SANI-CLOTH® BLEACH
GERMICIDAL DISPOSABLE WIPE

Technical Data Bulletin
PRODUCT DESCRIPTION

EPA-Registered Sani-Cloth® Bleach is a pre-moistened wipe designed to kill the most clinically relevant pathogens in healthcare. The stabilized 1:10 dilution of sodium hypochlorite is now tested effective against 50 microorganisms, including 14 Multi-Drug Resistant Organisms (MDROs) with an overall contact time of four minutes. Ideal for disinfecting hard, non-porous surfaces in high risk areas endemic with Multi-Drug Resistant Organisms and Clostridium difficile spores.

CHEMICAL COMPOSITION

<table>
<thead>
<tr>
<th>Active Ingredient</th>
<th>Sodium Hypochlorite</th>
<th>Other Ingredients (Does not include the weight of the wipe)</th>
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<tbody>
<tr>
<td></td>
<td>0.63%</td>
<td>99.37%</td>
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<tr>
<td>TOTAL</td>
<td>100.00%</td>
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EFFICACY

CLOSTRIDIUM DIFFICILE EFFICACY

**CLOSTRIDIUM DIFFICILE SPORES:** Clostridium difficile spores [ATCC 43598]

Test Method Used: Modified ASTM E 2197-02, Standard Quantitative Disk Carrier II Test Method for Determining the Bactericidal, Virucidal, Fungicidal, Mycobactericidal and Sporicidal Activities of Liquid Chemical Germicides, as specified by the U.S. EPA in Guidance for the Efficacy Evaluation of Products with Sporicidal Claims Against Clostridium difficile spores (February 5, 2009).

Organic Soil Load: None

Exposure Time: 4 minutes at 73.4° - 75.2°F

Incubation: 7 days at 82.4° - 89.6°F

Results: Met the performance criterion of a minimum reduction in viable spores of 6 Log10 for products with sporicidal claims against Clostridium difficile, in accordance with the U.S. EPA Guidance for the Efficacy Evaluation of Products with the Sporicidal Claims Against Clostridium difficile spores (February 5, 2009).

BACTERIAL ORGANISM EFFICACY

MULTI-DRUG RESISTANT BACTERIA: Acinetobacter baumannii Multi-Drug Resistant [ATCC 19606] [Effective against organisms resistant to Ampicillin, Cefazolin, Gentamicin, Trimethoprim/Sulfa and Intermediate resistance to Cefotaxime, Ceftriaxone and Piperacillin]

Enterobacter cloacae - NDM-1 positive [CDC 1000654]

Escherichia coli - NDM-1 positive [CDC 1001728]

ESBL Resistant Escherichia coli [ATCC BAA-196]

ESBL Resistant Klebsiella pneumoniae [ATCC 700603]

Klebsiella pneumoniae - Carbapenem Resistant [ATCC BAA-1705]

Klebsiella pneumoniae - NDM-1 positive [CDC 1000527]

Community Acquired Methicillin Resistant Staphylococcus aureus (CA-MRSA) [NARSA NRS384] [Genotype USA 300]

Community Acquired Methicillin Resistant Staphylococcus aureus (CA-MRSA) [NARSA NRS123] [Genotype USA 400]

Staphylococcus aureus Methicillin Resistant (MRSA) [ATCC 35592]

Streptococcus pneumoniae - Penicillin Resistant [ATCC 700677]

Vancomycin Intermediate Staphylococcus aureus (VISA) [HIP 5836]

Vancomycin Resistant Staphylococcus aureus (VRSA) [NARSA VRS1]

Vancomycin Resistant Enterococcus faecalis (VRE) [ATCC 51575]

Test Method Used: Modified AOAC Germicidal Spray Method for Hard Surface Disinfection

Organic Soil Load: 5% Fetal Bovine Serum

Exposure Time: 1 minute at 68° - 69.8°F

Incubation: 2 - 8 days at 95° - 98.6°F

Results: No growth observed
**TB:**

*TB:* *Mycobacterium bovis* - BCG (TB)

**Test Method Used:** Modified AOAC Method for Pre-Saturated Towelettes for Hard Surface Disinfection to Determine Tuberculocidal Effectiveness

**Organic Soil Load:** 5% Horse Serum

**Exposure Time:** 2 minutes at 68°F

**Incubation:** 90 days at 98.6°F

**Results:** No growth observed

**BACTERIA:**

*Bordetella pertussis* [ATCC 12743]

*Burkholderia cepacia* [ATCC 25416]

*Campylobacter jejuni* [ATCC 29428]

*Escherichia coli* [ATCC 11229]

*Escherichia coli O157:H7* [ATCC 35150]

*Klebsiella pneumoniae* [ATCC 4352]

*Legionella pneumophila* [ATCC 33153]

*Listeria monocytogenes* [ATCC 19117]

*Pseudomonas aeruginosa* [ATCC 15442]

*Salmonella enterica* [ATCC 10708]

*Serratia marcescens* [ATCC 14756]

*Staphylococcus aureus* [ATCC 6538]

*Streptococcus pyogenes* [ATCC 12344]

**Test Method Used:** Modified AOAC Germicidal Spray Method for Hard Surface Disinfection

**Organic Soil Load:** 5% Fetal Bovine Serum

**Exposure Time:** 1 minute at 68°F

**Incubation:** 2 - 5 days at 95° - 98.6°F

**Results:** No growth observed.

**VIRAL ORGANISM EFFICACY**

**ENVELOPED VIRUSES:**

Avian Influenza A H5N1 virus [Strain VN/551-PR8/CDC-RG CDC #2006719965]

Cytomegalovirus (CMV) [ATCC VR-538]

Herpes simplex virus type 2 [ATCC VR-734], Strain G

Human Coronavirus [ATCC VR-740], Strain 229E

Influenza A virus/Hong Kong Strain [ATCC VR-544]*

* Pandemic 2009 H1N1 Influenza A virus (Kill claim included)

Influenza B virus/Strain B/Hong Kong/5/72, [ATCC VR-823]

Respiratory syncytial virus (RSV) [ATCC VR-26], Strain Long

**NON-ENVELOPED VIRUSES:**

Adenovirus type 2 [ATCC VR-846], Adenoid 6 Strain

Canine Parvovirus [ATCC VR-2017], Cornell Strain

Hepatitis A virus (Human) (HAV) [Strain HM-175]

Norovirus (Feline Calicivirus) [ATCC VR-782]

Poliovirus type 1 [ATCC VR-1562], Chat Strain

Rhinovirus type 37 [ATCC VR-1147], Strain 151-1

Rotavirus [Strain WA]

**Test Method Used:** Tests were conducted according to U.S. Environmental Protection Agency guidelines in effect at the time for determining virucidal efficacy of disinfectants intended for use on dry inanimate surfaces.

**Organic Soil Load:** 5% Fetal Bovine Serum

**Exposure Time:** 1 minute at 68°F

**Results:** Virucidal according to the criteria established by the U.S. Environmental Protection Agency for registration and labeling of a disinfectant product as a virucide.
BLOODBORNE PATHOGENS:

Hepatitis B virus (HBV) - Duck HBV [Strain 7/31/07]
Hepatitis C virus (Human) (HCV) - Bovine Diarrhea virus [Strain Oregon C24-v-genotype 1]

Test Method Used:
Tests were conducted according to U.S. Environmental Protection Agency guidelines in effect at the time for determining virucidal efficacy of disinfectants intended for use on dry inanimate surfaces.

Organic Soil Load:
Hepatitis B virus (HBV) 100% Duck Serum
Hepatitis C virus (HCV) 5% Horse Serum

Exposure Time:
1 minute at 68°F

Results:
Virucidal against Hepatitis B and Hepatitis C viruses according to the criteria established by the U.S. Environmental Protection Agency for registration and labeling of a disinfectant product as a virucide.

HIV-1 (AIDS virus) [Strain HTLV-IIIg]

Test Method Used:
This test was conducted according to U.S. Environmental Protection Agency guidelines in effect at the time for determining virucidal efficacy of disinfectants intended for use on dry inanimate surfaces.

Organic Soil Load:
5% Fetal Bovine Serum

Exposure Time:
1 minute at 68°F

Results:
Virucidal against Human Immunodeficiency Virus Type 1 according to the criteria established by the U.S. Environmental Protection Agency for registration and labeling of a disinfectant product as a virucide.

PATHOGENIC FUNGI EFFICACY

YEAST ORGANISMS:

Candida albicans [ATCC 10231]
Aspergillus brasiliensis [ATCC 16404]
Trichophyton mentagrophytes [ATCC 9533]

Test Method Used:
Modified AOAC Germicidal Spray Method for Hard Surface Disinfection

Organic Soil Load:
5% Fetal Bovine Serum

Exposure Time:
4 minutes at 70° - 77°F

Incubation:
2 - 10 days at 77° - 86°F

Results:
No growth observed

TOXICITY

ACUTE ORAL TOXICITY OF SANI-CLOTH BLEACH®

Conclusion: A single dose of Sani-Cloth® Bleach solution was administered and observed for 14 days. No signs of toxicity were observed during the 14 day period of this study. Based on the results of this study, the acute oral toxicity LD50 of Sani-Cloth® Bleach was greater than 5 g/Kg of body weight.

ACUTE EYE IRRITATION OF SANI-CLOTH BLEACH®

Conclusion: One eye of each subject was instilled with the undiluted solution, while the contralateral eye remained untreated and served as a control. Under the conditions of the test, Sani-Cloth® Bleach produced eye irritation clearing in 7 days or less.

ACUTE DERMAL TOXICITY OF SANI-CLOTH BLEACH®

Conclusion: Following the single dermal administration, the subjects were observed for 14 days. Under the conditions of the test, the acute dermal LD50 of Sani-Cloth® Bleach was found to be greater than 5 g/Kg of body weight.

ACUTE SKIN IRRITATION OF SANI-CLOTH BLEACH®

Conclusion: This test was conducted according to U.S. Environmental Protection Agency guidelines in effect at the time of testing. The subjects were exposed to the undiluted solution for 4 hours. Under the conditions of the test, Sani-Cloth® Bleach produced minimal skin irritation.

SKIN SENSITIZATION OF SANI-CLOTH BLEACH®

Conclusion: This test was conducted according to U.S. Environmental Agency guidelines in effect at the time of testing to determine the potential for Sani-Cloth® Bleach to produce sensitization after repeated topical applications. Based on the results of this test, Sani-Cloth® Bleach would not be considered a dermal sensitizing agent.

ACUTE INHALATION TOXICITY OF SANI-CLOTH BLEACH®

Conclusion: This test was conducted according to U.S. Environmental Protection Agency guidelines in effect at the time of testing. The subjects were exposed to the aerosolized product for a four hour period. Based on the results of this study, the acute inhalation toxicity LC50 of Sani-Cloth® Bleach is greater than 2.35 mg/L of air.