts; (5) routine serological testing and immunization of patients and staff should be performed; (6) infection surveillance should be conducted and the data
used for continuous quality improvement; and (7) systems should be in place for public health reporting. An excellent review of those essential components of an infection prevention program is available elsewhere. The CDC has also published recommendations describing these components in detail.

Future Directions

Infection control strategies that prevent HBV infection among hemodialysis patients have been well established; however, some questions remain. More work is needed to determine the ideal hepatitis B vaccine dosage regimen for pre- and postdialysis pediatric patients and for predialysis adult patients, as well as the optimal timing for follow-up testing and administration of booster doses among vaccine responders. Also, reports of patients with mutant HBV, patients with reverse seroconversion, and patients with atypical HBV serological test results highlight the need for more research to evaluate their significance in dialysis population. Further work is needed to clarify the specific factors responsible for transmission of HCV among hemodialysis patients and to evaluate the effect of current prevention recommendations, HCV treatment, and other strategies on prevention and control of HCV infection in this setting.

VAIs continue to be a devastating complication among patient receiving maintenance hemodialysis; additional interventions are needed to further reduce rates of these infections. Finally, other important questions about the role dialysis centers play in the spread of MDROs and the effectiveness of interventions designed to prevent MDRO transmission require further investigation.

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REFERENCES

1. United States Renal Data System. USRDS 2016 Annual Data Report: Atlas of Chronic Kidney Disease and End-Stage Renal Disease in the United States. Bethesda, MD: National Institutes of Health, National Institute of Diabetes and Digestive and Kidney Diseases; 2016.

2. Khan IH, Catto GR. Long-term complications of dialysis: infection. Kidney Int Suppl. 1993;41:S143–S148.

3. Vanholder R, Ringoir S. Polymorphonuclear cell function and infection in dialysis. Kidney Int Suppl. 1992;38:S91–S95.

4. Hindmash SH, et al. Pyrogenic reactions during haemodialysis caused by extramural endotoxin. Lancet. 1975;2(7938):732–734.

5. Gordon SM, et al. Pyrogenic reactions in patients receiving conventional, high-efficiency, or high-flux hemodialysis treatments with bicarbonate dialysate containing high concentrations of bacteria and endotoxin. J Am Soc Nephrol. 1992;2(9):1436–1444.

6. Favero MS, et al. Gram-negative water bacteria in hemodialysis systems. Health Lab Sci. 1975;12(4):321–334.

7. Archibald LK, et al. Pyrogenic reactions in hemodialysis patients, Hanoi, Vietnam. Infect Control Hosp Epidemiol. 2006;27(4):424–426.

8. Rudnick JR, et al. An outbreak of pyrogenic reactions in chronic hemodialysis patients associated with hemodialyzer reuse. Artif Organs. 1995;19(4):289–294.

9. Jackson BM, et al. Outbreak of pyrogenic reactions and gram-negative bacteremia in a hemodialysis center. Am J Nephrol. 1994;14(2):85–89.

10. Beck-Sague CM, et al. Outbreak of gram-negative bacteremia and pyrogenic reactions in a hemodialysis center. Am J Nephrol. 1990;10(5):397–403.

11. Gordon SM, et al. Pyrogenic reactions associated with the reuse of disposable hollow-fiber hemodialyzers. JAMA. 1988;260(14):2077–2081.

12. Roth VR, Jarvis WR. Outbreaks of infection and/or pyrogenic reactions in dialysis patients. Semin Dial. 2000;13(2):92–96.

13. Pollak VE. Adverse effects and pyrogenic reactions during hemodialysis. JAMA. 1988;260(14):2106–2107.

14. Lefton C. Patients suffer pyrogenic reactions in Philadelphia dialysis units. Nephrol News Issues. 1994;8(12):10.

15. Nystrand R. The microbial world and fluids in dialysis. Biomed Instrum Technol. 2008;42(2):150–159.

16. Petersen NJ, Carson LA, Favero MS. Bacterial endotoxin in new and reused hemodialyzers: a potential cause of endotoxemia. Trans Am Soc Artif Intern Organs. 27:155–160.

17. Rai J, Shapiro FL, Michael AF. Endotoxemia in febrile reactions during hemodialysis. Kidney Int. 1973;4(1):57–60.

18. Favero MS, et al. Factors that influence microbial contamination of fluids associated with hemodialysis machines. Appl Microbiol. 1974;28(5):822–830.

19. Favero MS, et al. Pseudomonas aeruginosa: growth in distilled water from hospitals. Science. 173(999):836–838.

20. Favero, M.S. and Bland, L.A. eds. Microbiologic principles applied to reprocessing hemodialyzers. Guide to reprocessing of hemodialyzers. ed. Deane N, Wineman R, and Bemis J. 1986, Martinus Nijhoff: Boston. 63–73.

21. Carson LA, et al. Prevalence of nontuberculous mycobacteria in water supplies of hemodialysis centers. Appl Environ Microbiol. 1988;54(12):3122–3125.

22. Bolan G, et al. Infections with Mycobacterium chelonae in patients receiving dialysis and using processed hemodialyzers. J Infect Dis. 1985;152(5):1013–1019.

23. Gomila M, Ramirez A, Lalucat J. Diversity of environmental Mycobacterium isolates from hemodialysis water as shown by a multigene sequencing approach. Appl Environ Microbiol. 2007;73(12):3787–3797.

24. Basok A, et al. Spectrum of mycobacterial infections: tuberculosis and Mycobacterium other than tuberculosis in dialysis patients. Isr Med Assoc J. 2007;9(6):448–451.

25. Centers for Disease Control and Prevention. Nontuberculous mycobacterial infections in hemodialysis patients–Louisiana, 1982. MMWR Morb Mortal Wkly Rep. 1983;32(18):244–246.

26. Lowry PW, et al. Mycobacterium chelonae infection among patients receiving high-flux dialysis in a hemodialysis clinic in California. J Infect Dis. 1990;161(1):85–90.

27. Bland LA, Favero MS, eds. Microbial contamination control strategies for hemodialysis systems. Plant, Technology, and Safety Management Series. vol. 3. Oakland Terrace: Joint Commission on Accreditation of Healthcare Organizations; 1989.

28. Favero MS, et al. Microbial contamination of renal dialysis systems and associated health risks. Trans Am Soc Artif Intern Organs. 1974;20A:175–183.

29. Carson LA, et al. Factors affecting endotoxin levels in fluids associated with hemodialysis procedures. Prog Clin Biol Res. 1987;231:223–234.

30. AAMI. American National Standard: Reuse of Hemodialyzers. Arlington, VA: Association for the Advancement for Medical Instrumentation; 2008.

31. AAMI. Recommended Practice: Water for hemodialysis and related therapies. Arlington, VA: Association for the Advancement of Medical Instrumentation; 2014.

32. AAMI. Guidance for the preparation and quality management of fluids for hemodialysis and related therapies. Arlington, VA: Association for the Advancement of Medical Instrumentation; 2014.

33. AAMI. Quality of dialysis fluids for hemodialysis and related therapies. Arlington, VA: Association for the Advancement of Medical Instrumentation; 2014.

34. Centers for Medicare & Medicaid Services. Medicare and Medicaid programs; conditions for coverage for end-stage renal disease facilities. Final rule. Fed Regist. 2008;73(73):20369–20484.

35. Stamm JM, Engelhard WE, Parsons JE. Microbiological study of water-softerner resins. Applied Microbiology. 1969;18:376–386.

36. Anderson RL, Holland BW, Carr JK. Effect of disinfectants on pseudomonads colonized on the interior surfaces of PVC Pipes. Am J Public Health. 1990;80:17–21.

37. AAMI. American National Standard: Water treatment equipment for hemodialysis and related therapies. Arlington, VA: Association for the Advancement of Medical Instrumentation; 2014.

38. Arduino MJ. Microbiologic quality of water used for hemodialysis. Contemp Dial Nephrol. 1996;17:17–19.

39. Arduino MJ. What's new in water treatment standards for hemodialysis. Contemp Dial Nephrol. 1997;18:21–24.

40. Petersen NJ, et al. Pyrogenic reactions from inadequate disinfection of a dialysis fluid distribution system. Dialysis Transplant. 1978;7:52–57.
REFERENCES

41. Jochimsen EM, et al. A cluster of bloodstream infections and pyrogenic reactions among hemodialysis patients traced to dialysis machine waste-handling option units. Am J Nephrol. 1998;18(6):485–489.

42. Wang SA, et al. An outbreak of gram-negative bacteremia in hemodialysis patients traced to hemodialysis machine waste drain ports. Infect Control Hosp Epidemiol. 1999;20(11):746–751.

43. Rao CY, et al. An Outbreak of Phialeonium Mold Infections in Hemodialysis Patients: When Purified Water Is Not So Pure – Illinois, 2005, 55th Annual Epidemic Intelligence Service Conference. Atlanta, GA: CDC; 2006.

44. Favero MS, P.N., Boyer KM, Carson LA, Bond WW. Microbial contamination of renal dialysis systems and associated health risks. Trans Am Soc Artif Intern Organs. 1974;20A:175–183.

45. Centers for Disease Control and Prevention. Pyrogenic reactions and gram-negative blood-stream infections in hemodialysis patients. In: Services HaH, ed. Epidemiologic Investigation Report. Atlanta: Department of Health and Human Services; 1991.

46. Nguyen DB, et al. A large outbreak of hepatitis C virus infections in a hemodialysis clinic. Infect Control Hosp Epidemiol. 2016;37(2):125–133.

47. Novosad S, et al. Unusual Source of Gram-negative Blood-stream Infections in Hemodialysis Patients, in SHEA Spring 2017 Conference; 2017. St. Louis, MO.

48. Tokars JI, et al. National surveillance of dialysis-associated diseases in the United States, 2000. Semin Dial. 2002;15(3):162–171.

49. Finelli L, et al. National surveillance of dialysis-associated diseases in the United States, 2002. Semin Dial. 2005;18(1):52–61.

50. Toniolo Ado R, et al. Evaluation of the effectiveness of manual and automated dialyzers reprocessing after multiple reuses. Am J Infect Control. 2016;44(6):719–720.

51. Prasad N, Jha V. Hemodialysis in Asia. Kidney Dis (Basel). 2015;1(3):165–177.

52. Edens C, et al. Hemodialyzer reuse and gram-negative blood-stream infections. Am J Kidney Dis. 2017;69(6):726–733.

53. AAMI. Reuse of hemodiayzers. Arlington, VA: Association for the Advancement of Medical Instrumentation; 1986.

54. Bland L, et al. Hemodialyzer reuse: practices in the United States and implication for infection control. Trans Am Soc Artif Intern Organs. 1985;31:556–559.

55. Arduino MJ. How should dialyzers be reprocessed? Seminars in Dialysis. 1998;11(5):282–284.

56. Favero MS. Distinguishing between high-level disinfection, reprocessing, and sterilization. In: Easterling RE, ed. Reuse of Disposables: Implications for Quality Health Care and Cost Containment. Arlington, VA: AAMI; 1983:19–23.

57. Oyong K, et al. Outbreak of bloodstream infections associated with multiuse dialyzers containing O-rings. Infect Control Hosp Epidemiol. 2014;35(1):89–91.

58. Rosenberg, J. Primary Bloodstream Infections Associated with Dialyzer Reuse in California Dialysis Centers in IDSA 2005; October 6-9, 2005; San Francisco, CA. 2005: IDSA 2005; October 6-9, 2005; San Francisco, CA.

59. Welbel SF, et al. An outbreak of gram-negative bloodstream infections in chronic hemodialysis patients. Am J Nephrol. 1995;15(1):1–4.

60. Bland LA, et al. Potential bacteriologic and endotoxin hazards associated with liquid bicarbonate concentrate. ASAIO Trans. 1987;33(3):542–545.

61. Tokars JI, et al. National surveillance of dialysis-associated diseases in the United States, 1995. Atlanta, GA: US Department of Health and Human Services; 1998.

62. Petersen NJ, et al. Microbiologic evaluation of a new glutaraldehyde-based disinfectant for hemodialysis systems. Trans Am Soc Artif Intern Organs. 1982;28:287–290.

63. Townsend TR, Wei SB, Bartlett J. Disinfection of hemodialysis machines. Dialysis and Transplantation. 1985;14:274–287.

64. Centers for Disease Control and Prevention. Occupational exposures to formaldehyde in dialysis units. MMWR Morb Mortal Wkly Rep. 1986;35(24):399–401.

65. AAMI. Recommended Practice: Dialysate for Hemodialysis. Arlington, VA: Association for the Advancement of Medical Instrumentation; 2004.

66. AAMI. American National Standard: Water Treatment Equipment for Hemodialysis Applications. Arlington, VA: Association for the Advancement of Medical Instrumentation; 2006.

67. Favero MS, Petersen NJ. Microbiologic guidelines for hemodialysis systems. Dialysis and Transplantation. 1977;6:34–36.

68. Bommer J, Jaber BL. Ultrapure dialyse: facts and myths. Semin Dial. 2006;19(2):115–119.

69. Ward RA. Ultrapure dialyse. Semin Dial. 2004;17(6):489–497.

70. Izuhara Y, et al. Ultrapure dialyse decreases plasma pentosidine, a marker of “carbonyl stress”. Am J Kidney Dis. 2004;43(6):1024–1029.

71. Johnson DW, Pollock CA, Macdougall IC. Erythropoiesis-stimulating agent hyporesponsiveness. Nephrology (Carlton). 2007;12(4):321–330.

72. Go I, et al. The effect of ultrapure dialyse on improving renal anemia. Osaka City Med J. 2007;53(1):17–23.

73. Lacson Jr L, Levin NW. C-reactive protein and end-stage renal disease. Semin Dial. 2004;17(6):438–448.

74. Sitter T, Bergner A, Schiff H. Dialysate related cytokine induction and response to recombinant human erythropoietin in haemodialysis patients. Nephrol Dial Transplant. 2000;15(8):1207–1211.

75. Schiff H, et al. Effects of ultrapure dialysis fluid on nutritional status and inflammatory parameters. Nephrol Dial Transplant. 2001;16(9):1863–1869.

76. Hsu PY, et al. Ultrapure dialyse improves iron utilization and erythropoietin response in chronic hemodialysis patients—a prospective cross-over study. J Nephrol. 2004;17(5):693–700.

77. Furuya R, et al. Ultrapure dialyse reduces plasma levels of beta2-microglobulin and pentosidine in hemodialysis patients. Blood Purif. 2005;23(4):311–316.

78. Schiff H, Lang SM, Fischer R. Ultrapure dialysis fluid slows loss of residual renal function in new dialysis patients. Nephrol Dial Transplant. 2002;17(10):1814–1818.

79. Schiff H, Wendinger H, Lang SM. Ultrapure dialysis fluid and response to hepatitis B vaccine. Nephron. 2002;91(3):530–531.

80. Schiff H, Lang SM, Bergner A. Ultrapure dialyse reduces dose of recombinant human erythropoietin. Nephron. 1999;83(3):278–279.

81. Tielmans C, et al. Effects of ultrapure and non-sterile dialyse on the inflammatory response during in vitro hemodialysis. Kidney Int. 1996;49(1):236–243.

82. Baz M, et al. Using ultrapure water in hemodialysis delays carpal tunnel syndrome. Int J Artif Organs. 1991;14(11):681–685.
REFERENCES 410.e3

83. Upadhyay A, Jaber BL. We use impure water to make dialysate for hemodialysis. *Semin Dial.* 2016;29(4):297–299.
84. Upadhyay A, Susantitaphong P, Jaber BL. Ultrapure versus standard dialysate: a cost-benefit analysis. *Semin Dial.* 2017.
85. Hasegawa T, et al. Dialysis fluid endotoxin level and mortality in maintenance hemodialysis: a nationwide cohort study. *Am J Kidney Dis.* 2015;65(6):899–904.
86. AAML. *Water testing methodologies.* Arlington, VA: Association for the Advancement of Medical Instrumentation; 2014.
87. Pegues DA, et al. A prospective study of pyrogenic reactions in hemodialysis patients using bicarbonate dialysis fluids filtered to remove bacteria and endotoxin. *J Am Soc Nephrol.* 1992;3(4):1002–1007.
88. Alter MJ, et al. National surveillance of dialysis-associated diseases in the United States, 1989. *ASAIO Trans.* 1991;37(2):97–109.
89. Tokars JI, et al. National surveillance of dialysis associated diseases in the United States–1994. *ASAIO J.* 1997;43(1):108–119.
90. Tokars JI, et al. National surveillance of dialysis associated diseases in the United States, 1993. *ASAIO J.* 1996;42(3):219–229.
91. Tokars JI, et al. National surveillance of dialysis associated diseases in the United States, 1992. *ASAIO J.* 1994;40(4):1020–1031.
92. Tokars JI, et al. National surveillance of dialysis associated diseases in the United States, 1991. *ASAIO J.* 1993;39(4):966–975.
93. Tokars JI, et al. National surveillance of hemodialysis associated diseases in the United States, 1990. *ASAIO J.* 1993;39(1):71–80.
94. Alter MJ, et al. National surveillance of dialysis-associated diseases in the United States, 1987. *ASAIO Trans.* 1989;35(4):820–831.
95. Tokars JI, et al. National surveillance of dialysis-associated diseases in the United States, 2001. *Semin Dial.* 2004;17(4):310–319.
96. Tokars JI, et al. National surveillance of dialysis associated diseases in the United States, 1995. *ASAIO J.* 1998;44(1):98–107.
97. Tokars JI, et al. National surveillance of dialysis-associated diseases in the United States, 1997. *Semin Dial.* 2000;13(2):75–85.
98. David S, et al. Production of platelet activating factor by human neutrophils after backfiltration of endotoxin contaminated dialysate. *ASAIO J.* 1993;39(3):M773–7.
99. Laude-Sharp M, et al. Induction of IL-1 during hemodialysis: transmembrane passage of intact endotoxins (LPS). *Kidney Int.* 1990;38(6):1089–1094.
100. Yamagami S, et al. Detection of endotoxin antibody in long-term dialysis patients. *Int J Artif Organs.* 1990;13(4):205–210.
101. Gazenfeld-Gazit E, Eliahou HE. Endotoxin antibodies in patients on maintenance hemodialysis. *Isr J Med Sci.* 1969;5(5):1032–1036.
102. Sundaram S, et al. Transmembrane passage of cytokine-inducing bacterial products across new and reprocessed polysulfone dialyzers. *J Am Soc Nephrol.* 1996;7(10):2183–2191.
103. Arduino MJ, et al. The effects of endotoxin-contaminated dialysate and polysulfone or cellulose membranes on the release of TNF alpha during simulated dialysis. *Artif Organs.* 1995;19(9):880–886.
104. Henderson LW, et al. Haemodialysis hypotension: the interleukin hypothesis. *Blood Purification.* 1993;1:3–7.
105. Port FK, et al. The role of dialysate in the stimulation of interleukin-1 production during clinical hemodialysis. *Am J Kidney Dis.* 1987;10(2):118–122.
106. Kantor RJ, et al. Outbreak of pyrogenic reactions at a dialysis center. Association with infusion of heparinized saline solution. *Am J Med.* 1983;74(3):449–456.
107. Oliver JC, et al. Bacteria and endotoxin removal from bicarbonate dialysis fluids for use in conventional, high-efficiency, and high-flux hemodialysis. *Artif Organs.* 1992;16(2):141–145.
108. Favero MS, Bolyard EA. Microbiological considerations. Disinfection and sterilization strategies and the potential for airborne transmission of bloodborne pathogens. *Surg Clin North Am.* 1995;75(6):1071–1089.
109. Favero MS, Bond, WW eds. Chemical disinfection of medical and surgical materials. In: Block SS, ed. *Disinfection, Sterilization, and Preservation.* 4th ed. Philadelphia: Lea & Febiger; 1991:617–641.
110. Schulster L, Chinn RY. Guidelines for environmental infection control in health-care facilities. Recommendations of CDC and the Healthcare Infection Control Practices Advisory Committee (HICPAC). MMWR Recomm Rep. 2003;52(RR-10):1–42.
111. Facility Guidelines Institute. *Guidelines for Design and Construction of Hospitals and Outpatient Facilities.* 2014.
112. Bergervoet PW, et al. Application of the forensic Luminol for blood in infection control. *J Hosp Infect.* 2008;68(4):329–333.
113. Bloemmenbergen WE, Port FK. Epidemiological perspective on infections in chronic dialysis patients. *Adv Ren Replace Ther.* 1996;3(3):201–207.
114. Dobkin JF, Miller MH, Steigbigel NH. Septicemia in patients on chronic hemodialysis. *Ann Intern Med.* 1978;88(1):28–33.
115. Hoen B, et al. EPIBACDIAL: a multicenter prospective study of risk factors for bacteremia in chronic hemodialysis patients. *J Am Soc Nephrol.* 1998;9(5):869–876.
116. Bonomo RA, et al. Risk factors associated with permanent access-site infections in chronic hemodialysis patients. *Infect Control Hosp Epidemiol.* 1997;18(11):757–761.
117. Kaplowitz LG, et al. A prospective study of infections in hemodialysis patients: patient hygiene and other risk factors for infection. *Infect Control Hosp Epidemiol.* 1988;9(12):534–541.
118. Keane WF, Shapiro FL, Raji L. Incidence and type of infections occurring in 445 chronic hemodialysis patients. *Trans Am Soc Artif Intern Organs.* 1977;23:41–47.
119. Kessler M, et al. Bacteremia in patients on chronic hemodialysis. A multicenter prospective survey. *Nephron.* 1993;64(1):95–100.
120. Stevenson KB, et al. Epidemiology of hemodialysis vascular access infections from longitudinal infection surveillance data: predicting the impact of NKF-DOQI clinical practice guidelines for vascular access. *Am J Kidney Dis.* 2002;39(3):549–555.
121. Tokars JI, et al. A prospective study of vascular access infections at seven outpatient hemodialysis centers. *Am J Kidney Dis.* 2001;37(6):1232–1240.
122. Tokars JI, Miller ER, Stein G. New national surveillance system for hemodialysis-associated infections: initial results. *Am J Infect Control.* 2002;30(5):288–295.
123. Ponce P, et al. A prospective study on incidence of bacterial infections in portuguese dialysis units. *Nephron Clin Pract.* 107(4):c133–8.
REFERENCES

124. Tokars JI. Bloodstream infections in hemodialysis patients: getting some deserved attention. Infect Control Hosp Epidemiol. 2002;23(12):713–715.

125. Centers for Disease Control and Prevention. Dialysis Event Protocol. 4/25/2016: Available from: http://www.cdc.gov/nhsn/PDFs/pscManual/8pscDialysisEventcurrent.pdf.

126. Patel PR, et al. Dialysis event surveillance report: national healthcare safety network data summary, January 2007 through April 2011. Am J Infect Control. 2016;44(8):944–947.

127. Nguyen DB, et al. National healthcare safety network (NHSN) dialysis event surveillance report for 2014. Clin J Am Soc Nephrol. 2017;12(7):1139–1146.

128. Padberg Jr FT, Lee BC, Curr GR. Hemoaccess site infection. Surg Gynecol Obstet. 1992;174(2):103–108.

129. Klevens RM, Tokars JI, Andrus M. Electronic reporting of infections associated with hemodialysis. Nephrol News Issues. 2005;19(7):37–38, 43.

130. Klevens RM, et al. Dialysis surveillance report: national healthcare safety network (NHSN)-data summary for 2006. Seminars in Dialysis. 2008;21(1):24–28.

131. Thomson PC, et al. Vascular access in haemodialysis patients: a modifiable risk factor for bacteraemia and death. Qjm. 2007;100(7):415–422.

132. Lafrance JP, et al. Vascular access–related infections: definitions, incidence rates, and risk factors. Am J Kidney Dis. 2008.

133. Albers FJ. Clinical considerations in hemodialysis access infection. Adv Ren Replace Ther. 1996;3(3):208–217.

134. Powe NR, et al. Septicemia in dialysis patients: incidence, risk factors, and prognosis. Kidney Int. 1999;55(3):1081–1090.

135. Fan PY, Schwab SJ. Vascular access: concepts for the 1990s. J Am Soc Nephrol. 1992;3(1):11.

136. Gulati S, et al. Role of vascular access as a risk factor for infections in hemodialysis. Ren Fail. 2003;25(6):967–973.

137. NKF, NKF-DOQI clinical practice guidelines for vascular access. National Kidney Foundation-Dialysis Outcomes Quality Initiative. Am J Kidney Dis. 1997;30(4 suppl 3):S150–S191.

138. NKF, III. NKF-K-DOQI Clinical Practice Guidelines for Vascular Access: update 2000. Am J Kidney Dis. 2001;37(1 suppl 1):S137–S181.

139. NKF. Clinical practice recommendation 8: vascular access in pediatric patients. Am J Kidney Dis. 2006;48(suppl 1):S274–S276.

140. NKF. Clinical practice guidelines for vascular access. Am J Kidney Dis. 2006;48(suppl 1):S248–S273.

141. NKF. Clinical practice guidelines for vascular access. Am J Kidney Dis. 2006;48(suppl 1):S176–S247.

142. Pisoni RL, et al. Trends in US vascular access use, patient preferences, and related practices: an update from the US DOPPS practice monitor with international comparisons. Am J Kidney Dis. 2015;65(6):905–915.

143. Arnow PM, et al. An outbreak of bloodstream infections arising from hemodialysis equipment. J Infect Dis. 1998;178(3):783–791.

144. Block C, et al. Outbreak of bloodstream infections associated with dialysis machine waste ports in a hemodialysis facility. Eur J Clin Microbiol Infect Dis. 1999;18(10):723–725.

145. Centers for Disease Control and Prevention. Outbreaks of gram-negative bacterial bloodstream infections traced to probable contamination of hemodialysis machines—Canada, 1995; United States, 1997; and Israel, 1997. MMWR Morb Mortal Wkly Rep. 1998;47(3):55–59.

146. Tokars JI, A.M., Arduino MJ, eds. Nosocomial infections in hemodialysis units: strategies for control. The Principles and Practices of Nephrology. St. Louis: Mosby; 1995: 337–357.

147. Grohskopf LA, et al. Serratia liquefaciens bloodstream infections from contamination of epoetin alfa at a hemodialysis center. N Engl J Med. 2001;344(20):1491–1497.

148. Centers for Disease Control and Prevention. Guidelines for the prevention of intravascular catheter-related infections. MMWR Morb Mortal Wkly Rep. 2002;51(RR-10):1–29.

149. O’Grady NP, et al. Guidelines for the prevention of intravascular catheter-related infections. Clin Infect Dis. 2011;52(9):e162–93.

150. Patel PR, et al. Bloodstream infection rates in outpatient hemodialysis facilities participating in a collaborative prevention effort: a quality improvement report. Am J Kidney Dis. 2013;62(2):322–330.

151. Centers for Disease Control and Prevention. Reducing bloodstream infections in an outpatient hemodialysis center—New Jersey, 2008-2011. MMWR Morb Mortal Wkly Rep. 2012;61(10):169–173.

152. Dinwiddie L. Cleansing agents used for hemodialysis catheter care. Nephrol Nurs J. 2002;29(6):599–613.

153. Yahav D, et al. Antimicrobial lock solutions for the prevention of infections associated with intravascular catheters in patients undergoing hemodialysis: systematic review and meta-analysis of randomized, controlled trials. Clin Infect Dis. 2008;47(1):83–93.

154. Jaffer Y, et al. A meta-analysis of hemodialysis catheter locking solutions in the prevention of catheter-related infection. Am J Kidney Dis. 2008;51(2):233–241.

155. Berns JS, Tokars JI. Preventing bacterial infections and antimicrobial resistance in dialysis patients. Am J Kidney Dis. 2002;40(5):886–898.

156. Mermel LA. Prevention of intravascular catheter infections—insights and prospects for hemodialysis catheters. Nephrologie. 2001;22(8):449–451.

157. Mermel LA, et al. Guidelines for the management of intravascular catheter-related infections. Clin Infect Dis. 2001;32(9):1249–1272.

158. Camins BC, et al. A crossover intervention trial evaluating the efficacy of a chlorhexidine-impregnated sponge in reducing catheter-related bloodstream infections among patients undergoing hemodialysis. Infect Control Hosp Epidemiol. 2010;31(11):1118–1123.

159. Onder AM, et al. Controlling exit site infections: does it decrease the incidence of catheter-related bacteremia in children on chronic hemodialysis? Hemodial Int. 2009;13(1):11–18.

160. Hymes JL, et al. Dialysis catheter-related bloodstream infections: a cluster-randomized trial of the clearguard hd antimicrobial barrier cap. Am J Kidney Dis. 2017;69(2):220–227.

161. Brunelli SM, et al. Use of the Tego needlefree connector is associated with reduced incidence of catheter-related bloodstream infections in hemodialysis patients. Int J Nephrol Renovasc Dis. 2014;7:131–139.

162. Centers for Disease Control and Prevention. Recommendations for preventing transmission of infections among chronic hemodialysis patients. MMWR Recom. Rep. 2001;50(RR-5):1–43.
163. Centers for Disease Control and Prevention. Infection control requirements for dialysis facilities and clarification regarding guidance on parenteral medication vials. MMWR Morb Mortal Wkly Rep. 2008;57(32):875–876.

164. United States Renal Data System. USRDS 2008 Annual Data Report: Atlas of Chronic Kidney Disease and End-Stage Renal Disease in the United States. Bethesda, MD: National Institutes of Health, National Institute of Diabetes and Digestive and Kidney Diseases; 2008.

165. Berman SJ, et al. Burden of infection in patients with end-stage renal disease requiring long-term dialysis. Clin Infect Dis. 2004;39(12):1747–1753.

166. Slinin Y, Foley RN, Collins AJ. Clinical epidemiology of pneumonia in hemodialysis patients: the USRDS waves 1, 3, and 4 study. Kidney Int. 2006;70(6):1135–1141.

167. Guo H, et al. Pneumonia in incident dialysis patients—the United States Renal Data System. Nephrol Dial Transplant. 2008;23(2):680–686.

168. Centers for Disease Control and Prevention. Use of 13-valent pneumococcal conjugate vaccine and 23-valent pneumococcal polysaccharide vaccine for adults with immunocompromising conditions: recommendations of the Advisory Committee on Immunization Practices (ACIP). MMWR Morb Mortal Wkly Rep. 2012;61(40):816–819.

169. Grohskopf LA, et al. Prevention and control of seasonal influenza with vaccines. MMWR Recomm Rep. 2016;65(5):1–54.

170. Gilbertson DT, et al. Influenza vaccine delivery and effectiveness in end-stage renal disease. Kidney Int. 2003;63(2):738–743.

171. Van Buynder PG, et al. Healthcare worker influenza immunization vaccine or mask policy: strategies for cost effective implementation and subsequent reductions in staff absenteeism due to illness. Vaccine. 2015;33(13):1625–1628.

172. Saxen H, Virtanen M. Randomized, placebo-controlled double blind study on the efficacy of influenza immunization on absenteeism of health care workers. Pediatr Infect Dis J. 1999;18(9):779–783.

173. Salgado CD, et al. Preventing nosocomial influenza by improving the vaccine acceptance rate of clinicians. Infect Control Hosp Epidemiol. 2004;25(11):923–928.

174. Lynch JR, et al. Correlates of change in health care worker seasonal influenza vaccination rates among dialysis facilities. Am J Infect Control. 2015;43(4):409–411.

175. Jensen PA, et al. Guidelines for preventing the transmission of Mycobacterium tuberculosis in health-care settings, 2005. MMWR Recomm Rep. 2005;54(RR-17):1–141.

176. Centers for Disease Control and Prevention. Tuberculosis transmission in a renal dialysis center—Nevada, 2003. MMWR Morb Mortal Wkly Rep. 2004;53(37):873–875.

177. Linquist JA, et al. Tuberculosis exposure of patients and staff in an outpatient hemodialysis unit. Am J Infect Control. 2002;30(5):307–310.

178. Zaki AM, et al. Isolation of a novel coronavirus from a man with pneumonia in Saudi Arabia. N Engl J Med. 2012;367(19):1814–1820.

179. Kallen AJ, Arduino MJ, Patel PR. Preventing infections in patients undergoing hemodialysis. Expert Rev Anti Infect Ther. 2010;8(6):643–655.

180. Utley AH, et al. High-level vancomycin-resistant enterococci causing hospital infections. Epidemiol Infect. 1989;103(1):173–181.

181. Gray JW, George RH. Experience of vancomycin-resistant enterococci in a children’s hospital. J Hosp Infect. 2000;45(1):11–18.

182. Brady JP, Snyder JW, Hasbargen JA. Vancomycin-resistant enterococci in end-stage renal disease. Am J Kidney Dis. 1998;32(3):415–418.

183. Roghmann MC, et al. Colonization with vancomycin-resistant enterococci in chronic hemodialysis patients. Am J Kidney Dis. 1998;32(2):254–257.

184. Fishbane S, et al. Vancomycin-resistant enterococci in hemodialysis patients is related to intravenous vancomycin use. Infect Control Hosp Epidemiol. 1999;20(7):461–462.

185. Zacharioudakis IM, et al. Vancomycin-resistant enterococci colonization among dialysis patients: a meta-analysis of prevalence, risk factors, and significance. Am J Kidney Dis. 2015;65(1):88–97.

186. D’Agata EM, et al. Vancomycin-resistant enterococci among chronic hemodialysis patients: a prospective study of acquisition. Clin Infect Dis. 2001;32(1):23–29.

187. Dopirak M, et al. Surveillance of hemodialysis-associated primary bloodstream infections: the experience of ten hospital-based centers. Infect Control Hosp Epidemiol. 2002;23(12):721–724.

188. Taylor G, et al. Prospective surveillance for primary bloodstream infections occurring in Canadian hemodialysis units. Infect Control Hosp Epidemiol. 2002;23(12):716–720.

189. McDonald LC, Hageman JC. Vancomycin intermediate and resistant Staphylococcus aureus. What the nephrologist needs to know. Nephrol News Issues. 2004;18(11):63–64, 66–7, 71–2 passim.

190. Fridkin SK. Vancomycin-intermediate and -resistant Staphylococcus aureus: what the infectious disease specialist needs to know. Clin Infect Dis. 2001;32(1):108–115.

191. Chang S, et al. Infection with vancomycin-resistant Staphylococcus aureus containing the vanA resistance gene. N Engl J Med. 2003;348(14):1342–1347.

192. Sievert DM, et al. Vancomycin-resistant Staphylococcus aureus in the United States, 2002-2006. Clin Infect Dis. 2008;46(5):668–674.

193. Finks J, et al. Vancomycin-resistant Staphylococcus aureus, Michigan, USA, 2007. Emerg Infect Dis. 2009;15(6):943–945.

194. Zhu W, et al. Vancomycin-resistant Staphylococcus aureus isolates associated with Inc18-like vanA plasmids in Michigan. Antimicrob Agents Chemother. 2008;52(2):452–457.

195. Walters MS, et al. Vancomycin-resistant staphylococci aureus—Delaware, 2015. MMWR Morb Mortal Wkly Rep. 2015;64(37):1056.

196. Walters M, et al. Investigation and Control of Vancomycin-resistant Staphylococcus aureus: A Guide for Health Departments and Infection Control Personnel, 2015. Available from https://www.cdc.gov/hai/pdfs/vrsa-investigation-guide-05_12_2015.pdf.

197. Centers for Disease Control and Prevention. Invasive methicillin-resistant Staphylococcus aureus infections among dialysis patients—United States, 2005. MMWR Morb Mortal Wkly Rep. 2007;56(9):197–199.

198. Nguyen DB, et al. Invasive methicillin-resistant Staphylococcus aureus infections among patients on chronic dialysis in the United States, 2005-2011. Clin Infect Dis. 2013;57(10):1393–1400.

199. Centers for Disease Control and Prevention. Active Bacterial Core Surveillance Report, Emerging Infections Program Network, Methicillin-Resistant Staphylococcus aureus, 2015; 2015.
REFERENCES

200. Fluck R, et al. The UK Vascular Access Survey—follow-up data and repeat survey (chapter 5). Nephrol Dial Transplant. 2007;22(suppl 7): vii, 51–7.

201. Phatharacharukul P, et al. The risks of incident and recurrent clostridium difficile-associated diarrhea in chronic kidney disease and end-stage kidney disease patients: a systematic review and meta-analysis. Dig Dis Sci. 2015;60(10):2913–2922.

202. Tirath A, et al. Clostridium difficile infection in dialysis patients. J Investig Med. 2017;65(2):353–357.

203. Sheth H, et al. Clostridium difficile infections in outpatient dialysis cohort. Infect Control Hosp Epidemiol. 2010;31(1):89–91.

204. Evans R, et al. UK renal registry 18th annual report: chapter 12 epidemiology of reported infections amongst patients receiving dialysis for established renal failure in england 2013 to 2014: a joint report from public health england and the UK renal registry. Nephron. 2016;132(suppl 1):279–288.

205. See I, et al. Outbreak of clostridium difficile infections at an outpatient hemodialysis facility-michigan, 2012–2013. Infect Control Hosp Epidemiol. 2015;36(8):972–974.

206. Pop-Vicas A, et al. Multidrug-resistant gram-negative bacteria among patients who require chronic hemodialysis. Clin J Am Soc Nephrol. 2008;3(3):752–758.

207. Snyder GM, et al. Antimicrobial use in outpatient hemodialysis units. Infect Control Hosp Epidemiol. 2013;34(4):349–357.

208. D’Agata EM. Antimicrobial use and stewardship programs among dialysis centers. Semin Dial. 2013;26(4):457–464.

209. Infectious Diseases Society of America, et al. Combating antimicrobial resistance: policy recommendations to save lives. Clin Infect Dis. 2011;52(suppl 5):S397–428.

210. Bendinelli M, et al. Blood-borne hepatitis viruses: hepatitis B, C, D, and G viruses and TT virus. In: Specter RLH, Young SA, eds. Clinical virology manual. 3rd ed. Washington, DC: ASM Press; 2000:306–337.

211. Alter H. Discovery of non-A, non-B hepatitis and identification of its etiology. Am J Med. 1999;107(6B):16S–20S.

212. Choo QL, et al. Isolation of a cDNA clone derived from a blood-borne non-A, non-B viral hepatitis genome. Science. 1989;244(4902):359–362.

213. Choo QL, et al. Identification of the major, parenteral non-A, non-B hepatitis agent (hepatitis C virus) using a recombinant cDNA approach. Semin Liver Dis. 1992;12(3):279–288.

214. Centers for Disease Control and Prevention. Recommendations for prevention and control of hepatitis C virus (HCV) infection and HCV-related chronic disease. MMWR Recomm Rep. 1998;47(RR19):1–39.

215. Moyer LA, Alter MJ. Hepatitis C virus in the hemodialysis setting: a review with recommendations for control. Semin Dial. 1994;7:124–127.

216. Favero MS, Alter MJ. The reemergence of hepatitis B virus infection in hemodialysis centers. Semin Dial. 1996;9:373–374.

217. Niu MT, Coleman PJ, Alter MJ. Multicenter study of hepatitis C virus infection in chronic hemodialysis patients and hemodialysis center staff members. Am J Kidney Dis. 1993;22(4):568–573.

218. Fabrizi F, et al. Incidence of seroconversion for hepatitis C virus in chronic hemodialysis patients: a prospective study. Nephrol Dial Transplant. 1994;9(11):1611–1615.

219. Fabrizi F, et al. Hepatitis C virus infection and diabetes mellitus in end-stage renal disease: evidence of a negative association. Int J Artif Organs. 2006;29(7):691–697.

220. dos Santos JP, et al. Impact of dialysis room and reuse strategies on the incidence of hepatitis C virus infection in haemodialysis units. Nephrol Dial Transplant. 1996;11(10):2017–2022.

221. Forns X, et al. Incidence and risk factors of hepatitis C virus infection in a haemodialysis unit. Nephrol Dial Transplant. 1997;12(4):736–740.

222. McLaughlin KJ, et al. Nosocomial transmission of hepatitis C virus within a British dialysis centre. Nephrol Dial Transplant. 1997;12(2):304–309.

223. Petrosillo N, et al. Prevalence of infected patients and underestimating have a role in hepatitis C virus transmission in dialysis. Am J Kidney Dis. 2001;37(5):1004–1010.

224. Bergman S, et al. Hepatitis C infection is acquired pre-ESRD. Am J Kidney Dis. 2005;45(4):684–689.

225. Rahnavardi M, Hosseini Moghaddam SM, Alavian SM. Hepatitis C in hemodialysis patients: current global magnitude, natural history, diagnostic difficulties, and preventive measures. Am J Nephrol. 2008;28(4):628–640.

226. Saxena AK, Panhotra BR, Sundaram DS. The role the type of vascular access plays in the transmission of hepatitis C virus in a high prevalence hemodialysis unit. J Vasc Access. 2002;3(4):158–163.

227. Yakaryilmaz F, et al. Prevalence of occult hepatitis B and hepatitis C virus infections in Turkish hemodialysis patients. Ren Fail. 2006;28(8):729–735.

228. El-Reshad K, et al. Hepatitis C virus infection in patients on maintenance dialysis in kuwait: epidemiological profile and efficacy of prophylaxis. Saudi J Kidney Dis Transpl. 1995;6(2):144–150.

229. Gohar SA, et al. Prevalence of antibodies to hepatitis C virus in hemodialysis patients and renal transplant recipients. J Egypt Public Health Assoc. 1995;70(5-6):465–484.

230. Kamili S, et al. Infectivity of hepatitis C virus in plasma after drying and storing at room temperature. Infect Control Hosp Epidemiol. 2007;28(5):519–524.

231. Paintsil E, et al. Hepatitis C virus maintains infectivity for weeks after drying on inanimate surfaces at room temperature: implications for risks of transmission. J Infect Dis. 2014;209(8):1205–1211.

232. Jeffers LJ, et al. Hepatitis C infection in two urban hemodialysis units. Kidney Int. 1990;38(2):320–322.

233. Carrera F, et al. Prevalence of non-A non-B hepatitis and anti-HCV antibodies in a Portuguese dialysis population. Nephrol Dial Transplant. 1992;7(9):913–916.

234. Manescalchi F, et al. Anti-hepatitis C virus epidemiological study in two dialysis centers in Florence. Nephron. 1992;61(3):342–343.

235. Schneeberger PM, et al. The prevalence and incidence of hepatitis C virus infections among dialysis patients in the Netherlands: a nationwide prospective study. J Infect Dis. 2000;182(5):1291–1299.

236. Sivapalasingam S, et al. High prevalence of hepatitis C infection among patients receiving hemodialysis at an urban dialysis center. Infect Control Hosp Epidemiol. 2002;23(6):319–324.

237. Dussol B, et al. Hepatitis C virus infection among chronic dialysis patients in the south of France: a collaborative study. Am J Kidney Dis. 1995;25(3):399–404.

238. Selgas R, et al. Prevalence of hepatitis C antibodies (HCV) in a dialysis population at one center. Perit Dial Int. 1992;12(1):28–30.
REFERENCES

239. Thompson ND, et al. Nonhospital health care-associated hepatitis B and C virus transmission: United States, 1998-2008. Ann Intern Med. 2009;150(1):33–39.

240. Shimokura G, et al. Patient-care practices associated with an increased prevalence of hepatitis C virus infection among chronic hemodialysis patients. Infect Control Hosp Epidemiol. 2011;32(5):415–424.

241. Thompson ND, et al. Hepatitis C virus transmission in the hemodialysis setting: importance of infection control practices and aseptic technique. Infect Control Hosp Epidemiol. 2009;30(9):900–903.

242. Halleck R, et al. Hepatitis C transmission at an outpatient hemodialysis unit—New York 2001-2008. MMWR Morb Mortal Wkly Rep. 2009;58(8):189–194.

243. Collier MG, et al. Detection, reporting, and treatment of hepatitis C infections among hemodialysis patients. Infect Control Hosp Epidemiol. 2017;38(4):493–494.

244. Rao A, et al. Outbreak of hepatitis C virus infections at an outpatient hemodialysis facility: the importance of infection control competencies. Nephrol Nurs J. 2013;40(2):101–110. 164; quiz 111.

245. Alter MJ. The epidemiology of acute and chronic hepatitis C. Clinics in Liver Disease. 1997;1:559–568.

246. Alter MJ. Prevention of spread of hepatitis C. Hepatology. 2002;26(5 suppl 1):S93–S98.

247. Roth D, et al. Grazoprevir plus elbasvir in treatment-naive and treatment-experienced patients with hepatitis C virus genotype 1 infection and stage 4–5 chronic kidney disease (the C-SURFER study): a combination phase 3 study. Lancet. 2015;386(10003):1537–1545.

248. Fabrizi F, et al. Hepatitis C in chronic kidney disease, dialysis, and transplant. Kidney Int. 2016;89(5):988–994.

249. Kamili S, et al. Laboratory diagnostics for hepatitis C virus infection. Clin Infect Dis. 2012;55(suppl 1):S43–S48.

250. Centers for Disease Control and Prevention. Testing for HCV infection: an update of guidance for clinicians and laboratorians. MMWR Morb Mortal Wkly Rep. 2013;62(18):362–365.

251. Mbaye C, Thompson ND. Hepatitis C virus screening and management of seroconversions in hemodialysis facilities. Semin Dial. 2013;26(4):439–446.

252. Kidney Disease. Improving Global Outcomes, KDIGO clinical practice guidelines for the prevention, diagnosis, evaluation, and treatment of hepatitis C in chronic kidney disease. Kidney Int Suppl. 2008;(109):S1–99.

253. Centers for Disease Control and Prevention. In: Welfare HEa, ed. Control Measures for hepatitis B in dialysis centers. Atlanta, GA: Department of Health, Education, and Welfare; 1977.

254. Alter MJ, Favero MS, Francis DP. Cost benefit of vaccination for hepatitis B in hemodialysis centers. J Infect Dis. 1983;148(4):770–771.

255. Alter MJ, Favero MS, Maynard JE. Impact of infection control strategies on the incidence of dialysis-associated hepatitis in the United States. J Infect Dis. 1986;153(6):1149–1151.

256. Centers for Disease Control and Prevention. Recommendations of the Immunization Practices Advisory Committee (ACIP): Inactivated hepatitis B vaccine. MMWR Morb Mortal Wkly Rep. 1982;31(24):317–322, 327–328.

257. Alter HJ, et al. Type B hepatitis: the infectivity of blood positive for e antigen and DNA polymerase after accidental needlestick exposure. N Engl J Med. 1976;295(17):909–913.

258. Shikata T, et al. Hepatitis B antigen and infectivity of hepatitis B virus. J Infect Dis. 1977;136(4):571–576.

259. Bond WW, et al. Survival of hepatitis B virus after drying and storage for one week. Lancet. 1981;1(8219):550–551.

260. Favero MS, et al. Detection methods for study of the stability of hepatitis B antigen on surfaces. J Infect Dis. 1974;129(2):210–212.

261. Favero MS, et al. Letter: Hepatitis-B antigen on environmental surfaces. Lancet. 1973;2(7843):1455.

262. Snyderman DR, et al. Transmission of hepatitis B associated with hemodialysis: role of malfunction (blood leaks) in dialysis machines. J Infect Dis. 1976;134(6):562–570.

263. Snyderman DR, et al. Hemodialysis-associated hepatitis: report of an epidemic with further evidence on mechanisms of transmission. Am J Epidemiol. 1976;104(5):563–570.

264. Niu MT, et al. Hemodialysis-associated hepatitis B: report of an outbreak. Dial Transplant. 1989;18:542–555.

265. Centers for Disease Control and Prevention. Outbreaks of hepatitis B virus infection among hemodialysis patients—California, Nebraska, and Texas, 1994. MMWR Morb Mortal Wkly Rep. 1996;45(14):285–289.

266. Alter MJ, Ahtone J, Maynard JE. Hepatitis B virus transmission associated with a multiple-dose vial in a hemodialysis unit. Ann Intern Med. 1983;99(3):330–333.

267. Carl M, Francis DP, Maynard JE. A common-source outbreak of hepatitis B in a hemodialysis unit. Dial Transplant. 1983;12:222–229.

268. Hutm YJ, et al. An outbreak of hospital-acquired hepatitis B virus infection among patients receiving chronic hemodialysis. Infect Control Hosp Epidemiol. 1999;20(11):731–735.

269. Public Health Laboratory Service Survey. Decrease in the incidence of hepatitis in dialysis units associated with prevention programme. Br Med J. 1974;4(5947):751–754.

270. Najem GR, et al. Control of hepatitis B infection. The role of surveillance and an isolation hemodialysis center. JAMA. 1981;245(2):153–157.

271. Rhea S, et al. Hepatitis B reverse seroconversion and transmission in a hemodialysis center: a public health investigation and case report. Am J Kidney Dis. 2016;68(2):292–295.

272. Rangel MC, et al. Vaccine recommendations for patients on chronic dialysis. The Advisory Committee on Immunization Practices and the American Academy of Pediatrics. Semin Dial. 2000;13(2):101–107.

273. Mast EE, et al. A comprehensive immunization strategy to eliminate transmission of hepatitis B virus infection in the United States: recommendations of the Advisory Committee on Immunization Practices (ACIP) Part II: immunization of adults. MMWR Recomm Rep. 2006;55(RR-16):1–33. quiz CE1–4.

274. Mast EE, et al. A comprehensive immunization strategy to eliminate transmission of hepatitis B virus infection in the United States: recommendations of the Advisory Committee on Immunization Practices (ACIP) part I: immunization of infants, children, and adolescents. MMWR Recomm Rep. 2005;54(RR-16):1–31.

275. Hoofnagle JH, Di Bisceglie AM. Serologic diagnosis of a acute and chronic hepatitis. Sem Liver Dis. 1991;11(2):73–83.

276. Hochberger S, et al. Fully automated quantitation of hepatitis B virus (HBV) DNA in human plasma by the COBAS AmpliPrep/COBAS TaqMan system. J Clin Virol. 2006;35(4):373–380.
REFERENCES

277. Kaneko S, et al. Hepatitis B virus DNA detection and comparison with hepatitis B surface antigen. Gastroenterol Jpn. 1990;25(supp1 2):57–61.

278. Dienstag JL, et al. A preliminary trial of lamivudine for chronic hepatitis B infection. N Engl J Med. 1995;333(25):1657–1661.

279. Bourne EJ, et al. Quantitative analysis of HBV cccDNA from clinical specimens: correlation with clinical and virological response during antiviral therapy. J Viral Hepat. 2007;14(1):55–63.

280. Dienstag JL, et al. Extended lamivudine retreatment for chronic hepatitis B: maintenance of viral suppression after discontinuation of therapy. Hepatology. 1999;30(4):1082–1087.

281. Dienstag JL, et al. Lamivudine as initial treatment for chronic hepatitis B in the United States. N Engl J Med. 1999;341(17):1256–1263.

282. Lai CL, et al. A one-year trial of lamivudine for chronic hepatitis B. Asia Hepatitis Lamivudine Study Group. N Engl J Med. 1998;339(2):61–68.

283. Hadler SC, et al. Epidemiological analysis of the significance of low-positive test results for antibody to hepatitis B surface and core antigens. J Clin Microbiol. 1984;19(4):521–525.

284. Mezzelani P, et al. [The significance of the isolated anti-HBc carrier. A study of 1797 drug addicts. The Intersert Group of Scientific Collaboration]. Recent Prog Med. 1994;85(9):419–424.

285. Levine OS, et al. Seroepidemiology of hepatitis B virus in a population of injecting drug users. Association with drug injection patterns. Am J Epidemiol. 1995;142(3):331–341.

286. Satake M, et al. Infectivity of blood components with low hepatitis B virus DNA levels identified in a lookback program. Transfusion. 2007;47(7):1197–1205.

287. Bhatti FA, et al. Anti-hepatitis B core antigen testing, viral markers, and occult hepatitis B virus infection in Pakistani blood donors: implications for transfusion practice. Transfusion. 2007;47(1):74–79.

288. Silva AE, et al. Hepatitis B virus DNA in persons with isolated antibody to hepatitis B core antigen who subsequently received hepatitis B vaccine. Clin Infect Dis. 1998;26(4):895–897.

289. McMahon BJ, et al. Response to hepatitis B vaccine of persons positive for antibody to hepatitis B core antigen. Gastroenterology. 1992;103(2):590–594.

290. Lai CL, et al. Significance of isolated anti-HBc seropositivity by ELISA: implications and the role of radioimmunoassay. J Med Virol. 1992;36(3):180–183.

291. Hadler, S.C. and H.A. Fields, eds. Hepatitis delta virus. Textbook of Human Virology, ed. B.B. Belshe. 1991, Mosby: St. Louis. MO. 749–765.

292. Lettou LA, et al. Nosocomial transmission of delta hepatitis. Ann Intern Med. 1986;104(5):631–635.

293. Velandia M, et al. Transmission of HIV in dialysis centre. Lancet. 1995;345(8962):1417–1422.

294. El Sayed NM, et al. Epidemic transmission of human immunodeficiency virus in renal dialysis centers in Egypt. J Infect Dis. 2000;181(1):91–97.

295. Arduino MJ, et al. Preventing health-care associated transmission of bloodborne pathogens in hemodialysis facilities. Seminars in Infect Control. 2001;1(1):49–60.

296. Favero MS. Transmission of HIV in dialysis units. Anna J. 1993;20(5):599–600.

297. Dyer E. Argentinian doctors accused of spreading AIDS. BMJ. 1993;307(6904):584.

298. Mashragi F, et al. HIV transmission at a Saudi Arabia hemodialysis unit. Clin Infect Dis. 2014;59(6):897–902.

299. Team WHOER, et al. Ebola virus disease in West Africa—the first 9 months of the epidemic and forward projections. N Engl J Med. 2014;371(16):1481–1495.

300. Vallabhaneni S, et al. Investigation of the first seven reported cases of candida auris, a globally emerging invasive, multi-drug-resistant fungus—United States, May 2013-August 2016. MMWR Mortal Wkly Rep. 2016;65(44):1234–1237.

301. Hess S, Bren V. Essential components of an infection prevention program for outpatient hemodialysis centers. Semin Dial. 2013;26(4):384–398.

302. Centers for Disease Control and Prevention. Pyrogenic reactions and gram-negative bacteremia in a hemodialysis center, in Epidemic Investigation Report. In: C.f.D.C.a.P. (Centers for Disease Control and Prevention), ed. Atlanta, GA: Author; 1991.

303. Centers for Disease Control and Prevention. Bacteremia in hemodialysis patients. In: Epidemiologic Investigations Report, CDC. Atlanta, GA: Centers for Disease Control and Prevention; 1992.

304. Clark T, et al. Outbreak of bloodstream infection with the mold Phialemonium among patients receiving dialysis at a hemodialysis unit. Infect Control Hosp Epidemiol. 2006;27(11):1164–1170.

305. Centers for Disease Control and Prevention. Bacteremia associated with reuse of disposable hollow-fiber hemodialyzers. MMWR Mortb Mortal Wkly Rep. 1986;35:417–418.

306. Murphy J, et al. Outbreaks of bacteremia in hemodialysis patients associated with alteration of dialyzer membranes [abstract]. Asaio J. 1987;16:51.

307. Centers for Disease Control and Prevention. Pyrogenic reactions in patients undergoing high-flux hemodialysis—California, in Epidemic Investigations Report. In: C.f.D.C.a.P. (Centers for Disease Control and Prevention), ed. Atlanta, GA: Author; 1987.

308. Centers for Disease Control and Prevention. Pyrogenic reactions in hemodialysis patients on high-flux hemodialysis—California, in Epidemic Investigations Report. In: C.f.D.C.a.P. (Centers for Disease Control and Prevention), ed. Atlanta, GA: Author; 1987.

309. Kantor RJ, et al. Outbreak of hepatitis B in a dialysis unit, complicated by false positive HBs test results. Dial Transplant. 1979;8:232–235.

310. Centers for Disease Control and Prevention. Outbreak of hepatitis B in a dialysis center. Epidemic Investigation Report EPI91-17. Atlanta, CDC, 1993, in Epidemic Investigations Report. Atlanta, GA: CDC; 1993.

311. Centers for Disease Control and Prevention. Non-A, Non-B hepatitis in a dialysis center, Nashville, Tennessee, in Epidemic Investigation Report. In: C.f.D.C.a.P. (Centers for Disease Control and Prevention), ed. Atlanta, GA: Author; 1979.

312. Niu MT, et al. Outbreak of hemodialysis-associated non-A, non-B hepatitis and correlation with antibody to hepatitis C virus. Am J Kidney Dis. 1992;19(4):345–352.

313. Lake J, et al. Improving Infection Control in Hemodialysis Centers After Acute Hepatitis C Infections, in National Kidney Foundation Spring Clinical Meeting; 2016. Boston, MA.