

M. smegmatis ATCC 607 shares a number of morphological traits with M. Tuberculosis including the distinctive waxy cell wall that provides a robust resistance to chemical disinfectants and sanitizers

Results of the Study

Test Microorganism	Run Type	Treatment Time Point	Replicate	CFU/m³	Percent Reduction Compared to Time Zero	Log ₁₀ Reduction Compared to Time Zero	Adjusted Log ₁₀ Reduction ¹ Compared to Baseline
<i>M. Smegmatis</i> ATCC 607		Time Zero	Replicate 1	5.29E+07		N/A	
	Baseline	15 Minute	Replicate 1	4.41E+07	16.7%	0.08	N/A
		30 Minute	Replicate 1	4.51E+07	14.78%	0.07	N/A
		60 Minute	Replicate 1	3.42E+07	35.27%	0.19	N/A
		90 Minute	Replicate 1	2.55E+07	51.75%	0.32	N/A
	Test	Time Zero	Replicate 1	3.10E+08		N/A	
		15 Minute	Replicate 1	4.19E+05	99.86%	2.87	2.79
		30 Minute	Replicate 1	2.96E+05	99.905%	3.02	2.95
		60 Minute	Replicate 1	9.59E+03	99.997%	4.51	4.32
		90 Minute	Replicate 1	< 8.56E+01	> 99.99997%	> 6.56	> 6.24

The limit of detection for this assay is 8.00E+01 CFU/m³ and values below the limit of detection are noted as "<8.00E+01" in the data table. ¹The Log reductions for the Test Runs are adjusted to account for natural die-off and gravitational settling observed in the Control Run.

Mycobaderium smegmatis ATCC 607: -

this bacteria is an acid-fast, bacillus-shaped, aerobic microorganism that is commonly used a surrogate model for M. tuberculosis and is found in soil, plants, and water. M. Smegma/is is non-pathogenic to humans except in rare cases, and is considered saprophytic. Unlike other pathogenic Mycobacterium, M. Smegma/is isn't dependent on living in animals. M. smegma/is shares a number of morphological traits with M. Tuberculosis including the distinctive waxy cell wall that provides a robust resistance to chemical disinfectants and sanitizers. The quick growth rate of this microorganism is ideal for in-vitro testing, as other bacteria in this Genus may take several weeks to demonstrate growth. Due to the non-pathogenic nature of this organism, it is used as a M. Tuberculosis model for aerosol disinfection testing.

Page 10 of 11



STUDY REPORT

Study Title

Evaluation of Bioaerosols and Antimicrobial Efficacy of SAM Air System's Test Device

<u>Test Method</u>

Custom Aerosol Study

Study Identification Number NG16192

Study Sponsor

Cynthia Bell cyn370@comcast.net Scientific Air Management 1301 West Copans Road suite C6 Pompano Beach, Florida

Test Facility

Microchem Laboratory 1304 W. Industrial Blvd Round Rock, TX 78681 (512) 310-8378 Report Author: Samuel Hanley, B.S.



Purpose of the Study

The purpose of this study is to document the antimicrobial efficacy of SAM Air System's Test Device

Study Timeline

Devices Received	Cultures Initiated	Chamber Run	Nebulization Initiated and Treatment	Enumeration Plates Evaluated	Report Delivered
	Ba	seline: <i>Bacillus s</i>	ubtilis ATCC 196	59	
01 SEP 2020	05 NOV 2020	05 NOV 2020	05 NOV 2020	06 NOV 2020	20 NOV 2020
		Test: <i>Bacillus sub</i>	tilis ATCC 19659	2	
01 SEP 2020	10 NOV 2020	10 NOV 2020	10 NOV 2020	12 NOV 2020	20 NOV 2020
	Baselin	e: <i>Mycobacteriur</i>	<i>m smegmatis</i> ATC	CC 607	
01 SEP 2020	03 NOV 2020	13 NOV 2020	13 NOV 2020	16 NOV 2020	20 NOV 2020
	Test:	Mycobacterium :	smegmatis ATCC	607	
01 SEP 2020	12 NOV 2020	16 NOV 2020	16 NOV 2020	18 NOV 2020	20 NOV 2020

Page 2 of 11



Test Device Information

Name of Test Device: SAM S400 Manufacturer: SAM Air Systems



Note: The image above depict the test device, SAM S400, which was provided by the Study Sponsor for use in testing.

Page 3 of 11



Test Microorganism Information

The following test microorganisms were selected for this test:



Bacillus subtilis

This bacteria is Gram-positive, rod shaped, capable of forming endospores. Endospores of *Bacillus subtilis* can tolerate harsh environmental conditions such as UV exposure and high temperatures. Typically found in soil, this species is not known to cause disease in healthy individuals, but can be considered an opportunistic pathogen among the immuno-compromised. *Bacillus subtilis* endopores serve as one of the models for evaluating the effectiveness of sporicides and sterilants.

Mycobacterium smegmatis ATCC 607: This bacteria is an acid-fast, bacillus-shaped, aerobic microorganism that is commonly used a surrogate model for *M. tuberculosis* and is found in soil, plants, and water. *M. Smegmatis* is non-pathogenic to humans except in rare cases, and is considered saprophytic. Unlike other pathogenic *Mycobacterium*, *M. Smegmatis* isn't dependent on living in animals. *M. smegmatis* shares a number of morphological traits with *M. Tuberculosis* including the distinctive waxy cell wall that provides a robust resistance to chemical disinfectants and sanitizers. The quick growth rate of this microorganism is ideal for in-vitro testing, as other bacteria in this Genus may take several weeks to demonstrate growth. Due to the non-pathogenic nature of this organism, it is used as a *M. Tuberculosis* model for aerosol disinfection testing.



Criteria for Scientific Defensibility of a Custom Device Study

For Microchem Laboratory to consider a Device Study study to be scientifically defensible, the following criteria must be met:

- 1. The average number of viable bacteria, fungi, or bacteriophage recovered from the time zero samples should be approximately 1 x 10⁵ cells/m³.
- 2. Positive/Growth controls must demonstrate growth of the appropriate test microorganism.
- 3. Negative/Purity controls must demonstrate no growth of test microorganism.
- 4. The neutralization test suspension must be \geq 70% of that recorded for the neutralization control suspension count.

Passing Criteria

Because of the nature of the study, passing criteria may be determined by the Study Sponsor.

Testing Parameters used in this Study

Volume of inoculum added to Nebulizer	20.0 ml	Nebulization Time	60 minutes
Sampler Media (Volume)	Phosphate buffered saline (20.0 ml)	Neck Rinse Media (Volume)	Phosphate buffered saline (5.0 ml)
Sampling Time	10 minutes	Contact Times	Time zero 15 minutes 30 minutes 60 minutes 90 minutes
Sampling Type	Impingers, SKC biosamplers	Enumeration Media	BHIA (MS607) TSA (BS19569)
Incubation Temperature	36±1°C	Incubation Time	24-72 hours



Summary of the Procedure

- •
- Test microorganisms are grown on appropriate media.
- Culture used for test inoculum are evaluated for sterility, washed and concentrated in sterile phosphate buffered saline upon harvesting.
- The test inoculum is split into two equal parts and added to the appropriate number of nebulizers. Liquid culture should not exceed 20 ml per nebulizer.
- The device is setup per protocol requirements and operated per manufacturer's instructions.
- The chamber is setup and the safety checklist is completed prior to test initiation.
- Test is initiated by aerosolizing the microorganisms per the nebulizers and allowing the concentration to reach the required CFU/m³. Once the concentration is reached, a time zero sample is taken then the device is run for the specified contact time and an additional sample is taken for each contact time.
- The decontamination process is run, 4 hours of UV exposure, prior to any scientists entering the testing chamber.
- Samples are enumerated using standard dilution and plating techniques.
- Microbial concentrations are determined after appropriate incubation times.
- Reductions of microorganisms are calculated relative to concentration of the time zero or corresponding control run sample as applicable.



Study Notes:

4.0 ml of *Bacillus subtilis* ATCC 19659 spore stock was added to 46.0 ml of Phosphate Buffered Saline and mixed. 20.0 ml of inoculum was added to each nebulizer on testing that occurred on 05 NOV 2020 and 10 NOV 2020.

30.0 ml of *Mycobacterium smegmatis* ATCC 607 culture was added to 15.0 ml of Phosphate Buffered Saline and mixed. 20.0 ml of inoculum was added to each nebulizer on testing that occurred on 13 NOV 2020.

32.0 ml of *Mycobacterium smegmatis* ATCC 607 culture was added to 12.0 ml of Phosphate Buffered Saline and mixed. 20.0 ml of inoculum was added to each nebulizer on testing that occurred on 16 NOV 2020.

Per Study Sponsor instruction a small fan was placed in the upper corner of the NPAC chamber and allowed to run during the chamber runs to simulate the air circulation of an AC system.





Study Photos



Image 1: Test set up on 10 NOV 2020.



Control Results

Neutralization Method: N/A Growth Confirmation: Confirmed

Media Sterility: Sterile

Calculations

 $CFU/mI = (Average plate count) \times 1:10$ serial dilution factor

CFU/m³=[(CFU/ml x V_s) ÷ (T_s x 12.5 L/min)] x(1000 L/m³)

Where: $V_s = Bio$ -sampler volume (ml) $T_s = Time \text{ sampled (min)}$

$$\log_{10} \text{Reduction} = \log(\frac{B}{A})$$

Percent Reduction = $\frac{(B - A)}{B} \times 100\%$

Where:

B = Number of viable test microorganisms at time zero after nebulization A = Number of viable test microorganisms after the contact time



Results of the Study

Test Microorganism	Run Type	Treatment Time Point	Replicate	CFU/m³	Percent Reduction Compared to Time Zero	Log ₁₀ Reduction Compared to Time Zero	Adjusted Log ₁₀ Reduction ¹ Compared to Baseline
<i>B. subtilis</i> ATCC 19659		Time Zero	Replicate 1	5.81E+07		N/A	
	Baseline	15 Minute	Replicate 1	4.97E+07	14.39%	0.07	N/A
		30 Minute	Replicate 1	4.22E+07	27.27%	0.14	N/A
		60 Minute	Replicate 1	4.14E+07	28.79%	0.15	N/A
		90 Minute	Replicate 1	2.07E+07	64.30%	0.45	N/A
	Test	Time Zero	Replicate 1	9.41E+07	N/A		
		15 Minute	Replicate 1	2.42E+06	97.429%	1.59	1.52
		30 Minute	Replicate 1	6.89E+04	99.9268%	3.14	3.00
		60 Minute	Replicate 1	8.40E+01	99.99991%	6.05	5.90
		90 Minute	Replicate 1	< 8.64E+01	> 99.99991%	> 6.04	> 5.59

The limit of detection for this assay is 8.00E+01 CFU/m³ and values below the limit of detection are noted as "<8.00E+01" in the data table. ¹The Log reductions for the Test Runs are adjusted to account for natural die-off and gravitational settling observed in the Control Run.

Test Microorganism	Run Type	Treatment Time Point	Replicate	CFU/m³	Percent Reduction Compared to Time Zero	Log ₁₀ Reduction Compared to Time Zero	Adjusted Log ₁₀ Reduction ¹ Compared to Baseline
<i>M. Smegmatis</i> ATCC 607		Time Zero	Replicate 1	5.29E+07	N/A		
	Baseline	15 Minute	Replicate 1	4.41E+07	16.7%	0.08	N/A
		30 Minute	Replicate 1	4.51E+07	14.78%	0.07	N/A
		60 Minute	Replicate 1	3.42E+07	35.27%	0.19	N/A
		90 Minute	Replicate 1	2.55E+07	51.75%	0.32	N/A
	Test	Time Zero	Replicate 1	3.10E+08	N/A		
		15 Minute	Replicate 1	4.19E+05	99.86%	2.87	2.79
		30 Minute	Replicate 1	2.96E+05	99.905%	3.02	2.95
		60 Minute	Replicate 1	9.59E+03	99.997%	4.51	4.32
		90 Minute	Replicate 1	< 8.56E+01	> 99.99997%	> 6.56	> 6.24

The limit of detection for this assay is 8.00E+01 CFU/m³ and values below the limit of detection are noted as "<8.00E+01" in the data table. ¹The Log reductions for the Test Runs are adjusted to account for natural die-off and gravitational settling observed in the Control Run.



Additional Observations

Table 1. Champer Temperature and Humidity	Table	1.	Chamber	⁻ Temperature	and Humidity
---	-------	----	---------	--------------------------	--------------

Chamber Run	Chamber Temperature (Start/End)	Humidity (Start/End)
Baseline <i>B. subtilis</i> ATCC 19659	23.6°C/23.6°C	31%/36%
Test <i>B. subtilis</i> ATCC 19659	23.5°C/23.4°C	33%/37%
Baseline <i>M. Smegmatis</i> ATCC 607	23.7°C/23.7°C	32%/36%
Test <i>M. Smegmatis</i> ATCC 607	21.1°C/21.8°C	26%/29%

The results of this study apply to the tested substances(s) only. Extrapolation of findings to related materials is the responsibility of the Sponsor.

Copyright © Microchem Laboratory, 2020. Reproduction and ordinary use of this study report by the entity listed as "Sponsor" is permitted. Other copying and reproduction of all or part of this document by other entities is expressly prohibited, unless prior permission is granted in writing by Microchem Laboratory.

Page 11 of 11