A method and an apparatus for detecting and quantifying bacterial spores on a surface. In accordance with the method: bacterial spores are transferred from a place of origin to a test surface, the test surface comprises lanthanide ions. Aromatic molecules are released from the bacterial spores; a complex of the lanthanide ions and aromatic molecules is formed on the test surface, the complex is excited to generate a characteristic luminescence on the test surface; the luminescence on the test surface is detected and quantified.
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Figure 1A

DPA-rich core.
Figure 1B

Absorption → Energy Transfer → Emission

DPA → Tb

Tb$^{3+}$
Figure 1C

The diagram illustrates the energy transfer process involving the states of DPA and Tb$^{3+}$ ions. The transitions are labeled as follows:

- $^1\pi^*$ to $^3\pi^*$: Absorption
- $^3\pi^*$: Energy Transfer
- $^1\pi$: Emission

The states are represented as:

- DPA $^7F_{J=6-0}$
- Tb$^{3+}$$^5D_4$

The figure shows the sequential steps in the energy transfer process.
Figure 2A
Figure 2B
Figure 3
Figure 4
Figure 7
METHOD AND APPARATUS FOR DETECTING AND QUANTIFYING BACTERIAL SPORES ON A SURFACE

CROSS REFERENCE TO RELATED APPLICATIONS


STATEMENT OF GOVERNMENT SUPPORT

The present invention was made with support from the United States Government under Grant number NAS7-1407 awarded by NASA. The United States Government has certain rights in the invention.

BACKGROUND

1. Field

The present disclosure relates to the field of chemical detection. In particular, a method and apparatus for detecting and quantifying bacterial spores on a surface is disclosed.

2. Description of Related Art

Lanthanide complexes, particularly those of terbium (Tb³⁺) and europium (Eu³⁺), exhibit luminescence properties for the detection of aromatic biomolecules. The detection scheme is based on the absorption-energy transfer-emission mechanism, which is triggered by the binding of aromatic ligands to lanthanide complexes under UV excitation. Recent efforts have been focused on the detection of dipicolinic acid (DPA) (2,6-pyridinedicarboxylic acid), which is a unique constituent of bacterial spores present at high concentrations (up to 1 M). Dipicolinic acid is also a commercially available product having the following characteristics: CAS #: 499-83-2, Synonyms: 2,6 Pyrydine Dicarboxylic Acid, Molecular Formula: C₁₁H₄NO₄, Molecular Weight: 167.12, Description: White crystalline powder, Sulphated Ash: 0.3% max, Moisture Content: 0.5% max, Melting Point: 242.0 to 245.0 degree C., Assay: 99.0% min.

Bacterial spores are generally accepted to be indicator species for validating sterile balance since they are the most resilient form of life against sterilization regimens (Hindle and Hall, 1999 Analyst, 124, 1599-1604). Sterility testing of surfaces is traditionally performed by either (1) swabbing the surface with a cotton applicator, resuspending the swabbed spores, and plating the spore suspension onto growth media; or (2) using Replicate Organism Detection and Counting (RODAC) growth plates that are pressed against a surface to be analyzed. Each of these two bacterial assay steps requires 3-5 days before results are available.

As mentioned, dipicolinic acid (DPA) is present in high concentrations (about 1 molar or about 15% of by weight) in the core of bacterial spores (Murell, 1969, Bact. Spore 1, 216). In its deprotonated state, DPA is dipicolinate (DP) and is found in a 1:1 complex with Ca⁺² inside the spore, as shown in FIG. 1A. For all known life-forms, DPA is unique to bacterial spores and is naturally released into bulk solution upon germination—the process of spore-to-vegetative cell transformation. DP can also be released upon lysis of the bacterial spore. Thus, DPA and/or DP are indicator molecules for the presence of bacterial spores. DPA is a classic inorganic chemistry ligand that binds metal ions with high affinity. As mentioned, DPA takes the form of dipicolinate (DP) in its deprotonated form that binds to Ca⁺². DPA binding to terbium ions or other lanthanides exhibits luminescence under UV excitation as shown in FIGS. 1B and 1C. The green luminescence turn-on signal indicates the presence of bacterial spores. The intensity of the luminescence can be correlated to the number of bacterial spores per milliliter.

U.S. patent application Publication No. 2003-0138876 for "Method bacterial endospore quantification using lanthanide dipicolinate luminescence" discloses a lanthanide complex that is combined with a medium to be tested for endospores. Dipicolinic acid released from the endospores bonds the lanthanides, which have distinctive emission (i.e., luminescence) spectra, and are detected using photoluminescence. The concentration of spores is determined by preparing a calibration curve that relates emission intensities to spore concentrations for test samples with known spore concentrations. A lanthanide complex is used as the analysis reagent, and is comprised of lanthanide ions bound to multidentate ligands that increase the dipicolinic acid binding constant through a cooperative binding effect with respect to lanthanide chloride. The resulting combined effect of increasing the binding constant and eliminating coordinated water and multiple equilibria increases the sensitivity of the endospore assay by an estimated three to four orders of magnitude over prior art of endospore detection based on lanthanide luminescence.

U.S. patent application Publication No. 2004-0014154 for "Methods and apparatus for assays of bacterial spores" discloses a sample of unknown bacterial spores which is added to a test strip. The sample of unknown bacterial spores is drawn to a first sample region on the test strip by capillary action. Species-specific antibodies are bound to the sample when the unknown bacterial spores match the species-specific antibodies, otherwise the sample is left unbound. DPA is released from the bacterial spores in the bound sample. Terbium ions are combined with the DPA to form a Tb-DPA complex. The combined terbium ions and DPA are excited to generate a luminescence characteristic of the combined terbium ions and DPA to detect the bacterial spores. A live/dead assay is performed by a release of the DPA for live spores and a release of DPA for all spores. The detection concentrations are compared to determine the fraction of live spores. Lifetime-gated measurements of bacterial spores to eliminate any fluorescence background from organic chromophores comprise labeling the bacterial spore content with a long-lifetime lumophore and detecting the luminescence after a waiting period. Unattended monitoring of bacterial spores in the air comprises the steps of collecting bacterial spores carried in the air and repeatedly performing the Tb-DPA detection steps above.

Exciting the combined terbium ions and DPA generates a luminescence characteristic of the combined terbium ions and DPA. This is achieved by radiating the combined terbium ions and DPA with ultraviolet light.

U.S. patent application Publication No. 2004-0014154 further discloses a method for live/dead assay for bacterial spores comprising the steps of: providing a solution including terbium ions in a sample of live and dead bacterial spores;
A method is provided for detecting and quantifying individual bacterial spores comprising: releasing DPA from viable bacterial spores by germination from a first unit of the sample; combining the terbium ions with DPA in solution released from viable bacterial spores; exciting the combined terbium ions and DPA released from viable bacterial spores to generate a first luminescence characteristic of the combined terbium ions and DPA; detecting the luminescence to determine the presence of the bacterial spores; and generating an alarm signal when the presence of viable bacterial spores is detected or the concentration thereof reaches a predetermined magnitude.

Currently, bioburden levels are determined using the culture-dependent methods, with which bacterial spores are quantified in terms of colony forming units (CFU's) that become visible on growth plates after incubation. There are several limitations for culture-dependent methods. First, this process requires 3-5 days to complete. Second, a large number of bacterial spores can aggregate on individual particulates giving rise to a single CFU, and thus a large underestimation of the bioburden. Third, colony-counting methods only account for cultivable spore-forming species, which constitute less than 1% in environmental samples. It is desirable to provide a more efficient and sensitive method and apparatus for transferring all spore-forming bacteria (most especially bacteria of the genus *Bacillus* and *Clostridium*) originating on a surface, in the air or in water to a test surface, quantifying the spores, and further characterizing these spores as viable or nonviable.

**SUMMARY**

According to a first aspect of the present disclosure, a method is provided for detecting and quantifying individual bacterial spores comprising: capturing bacterial spores; transferring the bacterial spores to a test surface; providing one or more lanthanide ions on the test surface; releasing aromatic molecules from the bacterial spores on the test surface; forming a complex of the one or more lanthanide ions and the aromatic molecules on the test surface; and an imaging apparatus is provided for detecting and quantifying bacterial spores comprising: an ultraviolet light radiation device to excite a complex of lanthanide ions and aromatic molecules; and generate a characteristic luminescence of the complex; a microscope for detecting and quantifying bacterial spores exhibiting the luminescence of the complex; and an imaging device connected with the microscope for imaging the luminescence.

**BRIEF DESCRIPTION OF THE DRAWINGS**

FIG. 1A is a microscopic image of a spore (about 1 µm in diameter) highlighting a DPA rich spore core.

FIG. 1B is a diagram of a Tb³⁺ ion (shaded ball) which by itself has a low absorption cross section (<10 M⁻¹·cm¹) and consequently has low luminescence intensity. The Tb³⁺ ion can bind the light harvesting DPA/DP (absorption cross section >10⁶ M⁻¹·cm⁻²) originating from the spore. DPA/DP binding gives rise to bright Tb luminescence.

FIG. 2A is a photograph of a Tb³⁺ ion added but no L-alanine.

FIG. 2B is a photograph of a Tb³⁺ ion added but no L-alanine.

FIG. 2C is a photograph of a water filter for capturing bacterial spores from water (Example 1).

FIG. 2D is a photograph of an air filter as described herein for capturing bacterial spores from air (Example 1).

FIG. 3 is a photograph of a water filter for capturing bacterial spores from water (Example 1).

FIG. 4 is a photograph of an air filter as described herein for capturing bacterial spores from air (Example 1).
FIG. 4 depicts four 1 mm thick pieces of PDMS inoculated with *B. subtilus* spores. (A) Plasma cleaned and placed onto Tb**³⁺**-doped agar (B) Plasma cleaned and placed onto Tb**³⁺**-doped agar. (C) Not plasma cleaned, placed onto Tb**³⁺**-doped agar. (D) Not plasma cleaned, place onto Tb**³⁺**-doped agar.

FIG. 5 depicts a schematic apparatus for imaging quantifying and counting of bacterial spores (Example 3).

FIG. 6 depicts Eu**²⁺** microspheres (1 μm) on fluorescent paper imaged with an ImageX-TGi gated CCD camera mounted on a Carl Zeiss fluorescence microscope with 40-times objective, excited with a 300 Hz Perkin Elmer flashlamp. Images are obtained (A) without gating, (B) with gating (100 µs delay, 2.7 µs gate), and (C) 100 µm reference graticule to estimate spatial resolution.

FIG. 7 depicts two lifetime-gated photographs showing bacterial spores on R2A agar before germination (left portion of the figure) and after germination (right portion of the figure).

DETAILED DESCRIPTION

Transfer of Bacterial Spores from Place of Origin

The lanthanide ion-DPA/DP luminescence assay can be employed to detect individual bacterial spores from a place of origin. DPA/DP refers to DPA and/or DP. In other words, DPA/DP means at least one between DPA and DP. A place of origin includes any solid surface, water and/or air. In order to posit the bacterial spores onto a test surface, the bacterial spores are first captured from a place of origin. A place of origin can include an infinite number of possibilities. Bacterial spores on solid surfaces are transferred from the solid surface onto a cotton swab (FIG. 2A), or an adhesive polymer, such as PDMS (polydimethyl siloxane) agar or agarose (FIG. 2B). Bacterial spores in water are transferred from water onto a water filter (e.g. membrane filter)(FIG. 2C). Bacterial spores in the air are transferred from the air onto a test surface (FIG. 2D). Examples of each of these types of transfer methods are described in Example 1.

The step of collecting bacterial spores carried in the air comprises capturing the bacterial spores with an aerosol sampler or impactor. Preferably, the step of collecting bacterial spores carried in the air comprises continuously sampling the air. In one embodiment, air is passed over quartz filter tape using an air sampler (Example 1). Alternatively, air can be passed over lanthanide-doped agar using an air sampler.

Test Surface

The test surface may be the same surface onto which the bacterial spores were transferred from the place of origin. In this case, a second transfer onto a test surface is not necessary. However, if the transferring surface is not to be the test surface, then the bacterial spores are transferred onto the test surface. The present invention provides a method of using a test surface on which bacterial spores are posited. Once the bacterial spores are located on the test surface, they can be induced to release DPA/DP by germination and/or physical lysis.

In one embodiment, the test surface contains an adhesive polymer. In another embodiment, the test surface contains (is "doped with") a germinating agent. In another embodiment, the test surface, onto which bacterial spores are transferred, contains (is "doped with") a lysing agent or is subjected to a method of lysis. In another embodiment, the test surface contains (is "doped with") lanthanide ions. In yet another embodiment, the test surface is transparent allowing for detection of the luminescence produced from the excited lanthanide-DPA complex. In a preferred embodiment, the test surface is an adhesive polymer (PDMS, agar, agarose) that contains a germinating agent, contains lanthanide ions, and is transparent, allowing for detection of lanthanide-DPA luminescence (FIG. 3).

In a second preferred embodiment, the test surface is an adhesive polymer that contains a germinating agent, contains or is subjected to a lysing agent or method of lysis, contains lanthanide ions, and is transparent, allowing for detection of lanthanide-DPA luminescence. Alternatively, a test surface is an adhesive polymer that contains or is subject to a lysing agent or a method of lysis, contains lanthanide ions and is transparent, allowing for detection of lanthanide-DPA luminescence (FIG. 4).

According to one embodiment of the present disclosure, bacterial spores captured from a solid surface using cotton swabs can be transferred onto a test surface by resuspending the spores on the cotton swab into water, and then plating the water suspension onto a test surface. The bacterial spores on the cotton swab could also be suspended into water followed by filtration of the water suspension through a membrane water filter. The spores embedded onto the membrane water filter are then streaked onto a test surface (Example 1, 2). Alternatively, the membrane filter is the test surface. Alternatively, the swab is not made of cotton, but is made of any suitable material.

Examples of adhesive polymers include but are not limited to: polydimethyl siloxane (PDMS). Alternatively, agar can be doped with PDMS. Similarly, agarose can be doped with PDMS. In a preferred embodiment the adhesive polymer, PDMS, used to capture the bacterial spores from the surface of origin is subsequently used as a test surface. PDMS has low chemical activity, it is hydrophobic, it is optically transparent above 250 nm and it is impermeable to water. With these characteristics, a PDMS test surface allows for induction of lanthanide-DPA/DP luminescence, detection and quantifying. In one embodiment of the present disclosure, the test surface is optically transparent greater than 250 nm. In an alternative embodiment, the test surface is partially transparent.

In a second embodiment, the spores captured from a place of origin using an adhesive polymer are subsequently transferred from the adhesive polymer. This transfer can be carried out using several methods easily envisioned by one skilled in the art. For example, the spores can be streaked onto a test surface (Example 2).

In another embodiment, spores captured on a membrane filter are transferred onto a test surface by a streaking method as disclosed in Example 2. Alternatively, spores on a membrane filter are physically lysed on the membrane filter and then pressed against an adhesive polymer such as PDMS containing lanthanide ions. Similarly, spores on a membrane filter can be streaked (Example 2) onto an adhesive polymer (e.g. PDMS, agar, agarose) that contains lanthanide ions and L-alanine for induction of germination. In another embodiment, the membrane filter embedded with the bacterial spores is used as the test surface.

The step of collecting bacterial spores carried in the air comprises capturing the bacterial spores with an aerosol sampler or impactor. Preferably, the step of collecting bacterial spores carried in the air comprises continuously sampling the air. In one embodiment, air is passed over quartz filter tape using an air sampler (Example 1). In one aspect the quartz filter tape is subsequently as the test surface. Alternatively, air can be passed over lanthanide-doped agar using an air sampler.

The agents used for germination and the agents used for lysing can be added to the test surface before or after the bacterial spores have been transferred onto the test surface.
Alternatively, the agents used for germination the agents used for lysing can be added in a mixture with the transfer of the bacterial spores. Examples of a germinating agent include but are not limited to: L-alanine, L-asparagine and D-glucose. Examples of lysing methods include but are not limited to: microwaving, plasma cleaning, dry heating, autoclaving, sonicating and hydrogen chloride gassing.

When the step of releasing DPA from the bacterial spores comprises microwaving the bacterial spores to heat the solution, the step of combining the lanthanide ions with the DPA in solution comprises cooling the heated solution to increase the fraction of bound lanthanide-DPA complex. One of skill in the art can envision several methods to prepare ("dope") the test surface for germination. Likewise, one of skill in the art can envision several methods to prepare ("dope") the test surface for lysing.

Lanthanide ions can be added to the test surface before the bacterial spores have been transferred onto said test surface, after the bacterial spores have been transferred onto said test surface, or in a mixture with the bacterial spores being transferred to the test surface. Lanthanide ions can be added before, after or in conjunction with the induced release of DPA/DP from the bacterial spores. Examples of lanthanide ions include, but are not limited to: terbium (Tb⁺), europium (Eu⁺) and dysprosium. In a preferred embodiment terbium (Tb⁺) ions are used.

Inducing the Lanthanide-DPA/DP Luminescence

A lanthanide ion-DPA/DP luminescence assay can be employed to detect individual bacterial spores on surfaces. For example, the lanthanide-DPA/DP luminescence assay can be combined with an optically transparent, adhesive polymer (PDMS, agar or agarose) to collect bacterial spores from surfaces to be tested. Once the bacterial spores are located on the test surface, they can be induced to release their DPA/DP content by germination (e.g. using L-alanine) or physical lysis, for example by autoclaving or microwaving. The highly concentrated DPA/DP from the spores spills into the surrounding area, generating a high concentration region around the spore body. The reagents used for detection and induction of germination, if that is the chosen method for DPA/DP release, can be added into the matrix before or after the spores are sampled. The lanthanide-DPA/DP luminescence arising from the region around the spore body is then imaged onto a camera. The bacterial spore regions manifest themselves as bright spots that can be counted. Due to the long-lived excited states of luminescent lanthanides, lifetime-gated detection enables any fluorescent background from interferences to be eliminated. Lifetime gating drastically reduces the background and enables much greater contrast between the lanthanide-DPA/DP luminescence regions and the background.

It is understood by one skilled in the art, that upon release of DPA and/or DP outside the bacterial spore, the DPA and/or DP molecules can interact with other substances in its environment, resulting in a derivative of DPA or DP.

The step of detecting the luminescence to determine the presence of the bacterial spores comprises monitoring the luminescence with a spectrometer or fluorimeter, and the step of detecting the luminescence to determine the presence of the bacterial spores comprises continuously monitoring the luminescence.

In one embodiment of the present invention, an adhesive polymer for the terbium-DPA/DP luminescence assay for bacterial spores on surfaces is polydimethyl siloxane (PDMS) doped with TbCl₃ and L-alanine. The L-alanine induces germination to release the DPA/DP from the core of the spore to the immediate surroundings. The TbCl₃ binds the DPA/DP, which triggers green luminescence (543.5 nm) under UV excitation (250-300 nm) that can be quantified with a photodetector. Individual germinating spores can be imaged within a microscope field of view using a lifetime-gated camera.

From the perspective of sensor design, the bacterial spore is essentially a 1 µm sphere containing about 10⁶ molecules of DPA. In previous experiments (U.S. patent No. 2004-0014154), spores were collected from surfaces using the standard cotton swabbing method, resuspended into water, and DPA/TbCl₃ was then released into a bulk solution by germination or physical lysing and a subsequent lanthanide (Tb⁺)-DPA luminescence assay was performed. This approach led to very dilute DPA solutions (e.g., 1 spore per ml of solution yields [DPA]=1 µM), which ultimately limits the sensitivity. As disclosed in the present invention, spores collected using the cotton swab can be suspended into water, and the water suspension can then be plated onto a testing surface for subsequent DPA/TbCl₃ release, lanthanide-DPA/DP complexing, excitation, luminescence and quantification. Alternatively, the water suspension can be filtered through a membrane filter and the spores on the filter can be streaked onto a testing surface.

The traditional culture-based assays require 3 days for colonies to grow and be counted. This traditional culture-based assay, also known as the NASA standard assay, is reported in colony forming units (CFU), since the quantification is based on the number of colonies. However, a significant fraction of bacterial spores can undergo stage-1 germination, during which DPA (i.e., the chemical marker that is unique to bacterial spores) is released, in less than 4 minutes. This type of quantification, is reported as germinating spore units (GSU). Experimental results shown herein (Table 1) show a comparison of the GSU calculated following the teachings disclosed in this application, versus the CFU calculation of the NASA standard assay for the same amount of starting spores (total spore units/TSU). FIG. 3 further shows an L-alanine induced germination of Bacillus subtilis spores on a TbCl₃ doped agar. The DPA/TbCl₃ released upon germination luminesces when complexed with the Tb⁺ ions. (Example 2).

Detection, Imaging and Quantification of Lanthanide-DPA/DP Luminescence

A salient feature of the present disclosure is the implementation of lifetime-gated imaging to obtain an image with good contrast of bacterial spores after germination and/or lysis. Fluorescence lifetime imaging uses special detectors and light source technology to generate images wherein the contrast is related to the fluorescence lifetime across a sample. Lifetime gating takes advantage of the fact that lanthanide ion (e.g. terbium) luminescence lifetimes are on the order of milliseconds, while fluorescence lifetimes from impurities generally are on the order of nanoseconds. Lifetime gating drastically reduces the chance of false negatives, which could arise if the lanthanide ion luminescence is masked by background fluorescence from impurities.

More specifically, the imaging method takes advantage of the fact that a bacterial spore is essentially a 1 µm diameter bag comprising 10⁶ molecules of DPA and/or DP. Releasing DPA/DP by thermal lysis or germination in the presence of lanthanide ions generates local high lanthanide-DPA/DP concentrations (in the millimolar range) with correspondingly high luminescence intensities. When the luminescence bands "halo" surrounding the spore body is imaged into individual lifetime-gated CCD detector elements, individual spores will be easily counted. Even when spores are clustered together,
the spore counts per cluster will be proportional to the intensity arising from a cluster. Thus, the resultant “bright spots” or “halos” are counted and the number of spores per bright spot is estimated by the luminescence of the spot (i.e., the spot intensity). The lifetime gating allows imaging of the long-lived lanthanide-DPA/DP excited state in the presence of short-lived fluorescence interferences (impurities, etc.).

Under UV (blacklight) illumination, the luminescence of the embedded Tb^{3+} ions increased dramatically upon germination within 40 minutes of the bacterial spores, while the embedded Tb^{3+} luminescence in the control sample that had no exposure to L-alanine remained weak (FIG. 3). An agar control sample without Tb^{3+} that was covered with bacterial spores also did not yield detectable luminescence. Note that the bright edges of the spots are artifacts of drying due to refraction from accumulated material, which would not appear in a lifetime-gated image.

An example of imaged Tb-DPA/DP complex representing spores on a PDMS test surface containing Tb^{3+} ions, which were subsequently lysed using plasma cleaning are shown in FIG. 4. Those spores that were not subject to plasma cleaning, and thus did not lyse and release DPA/DP, did not exhibit fluorescence (panel C and D of FIG. 4).

The pictures in FIG. 3 were taken without magnification, and thus the individual spores cannot be enumerated as they germinate. However, the present disclosure provides germinating bacterial spores imaged with a lifetime-gated microscope (FIG. 5, Example 3). As the spores germinate, DPA is released from the core to generate high, localized DPA/DP concentrations, which show up as bright green luminescent halos surrounding the spore body. These results demonstrate that viable bacterial spores on surfaces can be enumerated (quantified) according to methods of the present invention. In another embodiment, viable and nonviable bacterial spores on surfaces are enumerated according to the method of the present invention. A further embodiment, viable and nonviable spores on surfaces are enumerated according to the disclosed device of the present invention.

FIG. 6 shows lifetime-gated images of Eu^{3+} microspheres on highly fluorescent paper obtained with an ImageX-TGi lifetime-gated CCD camera mounted on a Carl Zeiss fluorescence microscope with 40x objective, excited with a 300 Hz Perkin elmer flashlamp (Example 3). Eu^{3+} microspheres were employed because they are commercially available and have analogous photophysical properties. The ImageX system effectively rejected all of the strong background fluorescence when a delay time of 100 µs was used. The present invention allows for microspheres exhibiting weak, long-lived luminescence immobilized on a highly fluorescent matrix to be imaged with high contrast against a silent background when gating is applied.

Another example of the invention is illustrated in FIG. 7. Bacterial spores were added onto the surface of R2A agar doped with 10 mM L-alanine (Example 1) to induce germination and 100 mM TdCl, to generate bright luminescent spots around the spore body as they germinated and released DPA/DP. A Xe-flash lamp firing at 300 Hz with a 275 nm interference filter provided excitation for the Tb-DPA complex, and the corresponding bright spots from the bacterial spore Tb-DPA luminescent halos were imaged with a lifetime-gated camera set at a delay time of 100 µs and an integration time of 2 ms. The individual bacterial spores become clearly visible as countable spots after germination. The images shown in FIG. 7 can be obtained by an apparatus as shown in FIG. 5. The apparatus of FIG. 5 comprises: 1) an ultraviolet light radiation device 10 (e.g., a Xenon flash lamp); 2) a first elliptical lens 20 and a second elliptical lens 30; 3) the light radiation device 10 and the lenses 20, 30 (40 represents the space in between the lenses) can have a 45 degrees inclination with respect to a stage or test surface 50 where the bacterial spores are located. The distance between lens 30 and stage 50 can be one inch. The distance between light radiation device 10 and stage 50 can be two inches. The light radiation device 10 is adapted to excite a complex of one or more lanthanide ions and aromatic molecules and generate a characteristic luminescence of the complex; 4) a microscope 60 for detecting and quantifying bacterial spores exhibiting the luminescence of the complex. 5) A red bandpass filter 70, suitable for Eu^{3+}, can be connected with the microscope 60; 6) an imaging device 80 (e.g., a nanoCCD camera) connected with the microscope 60.

Quantifying Viable Bacterial Spores

Instead of diluting the DPA/DP into bulk solution, bacterial spores can be immobilized onto a test surface such as an adhesive polymer (e.g., PDMS, agar with PDMS, agarose with PDMS), and then induced to germinate or lyse on the polymer test surface to generate local high DPA/DP concentrations (i.e., DPA and/or DP remains in the immediate surroundings of the spore body). To obtain viable counts, germination is induced by doping L-alanine (or other germination inducing agents) into the polymer matrix; lanthanide ions (e.g. TbCl3) also doped into the polymer, allow for imaging and quantification of bacterial spores by triggering luminescence in the presence of DPA/DP. To obtain total counts, the bacterial spores immobilized on the polymer test surface containing lanthanide ion are physically lysed (e.g., by dry heating, microwave, sonication, plasma cleaning, hydrogen chloride gassing or autoclaving) and the subsequent fluorescence emitted upon excitation of the lanthanide-DPA/DP complex is imaged and quantified resulting in the total number of live and dead bacterial spores.

The present disclosure also provides a method and apparatus to measure the fraction of bacterial spores that remain viable or alive, hence a live/dead assay for bacterial spores. The method combines dipicolinic acid/dipicolinate-triggered lanthanide luminescence and DPA/DP release from (1) viable bacterial spore through germination, and (2) DPA/DP release subsequent to lysis of all viable and nonviable bacterial spores. The ratio of the results from steps (1) to the sum of steps (1) and (2) yield the fraction of bacterial spores that are alive.

In one embodiment of the present disclosure, a method is provided for quantifying the percentage of viable spores in a population mixture of viable and inviable spores. In a preferred embodiment, the method for quantifying the percent viable spores in a mixed population of viable and inviable spores comprises transferring bacterial spores from their place of origin onto a test surface containing lanthanide ions, inducing germination of DPA/DP from the transferred bacterial spores, exciting the lanthanide-DPA/DP complex with UV radiation, quantifying the luminescence associated with the lanthanide-DPA/DP of germination, subsequently lysing the non-germinated bacterial spores on the test surface, exciting the lysis-induced lanthanide-DPA/DP complex with UV radiation, and quantifying the luminescence associated with the lanthanide-DPA/DP of lysis. Using the same test surface for germination and subsequent lysis allows for an accurate calculation of the percent viable spores in any given mixed population of viable and non-viable spores. The ability to rapidly quantify the fraction of viable bacterial spores from various origins (e.g. solid surfaces, water and air) is an essential feature of the present invention.
The method and apparatus of the present disclosure provide the imaging of the spherical resolution of the high concentrating region of DPA (the "halo") around each spore body, which has been germinated or lysed. The present method makes it possible to detect and quantify extremely low concentrations of bacterial spores in very short time. The method and apparatus for bacterial spore detection and quantification according to the present disclosure results within minutes and requires approximately an hour for quantifying the percent viability of bacterial spores on surfaces.

Bacterial spore capture/transfer methods

Capture from water: (FIG. 2C) For capture of bacterial spores from water, quartz filter tape (Whatman) is used in combination with an air sampler (Bioscience International: SAS Super 100/180/360). The quartz filter tape is then suspended in water, and the water suspension is then plated onto the testing surface, or the water suspension is filtered through a membrane filter which is then streaked onto the testing surface. Alternatively, the quartz filter can be used as the test surface (See Table 1).

Bacterial spores: *Bacillus* (*Bacillus subtilis, Bacillus cereus, Bacillus atrophaeus* etc.) spores from American Type Culture Collection (ATCC) were used in the examples provided herein. Stock solutions of purified endospores were prepared according to methods well known in the art. Plating of suspended spores was carried out by methods well known in the art (W. Nicholson and P. Setlow, "Sporulation, germination and outgrowth," *Molecular biology methods for bacilli*, S. Cutting, Ed. Sussex, England: John Wiley and Sons, 1990, 391-450).

| EXAMPLE 2 |

Test Surface

"Streaking": Spores on the surface of a membrane filter are transferred to a test surface by contacting the two surfaces at one end and dragging across to the other end to effect the transfer of spores from a membrane onto a test surface. This process is referred to as "streaking". Alternatively spores on an adhesive polymer such as PDMS can also be streaked from the polymer onto a test surface.

Bacteria spores were immobilized onto a test sample surface of thin, flexible, clear, adhesive polymer polysiloxan (PDMS) (Dow Corning). PDMS was doped with L-alanine (Aldrich) to induce germination and generate local high concentration of DPA/DP. TbCl$_3$ (Aldrich) was also doped into the PDMS sample. The bacterial spores immobilized on the L-alanine and TbCl$_3$-containing polymer were physically lysed by microwave irradiation (Vaid and Bishop, 1998?), wherein DPA/DP was released and luminescence was turned on.

The test surface in FIG. 3 was prepared by adding 100 μl of R2A agar (doped with 1 mM TbCl$_3$ onto a quartz slide and allowing it to solidify. On top of the agar, 10 μl of 10$^8$ spores/ml *Bacillus subtilis* spores were added (i.e., 10$^7$ spores), followed by 10 μl of 1-mM L-alanine to induce germination.

The test surface in FIG. 4 shows four 1 mm thick flat pieces of PDMS inoculated with *B. subtilis* spores. The PDMS pieces shown in panel A and B were placed into a plasma cleaner for 30 minutes. The pieces shown in panel C and D were not. Each of the four pieces were then placed onto Tb$^{3+}$-doped agar. The two plasma cleaned pieces produced bright spots corresponding to DPA/DP released from the *B. subtilis* spores during the plasma cleaning which complexed with the Tb$^{3+}$ ions in the agar. The two non-lysed PDMS test pieces did not produce bright spots because the *B. subtilis* spores on these pieces were not induced to release DPA/DP.

| EXAMPLE 3 |

Detection and Quantifying Spores

An apparatus for detecting and quantifying bacterial spores on a surface including lanthanide ions and aromatic molecules released from the bacterial spores on the surface. The apparatus in FIG. 5 comprises a UV-light radiation device for exciting a complex of a Tb$^{3+}$ ion and DPA/DP to generate a characteristic luminescence of the complex on a surface. The
source for the UV-light was a Xenon flash lamp, which was approximately 5 cm away the test surface. Between the Xenon flash lamp and the test surface were two C-amount elliptical lenses. The Xenon flash lamp and the test substrate were positioned at an angle of 45 degrees to each other. The area of irradiation by the Xenon flash lamp was observed by a microscope objective with a red bandpass filter suitable for Eu³⁺ for detecting and quantifying bacterial spores exhibiting the luminescence of the complex on the test surface. The image was transferred from the microscope to the imaging device for imaging bacterial spores exhibiting the luminescence, using an imageX nanoCCD camera (Photonic Research Systems Ltd, United Kingdom). The pixel size on the camera is 11.6 microns horizontal by 11.2 microns vertical and the camera has a chip with 752×582 pixels on a 10.25 mm×8.5 mm vertical area. Lifetime gated images were captured with a 100-µs delay integrating for 2 milliseconds. 6 to 13 images were taken over different areas of the medium. Each image captured an actual agarose area of 3.2 mm² at 40x magnification. The spatial resolution is a function of the camera, the microscope objective and the microscope camera port. The microscope image is projected onto the camera port that then determines the spatial resolution.

The invention claimed is:

1. A method for detecting and quantifying individual bacterial spores comprising:
   - capturing the bacterial spores;
   - transferring the bacterial spores to a test surface;
   - providing one or more lanthanide ions on the test surface;
   - releasing aromatic molecules from the bacterial spores on the test surface;
   - forming a complex of the one or more lanthanide ions and the aromatic molecules on the test surface;
   - exciting the complex to generate a characteristic luminescence of the complex on the test surface; and
   - detecting and quantifying the individual bacterial spores by imaging the bacterial spores via luminescence spots surrounding the bacterial spores, said luminescent spots generated on the test surface through lifetime-gated imaging, with a micrometric spatial resolution.

2. The method according to claim 1, wherein the capturing of the bacterial spores comprises transferring bacterial spores from a second surface by way of an adhesive polymer.

3. The method of claim 2, wherein the adhesive polymer is selected from the group consisting of PDMS, agar and agarose.

4. The method of claim 1, wherein the capturing of the bacterial spores comprises transferring bacterial spores from a second surface by way of a swab.

5. The method of claim 4, further comprising transferring the bacterial spores from the swab into water.

6. The method of claim 5 wherein the water is passed through a water filter.

7. The method of claim 1 wherein the capturing of the bacterial spores is from a place of origin selected from the group consisting of water and air.

8. The method of claim 7 wherein the transferring of the bacterial spores is from a place of origin to a filter selected from the group consisting of an air filter and a water filter.

9. The method of claim 1, wherein the test surface comprises at least one adhesive polymer selected from the group consisting of PDMS, agar, agarose, PDMS together with agar, and PDMS together with agarose.

10. The method of claim 1, wherein the test surface comprises at least one partially transparent adhesive polymer.

11. The method of claim 1, wherein the one or more lanthanide ions are provided to the test surface prior to the bacterial spores, after the bacterial spores, or as a mixture with the bacterial spores.

12. The method of claim 11, wherein the aromatic molecules are released from the bacterial spores by lysis of the bacterial spores.

13. The method of claim 1, wherein the aromatic molecules are selected from the group consisting of dipicolinic acid, dipicolinate, and dipicolinic acid together with dipicolinate.

14. The method of claim 1, wherein the aromatic molecules interact with the test surface resulting in a derivative of the aromatic molecules.

15. The method of claim 1, wherein the aromatic molecules are released from the bacterial spores by germination of the bacterial spores.

16. The method of claim 15, wherein the germination of the bacterial spores is induced by way of a germinating agent selected from the group consisting of L-alanine, L-asparagine and D-glucose.

17. The method of claim 1, wherein the aromatic molecules are released from the bacterial spores by lysis of the bacterial spores.

18. The method of claim 17, wherein the lysis of the bacterial spores is induced by way of a method of lysis selected from the group consisting of microwaving, autoclaving, sonication, plasma cleaning dry heating and hydrogen chloride gasing.

19. The method of claim 9 wherein the test surface further comprises at least one lanthanide ion and a germinating agent.

20. The method of claim 9 wherein the test surface further comprises at least one lanthanide ion and is subject to a method of lysis.

21. The method according to claim 1, wherein the complex is excited by UV-light.

22. The method of claim 2 wherein the adhesive polymer is the test surface.

23. The method of claim 3 wherein the adhesive polymer is the test surface.

24. The method of claim 8 wherein the filter is the test surface.

25. The method of claim 1, wherein the releasing of aromatic molecules from the bacterial spores on the test surface is by germination of the bacterial spores on the test surface; wherein the complex is a first complex; wherein the method further comprises:
   - releasing aromatic molecules from nongerminated spores on the test surface by lysis;
   - forming a second complex of the one or more lanthanide ions and lysis-released aromatic molecules on the test surface;
   - exciting the second complex to generate a characteristic luminescence of the second complex on the test surface; and
   - detecting and quantifying the bacterial spores by imaging the bacterial spores via the luminescence of the second complex on the test surface by lifetime-gated imaging.

26. The method of claim 25 wherein the test surface comprises terbium or europium ions;

27. A method of claim 26 wherein the quantifying of the bacterial spores represented by the generated luminescence
comprises counting the number of luminescent spots and the number of spores per luminescent spot can be estimated by the spot intensity.

28. An apparatus for detecting and quantifying individual bacterial spores according to the method of claim 1, comprising:

- a test surface, wherein the test surface comprises one or more adhesive polymers, one or more lanthanide ions, bacterial spores and aromatic molecule released from said bacterial spores
- an ultraviolet light radiation device adjacent to the test surface to excite a complex of lanthanide ions and aromatic molecules and generate a characteristic luminescence of the complex;
- an life-time gated imaging device for imaging the luminescence.

29. An apparatus of claim 28 wherein a microscope is connected to the life-time gated imaging device.

30. A method for detecting and quantifying bacterial spores comprising:

- providing the bacterial spores on a test surface, each bacterial spore having a spore body;
- releasing aromatic molecules from the spore body of a bacterial spore onto the test surface, the released aromatic molecules defining on the test surface a surrounding area around the spore body of the bacterial spore;
- forming a complex of the one or more lanthanide ions and the aromatic molecules concentration regions on the test surface by lysis the released aromatic molecules on said region, and detecting and quantifying the bacterial spores by imaging surrounding areas through lifetime-gated imaging with a micrometric spatial resolution.

31. The method of claim 30, wherein imaging surrounding areas is performed by imaging individual surrounding areas.

32. The method of claim 30, wherein imaging surrounding areas is performed by imaging surrounding areas of individual spores clustered together on the test surface.

33. The method of claim 30, wherein the bacterial spores are embedded in the test surface.

34. The method of claim 30, wherein the test surface is an adhesive polymer.

35. The method of claim 34 wherein the adhesive polymer is optically transparent.

36. The method of claim 30, wherein the releasing of aromatic molecules from the bacterial spores on the test surface is by germination of the bacterial spores on the test surface; and wherein the complex is a first complex;

wherein the method further comprises:

- releasing aromatic molecules from nongerminated spores on the test surface by lysis the released aromatic molecules defining on the test surface a surrounding area around the spore body;
- forming a complex of the one or more lanthanide ions and the aromatic molecules on the surrounding area;
- exciting the complex to generate a characteristic luminescence of the complex on the surrounding area; and

detecting and quantifying the bacterial spores by imaging surrounding areas through lifetime-gated imaging.

37. A method for detecting and quantifying bacterial spores, the method comprising:

- providing the bacterial spores, each bacterial spore having a spore body;
- providing one or more lanthanide ions;
- releasing aromatic molecules from the spore body forming a complex of the one or more lanthanide ions and the aromatic molecules;
- exciting the complex to generate a characteristic luminescence of the complex; and
- detecting and quantifying bacterial spores by imaging the bacterial spores through lifetime-gated imaging,

wherein releasing aromatic molecules from the spore body is performed on a test surface to generate on the test surface an aromatic molecules concentration region around the spore body

the complex of the one or more lanthanide ions and the aromatic molecules is formed on said region, and detecting and quantifying bacterial spores is performed by imaging the aromatic molecules concentration regions on the test surface with a micrometric spatial resolution.

38. The method of claim 37, wherein the imaged aromatic molecules concentration regions are individual aromatic molecules concentration regions.

39. The method of claim 37, wherein the imaged aromatic molecules concentration regions are aromatic molecules concentration regions of individual bacterial spores clustered together on the test surface.

40. The method of claim 37, wherein the bacterial spores are embedded in the test surface.

41. The method of claim 37, wherein the test surface is an adhesive polymer.

42. The method of claim 41, wherein the adhesive polymer is optically transparent.

43. The method of claim 37, wherein the releasing of aromatic molecules from the bacterial spores on the test surface is by germination of the bacterial spores on the test surface; and wherein the complex is a first complex;

wherein the method further comprises:

- releasing aromatic molecules from nongerminated spores on the test surface by lysis to generate on the test surface aromatic molecules concentration regions around each spore body;
- forming a second complex of the one or more lanthanide ions and lysis-released aromatic molecules on said concentration regions;
- exciting the second complex to generate a characteristic luminescence of the second complex on the concentration regions; and
- detecting and quantifying the bacterial spores by imaging the aromatic molecules concentration regions on the test surface.