METHODS FOR ORGANIZING THE DISINFECTION OF ONE OR MORE ITEMS CONTAMINATED WITH BIOLOGICAL AGENTS

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RFID Reader in Hallway's RFID Tag Responding Writing Signal
RFID Tagged Equipment Mobile Displaying RFID Reader
Slevage units that can attach to any WV or H2O2 System
RFID Reader in the Storage rooms and Choke points
UV Disinfection Device
Managerrent System Servers

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Field of Classification Search
None
See application file for complete search history.

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ABSTRACT
The invention provides method for organizing the disinfection of one or more items contaminated with biological agent(s) comprising (a) attaching a radio-frequency ID (RFID) tag to an item(s) to be disinfected; exposing the item(s) of (a) to a disinfecting means for a period sufficient to disinfect the item; and (c) obtaining a signal from the tagged item when disinfection is complete thereby organizing the disinfection of one or more items.

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RFID Tag Responding / Emitting Signal

RFID Tagged Equipment

RFID Reader in the Storage rooms and Choke points

Mobile Display / RFID reader / Storage units that can attach to any UV or H2O2 System

RFID Reader in Hallways

UV Disinfection Device

Management System

Servers

FIG. 1
TABLETS DISPLAY SCREEN

RFID TAG
RFID CONTROLLER
MEMORY
SENSOR SIGNAL PROCESSOR

RFID TAG 1
CIRCUIT BOARD
RFID TAG 2
RFID TAG n

RFID READER/INTERROGATOR
MICRO PROCESSOR

WI-FI TRANSMITTER/RECEIVER

RFID ASSET TRACKING DATA
SERVER/ HOUSE SERVICES

FIG. 9
FIG. 10
Operator Notified

Discharged Patient Room \( \rightarrow \) RME in Equipment Room

Operator runs Pulsed UV Device in Position #1

UV radiation sensed dose > 50 millijoules/cm² OR > 5 minutes

RFID on wall < 4 feet

RFID on RME

Operator runs Pulsed UV Device in Positions #2, #3, ...

* Once all required positions for a room are complete, Bed Management system display screen will display the room changing status as "Complete. > Ready to next occupant"
METHODS FOR ORGANIZING THE DISINFECTION OF ONE OR MORE ITEMS CONTAMINATED WITH BIOLOGICAL AGENTS

This patent application is a 371 application of PCT application No. PCT/US2013/053307, filed Aug. 2, 2013, claiming the benefit of the filing date of U.S. Ser. No. 61/678,558, filed Aug. 1, 2012, the contents of all of which are herein incorporated by reference in their entireties into the present patent application.

Throughout this application various publications are referenced. The disclosures of these publications in their entireties are hereby incorporated by reference into this application in order to more fully describe the state of the art to which this invention pertains.

BACKGROUND OF THE INVENTION

Overview of Hospital Associated Infections

Hospital acquired or associated infections (HAI) are an important cause of mortality and morbidity affecting an estimated 1.7 million patients and causing 100,000 deaths annually in the United States. As multiple drug resistant organisms (MDROs) represent an increasing challenge to successfully treat, they also significantly contribute to increasing health care costs, not to mention unnecessary patient burden. Evidence suggests a 28-58% higher risk of infection is attributable to surface contamination and cross transmission, especially for Methicillin resistant Staphylococcus aureus (MRSA) and Clostridium difficile (C.diff). Additionally, an estimated 20-40% of all HAI result from cross contamination via health care personnel either by direct patient contact or by touching contaminated environmental surfaces in the room to include reusable medical equipment's like IV poles, blood pressure cuffs, and monitors. In addition to MRSA infections and C.diff, the role of environmental contamination for transmission has been studied in several other important infectious microbes including vancomycin resistant enterococcus (VRE), and Acinetobacter baumannii. Contaminated surfaces contribute up to half of the risk for acquiring infections, as these organisms persist on environmental surfaces for many days. In the intensive care unit (ICU) setting, even improvements in cleaning visibly soiled surfaces can lead to reducing the rate of MRSA and VRE by 30-50%. However, this requires an evaluation and feedback system that is quite labor intensive, and the sustainability of such costly efforts would undoubtedly be limited. In fact, several studies show that manual cleaning performed after patient rooms are vacated is unreliable, with residual contamination rates ranging up to 50%. This residual environmental contamination with potentially resistant organisms endangers the health of the next room occupant, not to mention staff, and risks cross transmission throughout the hospital via hand transmission from health care workers as well as with reusable medical equipment. Even after appropriate room cleaning, patients subsequently admitted to rooms where MRSA, C.diff or VRE patients were previously isolated remain at increased risk from acquisition, due to environmental contamination. This has been well demonstrated in many prior studies where admission to a room previously occupied by an HAI-positive patient was associated with a 28-58% increased risk of acquisition of these pathogens.

There have been multiple studies in various countries that have documented lack of compliance with established guidelines for disinfection leading to numerous outbreaks.

How Current Cleaning Practices are Inadequate

Many HAI reduction initiatives, such as raising compliance of hand hygiene, antimicrobial stewardship programs and isolation/screening practices, have become standard practice. The environmental cleaning of patient rooms, however, has not evolved significantly in response to other HAI reduction efforts. Recent studies by Carling et al., demonstrate conclusively that manual cleaning is inadequate for effectively reducing the bio-burden in patient care areas because as many as 70% of high-touch surfaces (e.g., bed rails, call buttons, television remote controls) are missed during both standard discharge and isolation cleanings. This could also be applied to other reusable medical equipment that is present in the same patient room. This incomplete cleaning allows for organisms to remain on the room or equipment following patient discharge, placing the subsequent patient at a higher risk as organisms like MRSA, VRE and C.diff spores can survive in the environment from 3-12 months.

The novel methods of the invention were devised to alleviate the aforementioned problems and should radically and systematically reduce the contamination on equipment or supplies in in-patient rooms.

SUMMARY OF THE INVENTION

Various embodiments described herein are directed to methods for organizing (manually or through automated means) the disinfection of one or more items contaminated with biological agent(s) comprising attaching a radio-frequency ID (RFID) tag to an item(s) to be disinfected, exposing the item(s) to a disinfecting means for a period sufficient to disinfect the item, and obtaining a signal from the tagged item when disinfection is complete thereby organizing the disinfection of one or more items.

The invention also provides methods for disinfection (manually or through automated means) of one or more items contaminated with biological agent(s) comprising exposing the item(s) having RFID tags to a disinfecting means for a period sufficient to disinfect the item and obtaining a signal from the tagged item(s) when disinfection is complete thereby organizing the process of disinfection of one or more items.

The invention also provides methods for detecting (manually or through automated means) whether an item exiting or leaving a designated space is contaminated comprising attaching a radio-frequency ID (RFID) tag to the item(s) to be disinfected, attaching an RFID to a chokepoint, and obtaining a signal from the tagged item indicating when a contaminated item crosses the chokepoint.

The invention further provides methods for detecting (manually or through automated means) whether an item exiting a designated space is clean or disinfected comprising attaching a radio-frequency ID (RFID) tag to the item(s) to be disinfected, attaching an RFID to a chokepoint, and obtaining a signal from the tagged item indicating when a clean or disinfected item crosses the chokepoint.

The invention also provides methods for organizing (manually or through automated means) the disinfection of a designated space contaminated with biological agent(s) comprising attaching one or more radio-frequency ID (RFID) tags to a designated area to be disinfected, exposing the designated area or portion thereof to a disinfecting means for a period sufficient to disinfect the designated area,
and obtaining a signal from the tagged designated area when disinfection is complete thereby organizing the disinfection of the designated space.

The invention further provides methods for disinfection (manually or through automated means) of a designated space contaminated with biological agent(s) comprising exposing the designated space or portion thereof to a disinfecting means for a period sufficient to disinfect the designated space or portion thereof, said designated space or portion thereof having one or more radio-frequency ID (RFID) tags and obtaining a signal from the tagged designated space or portion thereof when disinfection is complete thereby organizing the disinfection of the designated space or portion thereof.

Additionally provided are apparatus and systems for use in the methods of this invention.

BRIEF DESCRIPTION OF THE FIGURES

FIG. 1 is a diagram showing the process for disinfection of reusable medical equipment.

FIG. 2 is a photograph of a UV Aluminum reflector.

FIG. 3 shows photographs of other Mercury-based UV disinfection machines.

FIG. 4 shows photographs of the different Hydrogen Peroxide Systems.

FIG. 5 shows drawings of different passive RFID tags.

FIG. 6 shows drawings of different RFID readers.

FIG. 7 shows graphs of threshold sensing values over time.

FIG. 8 shows drawing of a pulsed xenon UV device.

FIG. 9 shows an overall flow chart of the inner workings of the Jangama with optional memory feature.

FIG. 10 shows a drawing of an IV pole tagged with active RFID tags, which depict clean versus dirty equipment status. The drawing on the left is an active RFID tag with e-ink technology; on the right, is an active RFID tag with LED display technology.

FIG. 11 shows drawing of Tru-D, a mercury-based UV device.

FIG. 12 shows an overall flow chart demonstrating the workings of the Jangama with an UV-based cleaning (also referred to herein as disinfecting) system.

DETAILED DESCRIPTION OF THE INVENTION

Definitions

Healthcare associated/acquired infections (HAI's) or nosocomial infections are infections that may be acquired by a patient during the course of his/her hospital admission.

In accord with the practice of the invention, the disinfecting means may kill or inhibit biological contaminants including, but not limited to, micro-organisms, viral particles, fungal particles, spores to include Clostridium difficile, tuberculous particles, other infectious agents, microbes in human secretion, human excretion, dust and fecal material.

Patient room refers to a room occupied by a patient during his/her hospital stay.

Clean/Dirty utility or equipment room refers to a room used to store re-usable medical equipment such as IV poles, tray tables, wheelchairs, thermometer, console, computer on wheels (COW), blood pressure cuff/device, bedside commode, intravenous/infusion pump and pole (IV pole), SCD devices, cooling blankets, warming blankets (bear huggers), portable x-ray machines, equipment carts, monitors, various high dollar machines like Electrocencephalograms (EEG), Electrocardiogram (EKG) machines, wheelchairs, walkers or crutches.

UV light refers to ultraviolet light delivered by, for example, the pulsed xenon lamp or mercury based technologies or equivalents at the appropriate frequency. The Mercury based technologies like Tri-D (Lumalier, Tenn.) (FIG. 11) or V-360+ Room Sanitizer (UVDL, CA) uses a mercury lamp to produce UV light in the 254 nm range which is bactericidal. Pulsed xenon-UV germicidal irradiation produces a more broad-spectrum UV irradiation, but also produces large amounts of energy in the germicidal spectrum (200-320 nm) by using a xenon gas flash lamp (Xenex Healthcare Service, TX).

Hydrogen Peroxide refers to hydrogen peroxide dry mist, hydrogen peroxide vapor, vaporized Hydrogen peroxide, hydrogen peroxide fog or steam or spray or equivalent.

RFID refers to Radio-frequency Identification which is a wireless use of radio-frequency electromagnetic fields to transfer data for the purposes of identifying and/or tracking objects. There is no physical contact between communicating objects.

RFID tag[s] refers to an object that can be applied, imbedded or otherwise attached onto a product, e.g., re-usable medical equipment or room entrance for the purposes of tracking the disinfection of equipment or patient rooms or utility storage rooms. These tags can be either active or passive or a hybrid thereof.

RFID reader[s] also referred to as RFID interrogator refers to an object that contains a radio-frequency module, which transmits and/or receives radio-frequency signals. The reader’s function is to interact with RFID tag[s]. These readers can be either active or passive. They can also be fixed or mobile. Signals from the reader are further transmitted to either the mobile unit (also referred to here as Jangama) microprocessor or to the central data warehouse directly.

RFID system refers to the type of tag and reader used to track and interrogate objects. A Passive Reader Active Tag (PRAT) system includes a passive reader and a battery operated active tag wherein the reader only receives the signal from the active tags. An Active Reader Passive Tag (ARPT) system includes an active reader which transmits signals to the passive tags and also, optionally, receives signals from the passive tags. An Active Reader Active Tag (ARAT) system includes active tags that can respond to the interrogation from an active reader.

Fixed RFID readers used in this invention will be used near the entrance of re-usable medical equipment storage rooms or actual patient rooms which will allow for a highly defined reading area for when the tagged re-usable medical equipment go in and out of the storage area or patient room. Such readers could also function as choke point readers.

Mobile readers (which may be housed in a mobile unit, e.g., Jangama) in this invention may be used with the no-touch disinfection devices to alert environmental services staff of patient rooms or equipment in need of cleaning (also referred to herein as disinfecting) within an interrogation zone.

Cleaning includes terminally disinfecting a patient room after a patient has been discharged to keep it ready for the next patient. This refers to thoroughly disinfecting the patient bed; including the top, front, sides, headboard, side rails, mattress, between side rails; the nurse call light; the TV remote; all high touch surfaces such as tray tables, bedside tables and drawers, phone, armchairs, door handles, light switches, closet handles, etc.; disinfecting the bathroom to include the fixtures, support bars and other surfaces. Privacy
curtains may be removed and bagged for laundry. Window curtains, ceilings and walls also may need to be disinfected. In cases where the occupant had, e.g., MRSA, C-Diff or VRE, the room may be terminally disinfected from floor to ceiling with agents such as diluted bleach. Additionally, all re-usable medical equipment in the room may be thorougly cleaned or disinfected and returned to special processing department or the utility storage area. Cleaning may be generally done by the environmental services personnel in most hospitals and there is always a potential for missed opportunities, where a high touch area was not thoroughly cleaned per protocol. Such missed opportunities could result in HAI in patients or visitors. This is a tedious and time consuming process, thereby reducing the turnaround time for a patient room. The invention focuses on decreasing the room turnaround time by decreasing the time needed to terminally clean or disinfect the room and also electronically updating the availability of a room throughout the hospital system without relying on manual entry.

Re-usable medical equipment such as IV poles, wheelchairs, walkers, bedside commodes, etc. are used as needed. Between each patient use, this equipment may be stored in dirty equipment rooms and wheeled/transported to the special processing department where terminal cleaning of the equipment occurs per protocol. This also leads to a missed opportunity for cleaning, resulting in potential HAI. This process also decreases the availability of re-usable medical devices during high patient census weeks and in ERs where efficiency is the key to delivering patient care. To address the issues, the invention in one embodiment, focuses on RFID tagged equipment which are held in storage rooms and disinfected with UV rays. The equipment will feature active RFID tags which can communicate with respective readers about its location, clean/unclean status, etc. The active tags would also enable the equipment to be tracked by a software/application on the hospital network making it easy for a nurse in emergency room trying to locate a clean/unused IV pole which could be on the med-surgical unit.

No touch disinfection systems are means of cleaning enclosed areas. The disinfection can be achieved either by using UV disinfection systems (either mercury-based as in FIG. 3 or pulsed-xenon-based as in FIG. 8) or hydrogen peroxide systems (vapor, mist or aerosolized as on FIG. 4). In either scenario, after manual cleaning, the device is placed in a room (typically discharged empty patient room) and run for a specified amount of time (disinfection cycle). For some devices, like a pulsed-xenon-based device, the device may have 2-4 positions of disinfection.

A chokepoint refers to a point of entry or escape, where a reader may be installed in order to track objects or to warn of the exit or entry of unclean equipment. A Jangama is a mobile or integrated unit to be used for organizing disinfection of rooms and equipment (FIG. 9).

As used herein, the term “comprising” when placed before the recitation of steps in a method means that the method encompasses one or more steps that are additional to those expressly recited, and that the additional one or more steps may be performed before, between, and/or after the recited steps. For example, a method comprising steps a, b, and c encompasses a method of steps a, b, x, and c, a method of steps a, b, x, and c, as well as a method of steps x, a, b, and c. Furthermore, the term “comprising” when placed before the recitation of steps in a method does not (although it may) require sequential performance of the listed steps, unless the context clearly dictates otherwise. For example, a method comprising steps a, b, and c encompasses, for example, a method of performing steps in the order of steps a, c, and b, the order of steps c, b, and a, and the order of steps c, a, and b. Unless otherwise indicated, all numbers expressing quantities of ingredients, properties such as molecular weight, reaction conditions, and so forth as used herein, are to be understood as being modified in all instances by the term “about.” Accordingly, unless indicated to the contrary, the numerical parameters herein are approximations that may very depending upon the desired properties sought to be obtained by the present invention. At the very least, and without limiting the application of the doctrine of equivalents to the scope of the claims, each numerical parameter should at least be construed in light of the number of reported significant digits and by applying ordinary rounding techniques. Notwithstanding that the numerical ranges and parameters describing the broad scope of the invention are approximations, the numerical values in the specific examples are reported as precisely as possible. Any numerical value, however, inherently contains standard deviations that necessarily result from the errors found in the numerical value’s testing measurements.

In order that the invention herein described may be more fully understood, the following description is set forth:

The invention provides methods for organizing the process of disinfection of one or more items contaminated with biological agent(s). The method comprises attaching a radio-frequency ID (RFID) tag to an item(s) to be disinfected. The method further comprises exposing the item(s) of the a disinfecting means for a period sufficient to disinfect the item. Further, the method comprises obtaining a signal from the tagged item when disinfection is complete thereby organizing the disinfection of one or more items.

In one embodiment, the one or more items contaminated with biological agent(s) may be disinfected by the method of the invention.

The invention also provides methods for disinfection of one or more items contaminated with biological agent(s). The method comprises exposing the item(s) having RFID tags attached thereto or contained therein to a disinfecting means for a period sufficient to disinfect the item. The method further comprises obtaining a signal from the tagged item (e.g., via the RFID tag) when disinfection is complete.

In one embodiment, the method for organizing the disinfection of one or more items contaminated with biological agent(s) may be manual or an automated method. In another embodiment, the method for disinfection of one or more items contaminated with biological agent(s) may be manual or an automated method.

In one embodiment, the item to be disinfected is a device, machine or tool. The machine may be medical equipment, e.g., durable medical equipment or reusable medical equipment.

In an embodiment of the invention, the disinfecting means is a ultra-violet (UV) light system. In a further embodiment, the ultra-violet light is from pulsed xenon light. For example, disinfection systems are available that provide such pulsed light (Xenex Healthcare Services LLC) as shown in e.g. FIG. 8. In another example, the UV light may be from a mercury vapor UV lamp. In another example, the disinfection system may provide UV light such as Tru-D LLC’s SmartUVC room disinfection system as shown in e.g. FIG. 11. The Mercury based technologies like Tru-D (Lumilair, Tenn.) or V-360+ Room Sanitizer (UVDI, CA) uses a mercury lamp to produce UV light in the 254 nm range which is bactericidal. Pulsed Xenon-UV germicidal irradiation produces a more broad-spectrum UV irradiation, but also produces large amounts of energy in the germicidal
spectrum (200-320 nm) by using a xenon gas flash lamp (Xenex Healthcare Service, TX).19

In accordance with the practice of the invention, the space or room where the equipment will be cleaned may be coated with a reflective coating (paint)(UVC 360, Lumacpect, Tenn.). This coating enhances reflectivity of the UVC, thus improving the overall disinfection and thoroughness of disinfection.20 Additionally, the space or room to be cleaned may be lined with reflective aluminum panels with arching overheads as shown in e.g. FIG. 2. These panels enhance reflectivity of the UVC thus improving the overall disinfection and thoroughness of disinfection.

In a further embodiment, the disinfecting means may be a hydrogen peroxide system. Hydrogen peroxide systems may include hydrogen peroxide vapour (HPV), activated hydrogen peroxide, dry mist hydrogen peroxide or vaporized hydrogen peroxide.21-22 Examples of hydrogen peroxide systems include, but are not limited to, Bioquell Aeration Units (Bioquell, UK), VHI ARD Biodecontamination System (Steris Corporation, OH), and GLOSAIR™ Environmental Decontamination Systems (ASP Inc., CA) as shown in e.g. FIG. 4.

Examples of biological contaminants include, but are not limited to, micro-organisms, viral particles, fungal particles, spores to include Clostridium difficile, tuberculous particles, other infectious agents, microbes in human secretion, human excretion, dust and fecal material.

Further, the micro-organism may be a bacterium, fungus, protozoa, spore or virus. In another embodiment, the bacterium may be Clostridium difficile, Clostridium botulinum, or Clostridium perfringens and its spores. Additionally, the bacterium may be a drug resistant bacterium. The drug resistant bacterium may be a methicillin-resistant Staphylococcus aureus (MRSA). Also, in another example, the bacterium may be Staphylococcus epidermidis or Staphylococcus saprophyticus. Further example of the drug resistant bacterium may be a Carbapenem-resistant gram-negative bacterium, or the gram negative bacteria Hemophilus influenzae, Klebsiella pneumoniae, or Legionella pneumophila. Additional examples of microorganisms include viruses such as human rhinovirus, poliovirus, norovirus, coxsackievirus or echovirus.

In one embodiment of the invention, the RFID tag may communicate with the disinfection means. In another embodiment, the RFID tag may communicate the initiation or completion of the cleaning or disinfection cycle to a computer. In a further embodiment, the method comprises processing the signal from the tagged item to create a log of the cleaning or disinfecting history of each RFID tagged item through a computer. In yet another embodiment, the method comprises processing the signal from the item(s) having an RFID tag to create a log of the cleaning or disinfecting history of each RFID tagged item through a computer.

The invention also provides methods for detecting whether an item exiting a designated space is clean or disinfected. The method comprises attaching a radio-frequency ID (RFID) tag to the item(s) to be disinfected. The method further comprises attaching an RFID detector or reader to a checkpoint or transit point. Further, the method comprises obtaining a signal from the tagged item indicating when a clean or disinfected item crosses the checkpoint or transit point.

In one embodiment, the method for detecting whether an item exiting a designated space is clean or disinfected may be an automated method.

In accord with the practices of the invention, the method further comprises processing the signal from the tagged item to create a log of the RFID tagged item through a computer. In a further embodiment, the RFID tag permits the creation of a log of the cleaning or disinfecting history of each RFID tagged item through a computer. In another embodiment, the RFID tag may be connected to a checkpoint, and may provide notification when a piece of dirty equipment crosses the checkpoint. In another embodiment, the alert and notification system may be computer based, wherein a signal is sent to the data warehouse and appropriate personnel may be notified via cell phone, pager, or other means of communication, include any vibrating or flashing wireless device.

In another embodiment, the notification device may be located on the door or other checkpoint itself and may be audible or visual.

In one embodiment of the invention, suitable RFID tags may include passive RFIDs as shown in FIG. 5. In a further embodiment, active RFID tags, for example, as shown in FIG. 11 may be used (Supply Insight Co., Intermec Co., AeroScout or AiRISTA). The RFID with display could be of many varieties. In one embodiment, the active RFID tag may include one LED light or 2 LED lights or 2 LED lights of different colors (US Publication No. US 2010/0171586). In another embodiment, the active RFID tag may be used to provide an audible or visual warning.

The invention also provides methods for organizing (manually or through automated means) the disinfection of a designated area(s) contaminated with biological agent(s) comprising attaching one or more radio-frequency ID (RFID) tags to a designated area to be disinfected. The method further comprises exposing the designated area or portion thereof to a disinfecting means for a period sufficient to disinfect the designated area. Further, the method comprises obtaining a signal from the tagged designated area when disinfection is complete thereby organizing the disinfection of the designated space.

In one embodiment, the designated area(s) contaminated with biological agent(s) may be disinfected by the method of the invention.

The invention also provides methods for organizing (manually or through automated means) the process of disinfection of a designated area(s) contaminated with biological agent(s) comprising exposing the designated area or portion thereof having one or more RFID tags to a disinfecting means for a period sufficient to disinfect the designated area. The method further comprises obtaining a signal from the tagged designated area or portion thereof when disinfection is complete.
In one embodiment, the method for organizing the disinfection of a designated space contaminated with biological agent(s) may be an automated method.

In accordance with the practice of the invention, the space or room to be cleaned may be coated with a reflective coating of paint (UVC 360, Lumacept, Tenn.). This coating enhances reflectivity of the UVC thus improving the overall disinfection and thoroughness of disinfection.

Additionally, the designated space to be disinfected may be a room or portion thereof. The room may be enclosed or open. For example, the room may be a hospital room (for humans and veterinary animals). In yet another example, the room may be in a day care center or nursery or school.

In one embodiment, the disinfecting means may be an ultra-violet light or hydrogen peroxide system as mentioned previously.

In various embodiments, the micro-organism or biological contaminant may be as mentioned previously.

In various embodiments, the RFID tag may communicate with the disinfection means. In another embodiment, the RFID tag may communicate the initiation or completion of the cleaning or disinfection cycle to a computer or data warehouse. In a further embodiment, the method comprises processing the signal from the tagged designated area to create a log of the cleaning or disinfecting history of each RFID tagged item through a computer. In yet another embodiment, comprising processing the signal from the tagged designated space or portion thereof to create a log of the cleaning or disinfecting history of each RFID tagged item through a computer. In yet another embodiment, the RFID tag permits the creation of a log of the cleaning or disinfecting history of each RFID tagged space or patient room or reusable medical equipment through a computer or data warehouse.

In another embodiment, the RFID tag may be attached to a patient or located in a patient room or uncleen piece of equipment and resets the status of the designated space for future disinfection. In another embodiment, the disinfection means may communicate with multiple RFID tags and automatically cleans the entire room.

In an embodiment of the invention, the RFID tag(s) in the designated area may be passive. In another embodiment, the RFID tags in the designated area are active.

The advantage of the invention includes the ability to keep track of clean and dirty equipment log; ability to distinguish clean and dirty equipment at the end user level with or without having to manually scan equipment; ability to prevent errors by introducing various safety mechanisms like alarms if equipment that is not clean is going to be used; ability to communicate with the environmental services personnel on a real time basis; ability for the hospital management and infection prevention and control teams to keep track of cleaning processes on a real time basis, for both equipment and patient rooms; and/or providing complete automation of the process thus preventing or reducing medical errors that can be fatal.

The following examples are provided to further illustrate aspects of the invention. These examples are non-limiting and should not be construed as limiting any aspect of the invention.

**EXAMPLES**

**Example 1**

For example, patient X is admitted to the hospital and is in Room A. Patient X has MRSA and has spent 5 days in Room A. During the course of his/her hospitalization, patient X touches several surfaces in Room A and also requires IV antibiotics which are delivered by an IV pump attached to an IV pole. Patient X is ready for discharge on day 5. This information is electronically entered in the hospital medical records. This electronic entry updates the housekeeping database alerting the environmental services personnel that Room A will soon be ready for cleaning. Once patient X has left the hospital, the software on the Jangama’s display unit or Jangama integrated into a no-touch device such as Xenex or Bioquell displays that Room A is now available for cleaning. The environmental services personnel are alerted, and one then takes the no-touch device with/without Jangama (depending whether Jangama is integrated or standalone) to Room A for cleaning. Room A is tagged with an active RFID tag, which updates the time of entry of device for cleaning. Once cleaning of Room A is in progress, the hospital bed database indicates the same. Once Room A is terminally cleaned per protocol and the no-touch device is taken out of the room, the room RFID tag updates the hospital bed database regarding the availability of that room. This interrogation upon entry and exit is provided by the RFID reader in the Jangama or no-touch device with Jangama incorporated within it. The bed management system may further have several disinfection positions that need to be completed before the room is marked as completely disinfected for certain type of disinfection systems; e.g., the Xenex device has three to four positions and the Tru-D has two positions. Merely by way of example, the disinfecting means may be arranged as follows: one for a bathroom, two in living rooms (one on either side of the bed) and one in an anteroom, if present. The bed management system shows the progress on the display as each position is completed.

A protocol for disinfecting medical equipment and obtaining a signal that disinfection is complete is as follows:

1. **Medical equipment of interest may be tagged with a unique identifier.** Examples of suitable identifiers include: RFID tag or tags (active, passive or combined), or RFID equivalents like: RTLS tags, Bluetooth ID, Wi-Fi tags, near field communication technologies or infrared tags. Active RFID tags may include LED display technology or E-ink display technology.

2. **The tagged equipment may be able to communicate with both a mobile RFID reader(s) and/or fixed RFID reader(s) as shown in FIG. 61, each of which in turn may communicate with a central data warehouse (management system).** The mobile and the fixed RFID reader(s) may also have the capacity to communicate with each other. A facility’s existing Wi-Fi technology may be used to enable such communications.

3. **Equipment holding rooms may have a fixed RFID reader.**

4. **Each UV device or equivalent device of interest may incorporate a mobile RFID reader in the form of an incorporated Jangama or standalone Jangama (FIG. 9).**

5. **This uniquely tagged RME may have a communication range of 20-30 feet with a mobile RFID reader and a range of e.g. 4-6 feet for the fixed reader.**

6. The mobile reader may also have the capacity to transmit or emit signals back to the RFID tagged RME. It also may have storage and display capacity.

7. **The fixed readers may also have the capacity to transmit or emit signals back to the RFID tagged equipment.**

8. **The portable reader/transmitter/storage/display device (Jangama) may have many functions:**

   a. **Receive and recognize each of these uniquely tagged ID equipment;**
b. Ability to measure UV radiation around it or Hydrogen peroxide levels around it;
c. Ability to send signals to the uniquely tagged equipment while in the vicinity of e.g. 4-6 feet;
d. Ability to communicate with fixed reader/transmitter;
e. Ability to communicate with management systems (data warehouse);
f. Ability to store information in its hardware to later transmit to the management system in case there are Wi-Fi issues; and/or
g. Ability to display live information about bed management as well as the ability to identify unclean equipment in the vicinity (e.g., about 20-30 feet).

9. The fixed sensor/emitter may have these functions:
   a. Receive and recognize each uniquely tagged RME within the room, at entry and exit (choke holder);
   b. Communicate with the RME being removed from/brought in the holding area, and changing the signal on the RME appropriately;
   c. Communicate with the mobile reader; and/or
   d. Communicate with management to provide a real time feed about equipment cleaning necessity.

10. The tagged equipment may have a receiver for a unique signal. Once the unique signal is received it may display e.g. a green light and or display the word “CLEAN”. The signal may be valid for 24 hours but may be customizable, for example, up to 72 hrs.

11. The tagged RME may return to unclean or red display representing unclean in the following exemplary scenarios:
   a. No unique signal from mobile reader/transmitter or fixed reader/transmitter;
   b. 24 hours have elapsed since the last signal (automatically change status to either red light or display “Unclean” at the end of 24 hours or the customized time);
   c. The equipment leaves a designated area (i.e., holding area); and/or
   d. The equipment was not within 4-6 feet of the UV unit or in the case of Hydrogen peroxide within the contained room.

12. The room itself may be lined with unique aluminum screen with panels at 45 degrees (FIG. 2) or coated with UV reflective paint. The aluminum and paint are great reflectors of UV light and may make the process of disinfection more effective as the equipment may be cleaned through the reflectivity of the panels or paint.

13. The mobile reader/transmitter/recorder (Jangama) may be equipped with an activity log for example. If equipment #1 was in the vicinity (e.g., 4 feet) of the sensor at this time date, it receives a total radiation of X units (0-5000 milliJoules/cm²) or other comparable units) for Y time (0-300 minutes) and receives the clean signal at z time. If taken out of the area, the central database may record the last time taken out of clean area and record it and merge the data with the mobile unit. A data log may be maintained in MS Excel format or equivalent tabular format in the central data warehouse and may be printable for regulatory (Joint Commission/Center for Medicaid and Medicare Services) compliance and available on request during an inspection.

14. The central data warehouse may be equipped with appropriate software to recognize all tagged equipment, their location and searchable properties like some of the current RFID vendors. The database may also be able to locate each UV or equivalent device and be able to update device maintenance logs.

15. If unclean equipment is accidentally wheeled out of the designated home area then it may produce an auditory signal (e.g., beeping noise) for designated period of time and a visual signal (e.g., a light signal such as a red light signal) may blink for a designated period to alert the handler that the equipment is not clean.

Active Management of Clean and Unclean Inventory using Pulsed Xenon UV Device: (FIGS. 1 and 12)

Each RFID tagged equipment is managed by a Wi-Fi or similar technology enabled central data warehouse through a fixed reader in each of the equipment holding rooms. Once the UV device equipped with the Jangama is used to clean the RME it records each uniquely tagged RME and transmits that information to the central database/Management system. This database may be updated on a real time basis, meaning, that it updates with each cleaning and each movement of the RFID tagged RME. An UV operator may be able to access this information on the UV device using the mobile reader/transmitter/storage/display unit (Jangama) on a real time basis as well, thereby alerting the operator to the holding areas in need of cleaning. Also, if the UV device is in the vicinity of a holding area with a high volume of unclean equipment, an instant beep/message (like a text message) may alert the operator handling the device. This may help with active management of clean/unclean equipment especially near high turnover areas such as Operating Rooms. The same device may also display a feed as to where the next room needs to be cleaned. This feed may be fed to the display screen on Jangama from bed control or bed management as well as central data warehouse servers.

The mobile sensor/emitter/storage device (Jangama) may also communicate with the fixed readers and all the RMEs in disinfection range (e.g., 4-6 feet) which may be targeted for cleaning. The fixed readers as shown in FIG. 6 may change the status of these visible RME in the central database to “BEING DISINFECTED”. Once the disinfection process is completed, the portable sensor of Jangama may communicate the completed cycle to the fixed readers and the database may be updated to “CLEAN” for these visible equipment and the resets the timer on the database and the tagged RME. It may be necessary to relocate the UV device to different sections of the holding room to effectively clean all the equipment (as the room may be bigger than e.g. 6x6x feet), as all equipment in need of cleaning may not be within UV range. At the pre-determined time, the fixed sensor may be able to turn off the “green signal” on all equipment that would no longer be considered clean. Alternatively, the equipment may be manually reset for cleaning. All this communication may occur over the pre-existing Wi-Fi communications available at the facility.

For example, if patient Y in room B needs IV fluids, the nurse may go to the utility/equipment storage room to get an IV pole that can be used for patient Y. All IV poles stored in the storage room have active RFID tags which illuminate “GREEN” for clean and “RED” for unclean as shown in FIG. 10. The storage room entrance/exit area has an active RFID reader. The nurse takes an IV pole which has the “RED” illuminated on it, but fails to notice it. As this IV pole is taken out of the utility room, the RFID reader interrogates the RFID tag on the IV pole and sounds out an alarm that an unclean pole is being taken out of the room. Should the “GREEN” illuminated pole be taken, no alarm will sound. The active tag on the IV pole now is in use in room B, which will result in software update of its exact location.

In another example, patient Y in room B needs IV fluids. The nurse goes to the utility/equipment storage room to get an IV pole that can be used for patient Y. All IV poles stored
in the storage room have active RFID tags which display “Clean” and “Dirty” on an E-ink-based RFID tag as shown in FIG. 10. The storage room entrance/exit area has an active RFID reader. The nurse takes an IV pole which has the word “DIRTY” displayed on it, but fails to notice it. As this IV pole is taken out of the utility room, the RFID reader interrogates the RFID tag on the IV pole and sounds out an alarm indicating that an unclean pole is being taken out of the room. Should a “Clean” displayed pole be taken, no alarm will sound. The active tag on the IV pole now is in use in room B, which will result in a software update of its exact location.

The fixed readers have ability to receive signals from 2 sources. One from fixed RFID reader or Blue tooth device and the other from the mobile reader/transmitter (Jangama).

The fixed reader may communicate all the information with a central server/management system through the facility’s Wi-Fi.

The mobile reader/transmitter (Jangama) may have the ability to communicate with RFID/Blue tooth equipment, fixed sensors, central server or management system through Wi-Fi. The received signal strength indication (RSSI) may be used to determine which room the equipment is in.

Real Time Location Systems (RTLS)

Real Time Location Systems (RTLS) are used to determine the physical proximity of active RFID tags and can be used in conjunction with this invention. This can be accomplished by installing fixed RFID readers at strategic locations where the active tag can be read simultaneously by two or more readers. Pre-determined algorithms can then determine the exact location of the active RFID tags by the RSSI level and triangulation. This would determine an approximate location within e.g. about 10 feet depending on the number of reader access points are used. For further accuracy, additional sensor technologies may be incorporated into the tag such as ultra-sound or infra-red.

Components:
1. RFID tag or equivalent for each equipment that needs to be tracked and disinfected;
2. Mobile sensor/reader/signal emitter/storage device/display device (Jangama);
3. Clean/Unclean display;
4. Green/Red light display;
5. Beep/this device is “unclean” announcer;
6. UV Reflectors/paint in the room;
7. Room readers;
8. Data warehouse master database/Management system.

Example 2

Another way to communicate with the RFID tagged RME may be through a no-touch disinfection (NTD) remote control mechanism where after the NTD device is done disinfecting, the NTD device sends a signal to the RFID tag and turns on a visual display (e.g., a LED light having a green color). In contrast, the default light signal may be a different color, such as red. The greed and red light can illuminate Clean and Unclean, respectively. The lights may be enclosed behind these wordings in a square box with these wording on the front and LED lights behind them. It may also display the words clean or dirty using the E-ink technology.

The mobile continuous RFID reader, mobile display (Jangama) and the tagged equipment may communicate with each other within e.g. 20-30 feet vicinity on a real time basis. Another mobile reader may be activated just like the light after UV radiation for 2 seconds. It may identify and transmit all the equipment ID for e.g. about less than 4 feet range and communicate that information to the management system to verify the equipment status as clean or dirty.

At checkpoints, a reader may be connected to the same mechanism that turns the light off or turns it red if it moves outside of the checkpoint area. The off mechanism (or turning it red/unclean) is triggered by the reader and the reader sends a message to the management system to change the log status as moved and now unclean. The same reader may have a voice alert/beep when unclean equipment is moved from the holding area away from checkpoint. Similarly, all the above information may display on E-ink based RFID tags.

Example 3

The following is a description of a process for creating status updates using the RFID tagged system of the invention.

Process:
1. The designated rooms will have unique identifiers: RFID tags (active, passive or combined), or RFID equivalents like: RTLS tags, Bluetooth ID, Wi-Fi tags, near field communication technologies or infrared tags. The number of tags may range from 2-3 (with at least one in the main room and 1 in the bathroom). The exact number of tags in each room will vary depending on the size of the room. For example, a room that is bigger than 6x6 feet may have at least 2 tags. Also, the central database management or bed management may have the information of number of tags and their locations (positions) in each room.
2. The portable sensor/emitter/storage (Jangama) may have 3 functions:
   a. Receive and recognize each of these unique tagged ID rooms.
   b. Ability to measure UV light amount/exposure around it or Hydrogen peroxide levels around it.
   c. A RFID reader may send a signal back to bed management system; via the existing facility’s Wi-Fi; once the disinfection cycle is completed, thereby indicating the area that has been recently disinfected; on a real time basis.
3. The mobile sensor (Jangama) may not recognize the RFID tags, which are not within e.g., 4-6 feet of the UV unit or in the case of Hydrogen peroxide within the contained room.
4. The portable sensor/emitter/recorder (Jangama) may log each of the activity. For example, that room #1 east wall was in the vicinity (about 4 feet) of the sensor at A time/date, may have received a total radiation of X many units for Y length of time, may have received the clean signal at Z time; room #1 west wall was in vicinity (about 4 feet) of the sensor at B time/date, may have received a total radiation of X many units for Y length of time, may have received the clean signal at Z time; room #1 bathroom was in vicinity (about 4 feet) of the sensor at C time/date, may have received a total radiation of X many units for Y length of time, may have received the clean signal at Z time. Combined data log may be maintained in Excel format or tabular format so it is printable and can be showed to regulatory authorities (JCCMS) on request or during inspection.
5. Once room #1 is thoroughly disinfected, the mobile sensor (e.g., including a Jangama) may update the central database/bed management system and display that particular room as “clean” for bed management and log when all the areas of the room are cleaned (it may remain in
“Unclean” status if the EMS person missed a position for UV disinfection devices like Xenex).
6. Once that particular room is assigned to a patient through bed management, the room may be displayed as “Occupied”.
7. Once the status of an Occupied room is changed to Unoccupied or discharged in bed management, the mobile sensor (Jangama) log may add the particular room to its “To be Cleaned” database, alerting the operator to rooms in need of cleaning. Alternatively, the status of a patient room may be reset to “To be Cleaned” manually or after a predetermined time.

Example 4

The “Jangama” may have the following components (FIG. 9):
1. RFID reader
2. RFID Controller
3. Display panel/unit
4. Wi-Fi network integrator (transmitter/receiver)
5. Microprocessor
6. Sensor signal processor
7. Sensors
8. Antennas
9. Memory

The Jangama can be integrated into an existing no-touch disinfection device circuit board instead of a stand-alone device.

Breakdown of the communication that occurs between different components of the system:

The RFID reader or interrogator or controller could be the same unit. The reader/interrogator reads and retrieves information, whereas the controller can send a signal back to RFID tag to rewrite memory or change display settings.

The RFID Reader sends an query which is received by a RFID tag. Active RFID tags store memory values corresponding to clean or dirty that are accessed by the reader. At the same time, each RFID tags unique ID is also accessed. This information is transmitted from the RFID reader to the mobile unit as well as the data warehouse through a Wi-Fi or wired network.

2. Communication between the RFID Controller and Active Tag:

The controller sends an query to a RFID tag, which replies to the controller the currently stored memory value (corresponding to clean or dirty) along with the unique ID for the tag. This information is also relayed to the microprocessor which in turn communicates with the stored information in the data warehouse through the existing Wi-Fi system.

Once the condition for disinfection is satisfied, the microprocessor asks the controller to issue a clean signal back to the active RFID tag and change the memory value to clean. The signal will only be used to active RFID tags within a set distance parameter of 4 feet, but this could also be anywhere from 0-8 feet. The controller confirms the RFID tags unique ID/new memory value and sends it back to the processor. The microprocessor then relays the new values to the data warehouse to clean status for those RFID tags which were in the set distance parameter with a time and date stamp.

3. Communication between Sensor Signal Processor and Sensors:

Sensors can sense various external parameters like UV, Hydrogen peroxide. Once a certain upper limit threshold is reached in the sensor (10-500 millijoules/cm² for UV or 10-20000 ppm for hydrogen peroxide), it sends a signal to the sensor signal processor, which changes the signal to a digital numerical signal for the microprocessor to know that certain values have been met (FIG. 7). This is referenced to the stored upper threshold limit for that sensor. If above the threshold, the microprocessor asks the RFID controller to issue a clean signal to the active RFID tags in a certain distance parameter as described above.

4. Sensors:

The sensors can sense various parameters like temperature, pressure, mercury based UV dose (continuous), pulsed xenon UV dose (cumulative dose of all pulses in a set time), or hydrogen peroxide concentration. The sensors transmit the sensed dose to the sensor signal processor.

5. Sensor Signal Processor:

The function of the sensor signal processor is to take the sensor input, convert it to a digital signal and reference it to the stored reference parameters for various external effects like UV or Hydrogen peroxide. Once the threshold is reached, the sensor signal processor will continue to send a signal of condition being met to the microprocessor for next 60 seconds or until the sensor keeps sending the signals whichever is later. For those 60 seconds, the sensor value will be set as met, even if the sensors sense no additional UV or H2O2. After 60 seconds, the parameters will be set to below threshold until another sensing event happens (FIG. 7). The sensor value could be set as met, for example, anywhere from 30 seconds to 300 seconds.

In yet another example, the sensors and sensor signal processors may be optional, when time is used as a parameter to fulfill condition being met algorithm for the RFID controller to transmit a signal to nearby RFID tags. This is usually possible if the circuit board is integrated into a no-touch disinfection system such as Pulsed UV or Mercury based UV or Hydrogen peroxide systems thus eliminating the need to sense an external signal.

6. Display Unit:

The display unit will have the capacity to display various things similar to a notebook or tablet. It can display information newly created in the microprocessor or the live information it receives from the data warehouse servers through Wi-Fi or live bed management feed through the bed management servers. It can also display information about any dirty RME in the near vicinity of the mobile unit. It can also perform other functions of a display unit like other tablets or phones receiving text or video chat again through a Wi-Fi network or a cellular network.

7. Microprocessor:

The microprocessor is like the brain of Jangama. Its function as a central processing unit for all external signals and is also the origin of digital signal either generated intrinsically or with the help of algorithms in the data warehouse. The digital signals are then relayed to RFID interrogator or controller for their actions to follow. Digital signals that originate from the microprocessor are continuously fed to the data warehouse for determining the location or position of a RFID tagged device much like a mobile RFID interrogator. Similarly, the microprocessor has the ability to process the data warehouse algorithms and convert them to RFID signals through controller and interrogator to read or write a RFID tag or change the memory in a RFID tag to change the clean or dirty labeling on the RFID tags that met the condition. In another example, the microprocessor may take the form of a FPGA (Field Programmable Gate Array), ASIC (Application Specific Integrated Circuit), or some other form of integrated analog or digital logic gates.
7. Data Warehouse:
   The data warehouse houses all the information required to function. Certain new algorithms will need to be incorporated into existing real-time location services algorithms. A new algorithm to match the sensor values, distance, exposure time and choke point integration will be added. Alternatively, these algorithms may be incorporated into the devices themselves.

8. The Active RFID with Display:
   The RFID with display could be of many varieties as previously described in the detailed description.

9. Memory:
   The memory component is optional and will mainly interact with the microprocessor and store data for a limited period of time. It is crucial to have data stored in this backup memory in case there are transmission issues with the facility Wi-Fi. Data could be stored until a proper Wi-Fi connection is established or until the unit can be manually synced with the central data warehouse.

Example 5

In another embodiment the invention would be used in conjunction with choke points. If tagged equipment is allowed to move from one area to another area after it has been designated as clean, or if clean equipment moves from one patient to another patient room for example, this may lead to cross contamination. To prevent this, the data warehouse will include a choke point algorithm.

For example, a piece of clean equipment is taken out of a current patient room as it was not being used. Doing so carries the risk of cross contamination. To prevent this, each patient room will have a choke point, where an active reader will sense the movement of the tagged equipment beyond a certain point and send a signal to the data warehouse, which confirms the equipment movement out an area and that it should not have been moved. The data warehouse will send a signal back to the controller at the choke point to change the RFID memory to dirty as soon as the choke point is breached. It will also sound an alarm to the end user about the equipment being dirty now. Such algorithms already exist in, for example, Avante’s RFID “Access-Trakker,” but such algorithms need to be modified to accommodate current aspects of invention as related to clean and dirty equipment. Integration of such concept will improve patient safety and prevent medical errors.

Similarly clean equipment could be in a clean equipment room. If the clean equipment were to travel outside the clean equipment room, the reader at the choke point will send a signal to the data warehouse that the clean equipment was removed from the clean equipment room. The data warehouses algorithm will track this equipment’s movements for next 5 minutes. If the equipment enters another patient room, then the readers at the patient rooms’ choke point will note this entry. Once this equipment enters the patient room, the data warehouse will change the entry to dirty and should the equipment be taken out again beyond the patient room’s choke point, a similar process as described above will happen.

Similarly in the clean equipment room, if dirty equipment is being placed back in the room, the choke point reader will confirm with the data warehouse of the equipment’s dirty status, record the movement and change the display on equipment to unclean, and keep the equipment handling person that the equipment is unclean.

Various such algorithms will be built to track the equipment at various stages of its movement alerting the end user both visually as well as by noise about the current status of the equipment.

Example 6

The following are algorithms that may be used in conjunction with the Jangama invention:

Algorithm 1: UV dose fulfillment algorithm
   1. Operator starts UV disinfection device
   2. Sensor starting sensing UV and calculates the cumulative dose of UV
   3. Once UV dose exceeds 100 milliwatts/cm² or millijoules/cm² then required dose for disinfection is fulfilled.

Algorithm 2: Hydrogen peroxide fulfillment algorithm
   1. Operator starts hydrogen peroxide disinfection device
   2. Sensor starting sensing hydrogen peroxide and calculates the cumulative dose of hydrogen peroxide.
   3. Once hydrogen peroxide dose exceeds 2000 parts per million (ppm) then required dose for disinfection is fulfilled. The ppm range is 2-20,000 as it varies on the size of the MOM.

Algorithm 3: Dirty to Clean Display Change Algorithm
   1. Once UV dose criteria is fulfilled the RFID controller will send a back to the RFID tags message and it will change the stored value that represents “dirty” to stored value for memory that represents “clean”.

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What is claimed is:
1. A method for organizing the disinfection of one or more items contaminated with biological agent(s), comprising:
   a) exposing an item to a disinfectant for a period sufficient to disinfect the item, said item having a wireless communication tag;
   b) sending a first signal to the wireless communication tag when disinfection is complete to update a display on the wireless communication tag to be indicative of a clean status for the item;
   c) subsequently detecting movement of the item past a chokepoint; and
   d) sending a second signal to the wireless communication tag to alter the display upon a lapse of a predetermined amount of time after detecting movement of the item past the chokepoint.
2. The method of claim 1, wherein the step of sending the first signal to the wireless communication tag comprises manually activating the signal transmission.
3. The method of claim 1, wherein the step of sending the first signal to the wireless communication tag is automated from a computer of a system that tracks disinfection of multiple items.
4. The method of claim 1, wherein the item is a device, machine or tool.
5. The method of claim 4, wherein the device, machine or tool is medical equipment.
6. The method of claim 5, wherein the medical equipment is a durable medical equipment or reusable medical equipment.
7. The method of claim 1, wherein the disinfectant is ultraviolet light.
8. The method of claim 7, wherein the ultraviolet light is from a pulsed xenon lamp.
9. The method of claim 7, wherein the ultraviolet light is from a mercury lamp.
10. The method of claim 1, wherein the disinfectant is hydrogen peroxide.
11. The method of claim 10, wherein the hydrogen peroxide is hydrogen peroxide vapor, activated hydrogen peroxide, dry mist hydrogen peroxide or vaporized hydrogen peroxide.
12. The method of claim 1, wherein the biological agent is selected from a group consisting of a micro-organism, viral particles, fungal particles, an infectious agent, microbes in human secretion, human excretion, dust and fecal material.
13. The method of claim 1, further comprising processing the first signal so as to create a log of the cleaning or disinfecting history of the item.
14. The method of claim 1, wherein the step of sending the signal to the wireless communication tag is automated from a disinfectant sensor.
15. The method of claim 14, wherein the disinfectant sensor is an ultraviolet light dose sensor.
16. The method of claim 14, wherein the disinfectant sensor is a sensor for detecting a fluidic disinfectant.
17. The method of claim 1, wherein the second signal is a signal to alter the display to be indicative of an unclean status.
18. The method of claim 17, further comprising sounding an alarm upon sending the signal.
19. The method of claim 1, wherein the first signal is to change visual information exhibited on an electronic visual display of the wireless communication tag.

20. The method of claim 1, wherein the first signal is to illuminate a light source on the display of the wireless communication tag.

21. The method of claim 1, wherein the wireless communication tag is a radio-frequency identification tag.

22. The method of claim 1, wherein the step of detecting movement of the item past the chokepoint comprises detecting the item entering a space associated with the chokepoint.

23. The method of claim 22, further comprising sending a third signal to the wireless communication tag to alter the display upon detecting movement of the item past the chokepoint to be indicative of entry of the item into the space associated with the chokepoint.

24. The method of claim 22, further comprising detecting the item exiting a holding area prior to detecting the item entering the space associated with the chokepoint.

25. The method of claim 24, further comprising sending a third signal to the wireless communication tag to alter the display upon detecting the item exiting the holding area.

26. The method of claim 1, wherein the step of detecting movement of the item past the chokepoint comprises detecting the item exiting a space associated with the chokepoint.

27. The method of claim 26, sending a third signal to the wireless communication tag to alter the display upon detecting movement of the item past the chokepoint to be indicative of the item exiting the space associated with the chokepoint.

28. The method of claim 1, wherein the predetermined amount of time is approximately 24 hours or longer.

29. The method of claim 1, wherein the chokepoint is associated with an area for storing medical equipment.

30. The method of claim 1, wherein the chokepoint is associated with a patient room.

31. The method of claim 30, further comprising:
   a) sending a fourth signal to the wireless communication tag to alter the display on the wireless communication tag upon receiving the third signal.
   b) determining an approximate location of the portable disinfection apparatus in the designated space from the obtained signals; and
   c) associatively recording the approximate location of the portable disinfection apparatus and a duration of the period the item is exposed to the disinfectant.

32. A method for disinfection of an item contaminated with biological agent(s) in a designated space, comprising:
   a) exposing a designated space to a disinfectant discharged from a portable disinfection apparatus for a period sufficient to disinfect an item in the designated space, wherein some non-portable surfaces of the designated space have a wireless communication tag;
   b) obtaining a signal from at least some of the wireless communication tags regarding their proximity to the portable disinfection apparatus before or during the period the item is exposed to the disinfectant and/or when the period the item is exposed to the disinfectant is complete;
   c) associatively recording the approximate location of the portable disinfection apparatus in the designated space from the obtained signals; and
   d) associatively recording the approximate location of the portable disinfection apparatus and a duration of the period the item is exposed to the disinfectant.

33. The method of claim 32, wherein the wireless communication tags communicate with the portable disinfection apparatus.

34. The method of claim 32, wherein the wireless communication tags communicate the initiation or completion of the cleaning or disinfection cycle to a computer of a system that tracks multiple designated spaces.

35. The method of claim 32, further comprising processing the signals so as to create a log of the cleaning or disinfecting history of the designated space.

36. The method of claim 32, wherein the wireless communication tags are radio-frequency identification tags.

37. The method of claim 32, wherein the wireless communication tags are passive.

38. The method of claim 32, wherein the wireless communication tags are semi-passive.

39. The method of claim 32, wherein the wireless communication tags are active.

40. The method of claim 32, further comprising detecting entry of the portable disinfection apparatus into the designated space prior to exposing the designated space to the disinfectant.

41. The method of claim 32, wherein the step of exposing the designated space to a disinfectant comprises successively exposing different portions of the designated space to a disinfectant for periods respectively sufficient to disinfect items in the different portions, wherein the step of determining the approximate location of the portable disinfection apparatus in the designated space comprises determining an approximate location of the portable disinfection apparatus in each of the different portions of the designated space from the obtained signals, and wherein the step of associatively recording comprises associatively recording the approximate location of the portable disinfection apparatus in each of the different portions of the designated space and a duration of each of the disinfection periods.

42. The method of claim 32, wherein the designated space is a patient room, an operating room or a storage room.

43. The method of claim 32, wherein the disinfection period is a period sufficient to disinfect items that are at least 1 foot away from the portable disinfection apparatus.

44. A method, comprising:
   a) exposing a designated space to ultraviolet light, said designated space having one or more wireless communication tags;
   b) receiving a signal that a predetermined amount of ultraviolet light has been detected; and
   c) subsequent to receiving the signal, sending a different signal of a predetermined strength to update displays only on a set of the one or more wireless communication tags that are within a set distance from the source of the ultraviolet light, wherein the predetermined strength is less than a signal strength constituting a maximum communication range of a transmitter sending the different signal, and wherein the different signal is to update the displays on the set of the one or more wireless communication tags to be indicative of a clean status for surfaces in the designated space comprising the respective set of one or more wireless communication tags.

45. The method of claim 44, further comprising exposing the designated space to a fluidic disinfectant prior to exposing the item to ultraviolet light.

46. The method of claim 44, wherein the different signal is to change visual information exhibited on an electronic visual display of the wireless communication tag.

47. The method of claim 44, wherein the different signal is to illuminate a light source on the display of the wireless communication tag.

48. The method of claim 44, wherein the set distance is equal to or less than an upper limit of a set disinfection range of a source of the ultraviolet light.
49. The method of claim 44, wherein the set distance is less than a dimension of the designated space.

50. The method of claim 44, wherein the designated space is an area suitable for human occupancy.

51. A method for organizing the disinfection of one or more items contaminated with biological agent(s), comprising:
   a) exposing an item to a disinfectant for a period sufficient to disinfect the item, said item having a wireless communication tag;
   b) sending a first signal to the wireless communication tag when disinfection is complete to update a display on the wireless communication tag to be indicative of a clean status for the item;
   c) subsequently detecting movement of the item past a chokepoint or outside a vicinity of the chokepoint, wherein the chokepoint is associated with a space in which the item is exposed to the disinfectant; and
   d) sending a second signal to the wireless communication tag to alter the display to be indicative of an unclean status for the item upon detecting movement of the item past the chokepoint.

52. The method of claim 51, wherein the step of detecting movement of the item past the chokepoint comprises detecting the item exiting a space associated with the chokepoint.

53. The method of claim 52, further comprising detecting movement of the item past the chokepoint that is indicative of the item entering the space associated with the chokepoint subsequent to the step of sending the first signal to the wireless communication tag and prior to the step of detecting movement of the item past the chokepoint that is indicative of the item exiting the space associated with the chokepoint.

54. The method of claim 53, further comprising detecting the item exiting a holding area subsequent to the step of sending the first signal to the wireless communication tag and prior to detecting the item entering the space associated with the chokepoint.

55. The method of claim 51, wherein the space associated with the chokepoint is a patient room.

56. The method of claim 55, further comprising:
   a) receiving a third signal indicating a patient has been discharged from the patient room prior to detecting movement of the item past the chokepoint; and
   b) sending a fourth signal to the wireless communication tag to alter the display on the wireless communication tag upon receiving the third signal.

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