NRT Quick Reference Guide: 
Brucella Species (Causes the disease Brucellosis)

For reference, please see "Key References Cited/Used in National Response Team (NRT) Quick Reference Guides (QRGs) for Bacterial 2011 Revision."

QRGs are intended for Federal On-Scene Coordinators (OSCs) and Remedial Program Managers (RPMs).

Agent Characteristics

Agent Classification: Biological Type: Bacteria (Four Species: B. melitensis, B. abortus, B. suis, B. canis)

Description: Brucellosis is a systemic, zoonotic disease, & transferable between different animal species and is caused by one of the 4 species of bacteria listed above. Virulence in humans decreases in the species order provided. They are small aerobic, non-motive bacteria. They reside in tissue & bone marrow, & are extremely difficult to eradicate, even with antibiotic therapy. In the US, the incidence of brucellosis is less than 0.5 cases/100,000 people (mostly with B. melitensis). Brucellosis is endemic in wildlife populations in WY & MT and in sheep, goats, cattle, bison, elk & several other animals in WY, MT, ND, SD. Brucellosis is also found sporadically in feral swine in the southeastern US (FL, AL, GA, TX). Most outbreaks are seen in wild ungulates (e.g., deer, elk, moose, bison). Humans can be exposed during the birthing & slaughtering of an animal, when handling animal viscera, and eating/drinking unpasteurized milk, cheese, ice cream, etc. Humans can also be exposed via inhalation of infectious aerosols or if mucous membranes are exposed to the agent. When responding to human cases, it should be determined if it is from a natural outbreak or a bio-terrorist event. If a natural event, infected herds may be culled. Note: This QRG focuses on intentional (bio-terrorism) release scenarios, but since brucellosis outbreaks occur naturally the natural exposure routes are also described.

Bio-Safety Level: 3
CDC Class: B
HHS/USDA Select Agent: Yes, excluding B. canis
Incubation Period: 5-60 days.
Duration of Illness: Weeks to years
Person-to-Person Transmission: Rare; from sexual contact or via breastfeeding

Other Forms of Transmission: Treatments: Treatment is difficult but some antibiotics have been shown to be effective.

Incubation Period: 5-60 days

Health Effects

CAUTION: REAEROSONIZATION IS A CONCERN FOR ALL RELEASE SCENARIOS

Air: If weaponized, Brucella is an inhalation threat to humans & wild & domestic animals. The area of initial release might be difficult to identify because symptoms may take days to appear.

Soil: Brucella is persisting in soil for up to 125 days. Decon precautions should be taken.

Water: Brucella is a probable water threat because the bacteria are stable for 20-72 days

Other: Brucella spp. are naturally occurring & endemic in the United States. Natural occurring exposure includes contact with infected animals or contaminated animal products; these includes eating contaminated meat products.

Release Scenarios

Onset Symptoms may occur 5-60 days after exposure.

Signs/Symptoms per Exposure Route

General: Brucellosis is a systemic bacterial disease with acute or gradual onset in the form of flu-like symptoms, with intermittent fevers, making it difficult to conclusively diagnose.

Inhalation: Unless it is weaponized, transmission of Brucella via inhalation is not a common route but it can be for laboratory, slaughterhouse, & large animal veterinarian occupations.

Skin: Infection is possible through abraded skin. Persons who work in slaughterhouses, meat-packing plants, hunters, veterinarians, etc. are at higher risk of exposure.

Ingestion: Infection is possible by eating/drinking contaminated, often unpasteurized, milk or dairy products.

Effect Levels

Infectivity: Brucella has high infectivity.
Infective Dose: Currently, there is no infective dose listed for Brucella spp. but it is estimated that inhaling only 10-100 weaponized bacteria is sufficient to cause human disease.
Lethality: Brucella has low lethality (0.5 - 6%).

Concerns

Check with the Health & Safety Officer regarding PPE, Medical Surveillance, & Health & Safety Plan (HASP). Level of PPE may vary depending upon the incident & site specific circumstances. The PPE Levels listed are general suggestions only & are appropriate only for Brucella; they may not provide protection for some decon & other chemicals that workers may be exposed to during response/recovery operations. For decon of workers, use warm soapy water, taking care to avoid abrading the skin.

Medical

Baseline: Annual physical & respiratory function exams. THERE IS NO FDA APPROVED VACCINE AGAINST BRUCELLOSIS.

Treatments Available: Treatment post exposure is supportive & is accompanied with Ciprofloxacin & Doxycycline antibiotics. Prophylactic use of antibiotics may be recommended.

First Aid

During Incident: Conduct medical monitoring; use PPE as designated by the HASP; record the PPE Levels used; monitor for fever & other signs/symptoms as listed under Health Effects & if necessary, ensure medical attention is obtained as soon as possible.

Post Incident: Monitor for signs/symptoms & if necessary, ensure medical attention is provided as soon as possible.

PPE

Emergency Response to a Suspected Biological Incident: Possible PPE Levels for emergency responders is based on scenario risks from highest level of protection to least: 1) Pressure-demand Self Contained Breathing Apparatus (SCBA) with Level A protective suit, when: a) Event is uncontrolled, b) The type(s) of airborne agent(s) is unknown, c) The dissemination method is unknown, d) Dissemination via an aerosol-generating device is still occurring, e) Dissemination via an aerosol-generating device has stopped, but there is no information on the duration of dissemination, or what the exposure concentration may be. 2) Pressure-demand SCBA with Level B protective suit, when: a) The suspected biological aerosol is no longer being released, b) Other conditions may present a splash hazard. 3) Full-facepiece respirator with P100 filter or PAPR with HEPA filters, when: An aerosol-generating device was not used to create high airborne concentrations. 4) Disposable hooded coveralls, gloves, & foot coverings, when: Dissemination was by a liquid, package, or material that can be bagged, contained, etc.

Other Workers: PPE recommendations for workers other than emergency responders must be developed in the HASP for the specific scenario. PPE recommendations will vary by job type (e.g., cleanup, decon, etc.), type of exposure (e.g., airborne or surface/liquid/solid soil), & any other site hazards (e.g., chemical, physical, etc.).

Field Detection

Fixed Aerosol Monitoring: An aerosol release of Brucella species may be detected using air samples & PCR. Results may be delayed as much as 2 days from time of release. In the absence of reliable detection, a Brucella release will only be confirmed once patients present with symptoms and are diagnosed or animal die-off from brucellosis is confirmed. Consult EPA/HQ-EOC at 202-564-3850 for more information.

Portable Aerosol Monitoring: Portable aerosol monitoring may use dry or wet sampling methods. Dry sampling (useful only for molecular analyses) includes gelatin, cellulose acetate & Teflon filters. Wet sampling methods include impingers (low flow) & impactors (single or six stage). Refer to the manufacturer’s aseptic sampling methods, flow rates, & sampling times. Ensure that the appropriate pump is used for the selected sampling method.

Sampling

Concerns: BEFORE OBTAINING SAMPLES: Identify sample transportation requirements; Contact EPA/HQ-EOC (202-564-3850) for ERLN contract laboratories able to analyze these types of samples; Clearly identify & coordinate with the laboratory to be used since most labs cannot analyze all types of media (e.g., wipes, swabs, and HEPA vacuum samples); Coordinate with the sample disposal facility for acceptance criteria (i.e., sample decon requirements); Coordinate with investigative units (EPA-CID & FBI) to ensure sample chain-of-custody is maintained between the groups. Note: Detection/analytical equipment & sampling techniques will be highly site-specific & depend on: 1) the characteristics of the agent; 2) the type of contaminated surfaces (e.g., porous v. nonporous); 3) the phases/purposes of sampling (initial ID v. post-decon sampling); 4) the way in which samples are handled so as not to adversely affect viability; 5) transportation regulations 6) the acceptance criteria of the analytical laboratory & 7) the sample decon requirements for the waste disposal facilities to be used. 

See LABORATORY ANALYSIS, below.

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CAUTION: ONLY MANUFACTURER CERTIFIED HEPA VACUUM EQUIPMENT SHOULD BE USED. A site-specific sampling plan should be reviewed & approved by appropriate Subject Matter Experts &/or through ICS channels.

Sample Location Plans: If release was limited to a small area due to opening a letter or container, start with an area thought to be free of contamination & work in circular patterns towards the initial point of contamination. Be concerned about other contaminated areas due to foot traffic/ventilation systems (elevator buttons, mail, corners of hallways, baseboards, light switches, door knobs, etc.). Based on site characteristics & laboratory capacity, the sample plan may be judgmental, probabilistic, or a combination thereof.

Consult EPA/HQ-EOC at 202-564-3850 for Environmental Response Laboratory Network (a.k.a. ERLN laboratory) contact information for personnel who can explain/describe the sampling procedure most compatible with their current analytical procedure.

Types of Samples: Air, water, soil, surfaces, dairy production/livestock

Note: While Brucella DNA can be detected long after the bacteria have perished & might be of forensic interest, the presence of the DNA says little about the potential human risk in the days following a release.

Air: Collect air samples with a gel filter, impinger or impactor. Refer to the manufacturer's aseptic sampling methods, flow rates, & sampling times. Ensure that the appropriate pump is used for the selected sampling method.

Water: Since Brucella can persist in water, any potable water source should be sampled. If the potable water is chlorinated, the chlorine needs to be neutralized immediately with a sodium thiosulfate or other neutralizer at the concentration specified by the analytical laboratory prior to shipment. As chlorine levels can vary substantially throughout a drinking water system, it is not always appropriate to assume that a sample is chlorinated based solely on the description of the water treatment processes in use.

Soil: For the localized areas where soil deposition of the agent is suspected (i.e., aerosol or liquid droplets), a surface soil sample from a depth of less than 1 inch (2.5 cm) should be obtained from a non-vegetated area.

Surfaces: 1) Wipe & Swab Sampling (for non-porous surfaces): Sterile macrofoam swabs moistened with 1X phosphate-buffered saline supplemented with 0.01% Tween-20 (PBST). If this solution is not available, use sterile de-ionized water (DI). Do NOT use dry wipes or swabs. 2) HEPA Vacuum Sampling (for both porous & non-porous surfaces): collect samples in a HEPA sock designed to fit into an inlet nozzle of a manufacturer certified HEPA vacuum cleaner. Good for screening & determining the extent & location of contamination in large areas.

Agriculture & Wildlife: Upon confirmation of an outbreak, ensure that’s agencies are notified immediately since brucella is a zoonotic vector borne disease; USDA at 202-720-5711 & National Center for Emerging and Zoonotic Infectious Diseases at 800-232-4636 (after hours call the Directors Emergency Operations Center at 770-498-7100).

Samples that test for Re-aerosolization: 1) Wipe sampling of the air duct system (filters, areas of particulate deposition) if exposure occurred indoors. 2) Air Sampling & Single Stage Impactors with settle plates for capturing airborne particulates of respirable size (1-5 microns) on a series of agar plates. Agar plates are then sent to laboratory for culture analysis.

Sample Packaging & Shipping: The packaging & shipping of samples are subject to strict regulations established by DOT, CDC, USPS, OSHA, & IATA. Contact the sample-receiving laboratory to determine if they have additional packaging, shipping or labeling requirements (e.g., DO NOT X-RAY). Samples should be packaged in an air-tight container & kept at temperatures of 40-50°F (4-10°C). Ensure samples are not placed directly on the ice used for cooling the shipping container.

CAUTION: Many labs may not be able to perform analysis on all matrices (e.g., wipes & soil). The goal of laboratory analysis for environmental sampling purposes is to determine if viable Brucella is present in the sample. Note: The selected laboratory may use a tiered approach. If a tiered approach is used, the initial analysis may only determine if select/particular components of the bacterium are present in the sample (e.g., presence or absence). It may take additional time (up to weeks depending on the laboratory) to determine if the bacterium are viable & still able to cause adverse effects.

Laboratory Information: Contact EPA/HQ-EOC (202-564-3850) for ERLN contract laboratories able to analyze these types of samples.

Decontamination/Cleanup

CAUTION: DECON SOLUTIONS SHOULD NOT BE DEPLOYED AS A SPRAY WHENEVER POSSIBLE.

Decon Planning: Site-specific decon/cleanup plan should be developed & approved by all necessary organizations/SMES via ICS channels. Responders should develop a plan that takes into account: 1) Nature of contamination including purity, physical properties, how it entered the facility, etc.; 2) Extent of contamination; 3) the amount & possible pathways that have spread the agent. It is advisable to isolate the contaminated area; & 3) Objectives of decon, including decon of critical items for re-use & the treatment, removal, or packaging of other items for disposal. Note: Crisis exemptions from EPA’s Office of Pesticide Programs might be necessary depending on decontaminating agents used.

CAUTION: ONLY MANUFACTURER CERTIFIED HEPA VACUUM EQUIPMENT SHOULD BE USED.

Decon Methods: Decon decisions will be site & situation specific but due to re-aerosolization concerns, under NO circumstances should a non-HEPA vacuum cleaner or a broom be used. EPA’s National Decon Team (800-329-1841) can provide specific decontamination parameters & requirements for using readily available commercial items such as household bleach.

Methods used on surfaces: 1) Source reduction steps, including HEPA vacuuming; 2) Liquid antimicrobial products such as pH-amended bleach (mixture of 1 part household bleach (5.25% to 6.0%) to 1 part white vinegar to 8 parts water, is recommended). This product affects surfaces differently in terms of corrosiveness, staining, & residue. The product will be most efficient a) at higher temperatures (i.e., >70°F or 21°C) b) when plain bleach (e.g., no added fragrance) is used to make the pH-amended bleach solution, c) when pH is < 7, d) when presence of other surface contaminants is minimal, & e) when surfaces remain wet with amended bleach solution for 60 minutes. Note: Store-bought bleach does degrade with time – check the expiration date. Alternate antimicrobial products include: chlorine dioxide, hydrogen peroxide, & peroxyacetic acid. Fumigation: Uses gas or vapor to decontaminate facilities in which there is evidence of high levels of contamination, re-aerosolization, or if decontamination of limited access areas is required (e.g., HVAC systems). Fumigants: chlorine dioxide, & vaporized hydrogen peroxide. Prior to use, the fumigant’s compatibility with materials, penetration capacity, method of removal at the end of fumigation, as well as it’s physical, chemical, & toxicological properties should be taken into account. Each chemical has a specified range for process variables (e.g., temperature, relative humidity, conc. & contact time) that must be followed. Other Decon: 1) Ethylene oxide sterilization is used to decontaminate items in an off-site sterilization chamber. 2) Irradiation uses cobalt-60 & electron beam technologies to destroy agents at off-site locations. This procedure may destroy magnetic media. Irradiation & chemical sterilization may be useful in decontaminating items that are intended to be returned to owners.

Verification of Decon: Site & situation specific. Please contact ERT (732-321-6660) and NDT (800-329-1841) for further assistance.

CAUTION: Hazardous waste transportation & disposal are regulated federally; however more stringent regulations may exist under state authority. These regulations differ from state-to-state. Detailed state regulations can be found at www.envcap.org.

Waste Disposal Planning: Waste generated from assessment & cleanup activities should be autoclaved, chemically disinfected, or fumigated & then tested to be sure the agent(s) were inactivated. Waste disposal for agent-contaminated wastes generated from decontamination & disposal activities will be problematic. Landfills willing to take these wastes may be limited & incineration may be prohibitively expensive or impractical. All waste disposal options should be investigated as early as possible in the response process as possible. Transportation of the agent contaminated wastes from the site to the landfill or incinerator may be problematic as well. Agreements must be reached between the waste generator & acceptor BEFORE transport. Information regarding the agreements may be required to be available to the public. Transportation of hazardous waste may cross several states and localities, which may exceed federal regulations. Requirements for transporting hazardous materials, & procedures for exemption, are specified in http://www.fmcas.dot.gov/safety-security/hazmat/comply/yrregs.htm#hmp. The EPA has developed a web-based Incident Waste Management Planning & Response Tool which contains guidance related to waste transportation, contact information for potential treatment, disposal facilities, & state regulatory offices, packaging guidance to minimize risk to workers, & guidance to minimize the potential for contaminating the treatment or disposal facility. Access to the EPA’s web based disposal tool requires preregistration: http://www2.ergweb.com/bdtool/login.asp.