US ERA ARCHIVE DOCUMENT

Environmental Technology Verification

Test Report of Control of Bioaerosols in HVAC Systems

Aeolus Corporation Synthetic Minipleat V-Cell, SMV-M13-2424

Prepared by

Research Triangle Institute



Under a Contract with U.S. Environmental Protection Agency





THE ENVIRONMENTAL TECHNOLOGY VERIFICATION PROGRAM





ETV Joint Verification Statement

TECHNOLOGY TYPE: VENTILATION MEDIA AIR FILTER

APPLICATION: FILTRATION EFFICIENCY OF BIOAEROSOLS IN

HVAC SYSTEMS

TECHNOLOGY NAME: Synthetic Minipleat V-Cell, SMV-M13-2424

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The U.S. Environmental Protection Agency (EPA) has created the Environmental Technology Verification (ETV) Program to facilitate the deployment of innovative or improved environmental technologies through performance verification and dissemination of information. The goal of the ETV Program is to further environmental protection by substantially accelerating the acceptance and use of improved and cost-effective technologies. ETV seeks to achieve this goal by providing high quality, peer-reviewed data on technology performance to those involved in the design, distribution, financing, permitting, purchase, and use of environmental technologies.

ETV works with recognized standards and testing organizations; stakeholder groups which consist of buyers, vendor organizations, permitters, and other interested parties; and with the full participation of individual technology developers. The program evaluates the performance of innovative and improved technologies by developing test plans that are responsive to the needs of stakeholders, conducting field or laboratory tests (as appropriate), collecting and analyzing data, and preparing peer-reviewed reports. All evaluations are conducted in accordance with rigorous quality assurance protocols to ensure that data of known and adequate quality are generated and that the results are defensible.

EPA's National Risk Management Research Laboratory contracted with the Research Triangle Institute (RTI) to establish a homeland-security-related ETV Program for products that clean ventilation air. RTI evaluated the performance of ventilation air filters used in building heating, ventilation and air-conditioning (HVAC) systems. This verification statement provides a summary of the test results for the Aeolus Corporation Synthetic Minipleat V-Cell, SMV-M13-2424 media air filter.

VERIFICATION TEST DESCRIPTION

All tests were performed in accordance with RTI's "Test/Quality Assurance Project Plan: Biological Testing of General Ventilation Filters," which was approved by EPA. Tests were performed for the following:

- Bioaerosol filtration efficiency tests of the clean and dust-loaded filter. Three bioaerosols were used in the testing:
 - o The spore form of the bacteria *Bacillus atrophaeus* (BG), a gram-positive spore-forming bacteria elliptically shaped with dimensions of 0.7 to 0.8 by 1 to 1.5 μ m,
 - o Serratia marcescens, a rod-shaped gram-negative bacteria with a size of 0.5 to 0.8 by 0.9 to 2.0 μ m, and
 - o The bacterial virus (bacteriophage) MS2 dispersed as a micrometer-sized polydisperse aerosol.
- Inert aerosol filtration efficiency tests consisting of an American National Standards
 Institute (ANSI)/American Society of Heating, Refrigerating and Air-Conditioning
 Engineers (ASHRAE) Standard 52.2-1999 type test (0.3 to 10 μm) and extended
 fractional efficiency measurements down to 0.02 μm particle diameter on both clean and
 dust-loaded filter.
- ASHRAE 52.2 test. This test provides filtration efficiency results (average of the minimum composite efficiency) given for three size ranges of particles: E1, 0.3 to 1.0 μ m; E2, 1.0 to 3.0 μ m; and E3, 3.0 μ m to 10 μ m.

VERIFIED TECHNOLOGY DESCRIPTION

As shown in Figure 1, the Aeolus Corporation Synthetic Minipleat V-Cell, SMV-M13-2424 media air filter is a V-cell filter with nominal dimensions of 0.61 by 0.61 by 0.31 m (24 by 24 by 12 in.). The filter has a plastic frame, and the filter media color is white and red stripes. The media is polypropylene. There are four V-cells with minipleated media. The Aeolus Corporation part number is SMV-M13-2424.

VERIFICATION OF PERFORMANCE

Verification testing of the Aeolus Corporation Synthetic Minipleat V-Cell, SMV-M13-2424 media air filter began on September 9, 2003 at the test facilities of RTI and was completed on October 8, 2003. The results for the bioaerosol filtration efficiency tests are presented in Figure 1 Table 1 for the clean and dust-loaded filter. Table 2 presents the results of the ASHRAE 52.2 test. All tests were conducted at an air flow of 0.93 m3/sec (1970 cfm).



Figure 1. Photograph of the Aeolus Corporation Synthetic Minipleat V-Cell, SMV-M13-2424 media filter.

Table 1. Bioaerosol Filtration Results

		Filtration	Filtration	Filtration
	Pressure Drop	Efficiency for	Efficiency for	Efficiency for
	Pa (in. H ₂ O)	Removal of	Removal of	Removal of
		B. atrophaeus, %	S. marcescens, %	MS2 phage, %
Clean	77 (0.31)	69	64	73
Dust loaded	348 (1.40)	99	99.5	99

Table 2. Summary of ASHRAE 52.2 Test

	E1 0.3 to 1.0 μ m,	E2 1.0 to 3.0 μm,	E3 3.0 to 10 μm, %	Minimum Efficiency Reporting Value (MERV)
Aeolus Corporation Synthetic Minipleat V- Cell, SMV-M13-2424	57	85	97	12 at 0.93m ³ /sec (1970 cfm)

The quality assurance officer reviewed the test results and the quality control data and concluded that the data quality objectives given in the approved test/QA plan were attained.

This verification statement addresses two performance measures of media air filters: filtration efficiency and pressure drop. Users of this technology may wish to consider other performance parameters such as service life and cost when selecting a media air filter for bioaerosol control. In accordance with the test/QA plan¹, this verification statement is applicable to filters manufactured from December 2003 through November 2006.

Original signed by E. Timothy Opp	pelt 12/8/2003
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ETV-HS

Research Triangle Institute

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Environmental Technology Verification

Test Report of Filtration Efficiency of Bioaerosols in HVAC Systems

Aeolus Corporation Synthetic Minipleat V-Cell, SMV-M13-2424

Prepared by:

Research Triangle Institute Engineering and Technology Research Triangle Park, NC 27709

GS10F0283K-BPA-1, EPA Task Order 1101 RTI Project No. 08787.001

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December 2003

Notice

This document was prepared by the Research Triangle Institute (RTI) with funding from the U.S. Environmental Protection Agency (EPA) through the General Service Administration Contract No. GS10F0283K per EPA's BPA-1, Task Order 1101. The document has undergone RTI and EPA peer and administrative reviews and has been approved for publication. Mention of corporation names, trade names, or commercial products does not constitute endorsement or recommendation for use of specific products.

Foreword

The Environmental Technology Verification (ETV) Program, established by the U.S. Environmental Protection Agency (EPA), is designed to accelerate the development and commercialization of new or improved environmental technologies through third-party verification and reporting of performance. The goal of the ETV Program is to verify the performance of commercially ready environmental technologies through the evaluation of objective and quality-assured data so that potential purchasers and permitters are provided with an independent and credible assessment of the technology that they are buying or permitting.

EPA's National Risk Management Research Laboratory contracted with the Research Triangle Institute (RTI) to establish a homeland-security related ETV Program for products that clean ventilation air. RTI developed (and EPA approved) the "Test/Quality Assurance Plan for Biological Testing of General Ventilation Filters¹." The test described in this report was conducted following this plan.

Availability of Report

Copies of this verification report are available from

- Research Triangle Institute
 Engineering and Technology Unit
 PO Box 12194
 Research Triangle Park, NC 27709-2194
- U.S. Environmental Protection Agency
 Air Pollution Prevention and Control Division, E305-01
 109 T.W. Alexander Drive
 Research Triangle Park, NC 27711

Web sites: http://www.epa.gov/etv/verifications

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Acronymns/Abbreviations

ANSI American National Standards Institute

ASHRAE American Society of Heating, Refrigerating and Air-Conditioning Engineers, Inc.

ASME American Society of Mechanical Engineers

B Bacillus

BG Bacillus atrophaeus (formerly B. subtilis var niger and Bacillus globigii)

cfm cubic feet per minute CFU colony forming unit

cm centimeter

d₅₀ median diameter (of particle)

DQO data quality objective

EPA U.S. Environmental Protection Agency ETV Environmental Technology Verification

F Fahrenheit fpm feet per minute HS homeland security

in. inch(es)

KCl potassium chloride kPa kilopascal(s) L liter(s)

MERV minimum efficiency reporting value

 $\begin{array}{lll} m & meter(s) \\ mm & millimeter(s) \\ mL & milliliter(s) \\ min & minute(s) \\ \mu m & micrometer(s) \end{array}$

NAFA National Air Filtration Association

nm nanometer(s)

OPC optical particle counter
QA quality assurance
QC quality control
Pa pascal(s)

PFU plaque forming units

psig pounds per square inch gauge
RTI Research Triangle Institute
SAE Society of Automotive Engineers
SMPS scanning mobility particle sizer

Acknowledgments

The authors acknowledge the support of all of those who helped plan and conduct the verification activities. In particular, we would like to thank Ted Brna, EPA's Project Manager, and Paul Groff, EPA's Quality Assurance Manager, both of EPA's National Risk Management Research Laboratory in Research Triangle Park, NC. We would also like to acknowledge the assistance and participation of our stakeholder group for their input, as well as Al Veeck and the National Air Filtration Association (NAFA), and Intertek ETL SEMKO, especially Theresa Peck, for their help in acquiring the filters, and Aeolus Corporation for donating the filters to be tested.

For more information on the Aeolus Corporation Synthetic Minipleat V-Cell, SMV-M13-2424 filter, contact

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1.0 Introduction

EPA's National Risk Management Research Laboratory contracted with the Research Triangle Institute (RTI) to establish a homeland-security related ETV Program for products that clean ventilation air. RTI convened a group of stakeholders representing government and industry with knowledge and interest in the areas of homeland security and building ventilation. The group met in December 2002 and recommended technologies to be tested. RTI then developed (and EPA approved) the "Test/Quality Assurance Plan for Biological Testing of General Ventilation Filters¹." The first round of tests included ten types of filters. The tests described in this report were conducted following this plan.

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2.0 Product Description

As shown in Figure 1, Aeolus Corporation Synthetic Minipleat V-Cell, SMV-M13-2424 media air filter is a V-cell filter with nominal dimensions of 0.61 by 0.61 by 0.31 m (24 by 24 by 12 in.). The filter has a plastic frame, and the filter media color is white and red stripes. The media is polypropylene. There are four V-cells with minipleated media. The Aeolus Corporation part number is SMV-M13-2424.

3.0 Test Procedure

The test program measured the culturable bioaerosol removal efficiency of general ventilation filters. Three tests were required to accomplish this goal. First, the American Society of Heating, Refrigerating and Air-Conditioning Engineers, Inc. (ASHRAE)

Standard 52.2² test was performed on one filter of the test filter type to determine the minimum

Figure 1. Photograph of the Aeolus Corporation Synthetic Minipleat V-Cell, SMV-M13-2424 Media Filter.

efficiency reporting value (MERV) of the filter. After determining the MERV, the biological test using three different bioaerosols and an inert aerosol test on both clean and fully dust-loaded filters were performed on a second filter. All tests were at an air flow rate of 0.93 m³/sec (1970 cfm) to conform to the conditions described in ASHRAE Standard 52.2.

All testing was performed in a test duct as specified in ASHRAE Standard 52.2. A schematic of the test duct is shown in Figure 2. The test section of the duct is 610 mm (24 in.) by 610 mm (24 in.) square. The locations of the major components, including the sampling probes, device section (filter holder), and the aerosol generator (site of aerosol injection) are shown.

The inert testing and the ASHRAE Standard 52.2 test were performed using a potassium chloride (KCl) aerosol. The filters were loaded using ASHRAE dust, composed of 72% Society of Automotive Engineers (SAE) fine, 23% powdered carbon, and 5% cotton linters. The final pressure drop was determined by the Standard's requirements.

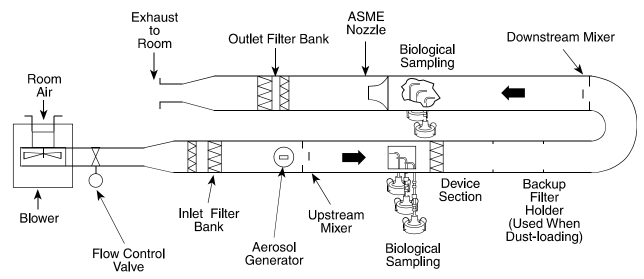


Figure 2. Schematic of Test Duct. Filter is placed in device section.

The bioaerosol tests were conducted using three microorganisms, two bacteria and one bacterial virus. The spore form of the bacteria *Bacillus atrophaeus* (formerly *B. subtilis var niger* and *Bacillus globigii* or BG) was used as the simulant for gram-positive spore-forming bacteria. The BG spore is elliptically shaped with dimensions of 0.7 to 0.8 by 1 to 1.5 μ m. *Serratia marcescens* was used as the surrogate for rod-shaped gram-negative bacteria. *S. marcescens* is 0.5 to 0.8 by 0.9 to 2.0 μ m.

The bacterial virus (bacteriophage) MS2 (0.02 to 0.03 μ m), having approximately the same aerosol characteristics as a human virus, was used as a surrogate for the viruses of similar and larger size and shape. Although the individual virus particles are in the submicrometer size range, the test particle size planned for the virus tests will span a range of sizes (polydispersed bioaerosol). This test was not designed to study the removal efficiencies for single individual virus particles; rather, it was designed to determine the removal efficiencies for virus particles as they are commonly found indoors. A representative challenge would be a micrometer-sized, polydispersed aerosol containing the phage because:

- The aerosols created from sneezing and coughing vary in size from < 1 to > 20 μ m, but the largest particles settle out and only the smaller sizes remain in the air for extended periods for potential removal by an air cleaner;
- Few viruses have been found associated with particles less than 1 μ m; and
- Nearly all 1 to 2 μ m particles are deposited in the respiratory tract, while larger particles may not be respired.

Bacteria suspension preparation for the aerosolization process required that the specific test organism be grown in the laboratory and the suspension prepared for aerosol generation in the test rig. The microbial challenge suspensions were prepared by inoculating the test organism on solid or liquid media, incubating the culture until mature, wiping organisms from the surface of the pure culture (if solid media), and eluting them into sterile diluent to a known concentration.

The bacterial virus challenge was prepared by inoculating a logarithmic phase broth culture of the host bacteria with phage and allowing it to multiply until the majority of the host bacteria were lysed. The mixture was centrifuged to remove the majority of the cell fragments. The resultant supernatant was the phage stock and was used as the challenge aerosol. The concentration of the phage stock was approximately 1×10^9 or higher plaque forming units per milliliter, (PFU) /mL.

The challenge organism suspensions were aerosolized using a Collison nebulizer (BGI, Waltham, MA) at 103.4 kPa (15 psig) air pressure. The nebulizer generates droplets with an approximate volume mean diameter of 2 μ m. The particle diameter after the water evaporates depends on the solids content of the suspension. Particle size was determined by the size of the suspended organism (if singlets).

Upstream and downstream sampling of the bacteria was accomplished using a one-stage Andersen viable bioaerosol sampler. The one-stage Andersen sampler is a 400-hole multiple-jet impactor operating at 28 L/min. The d_{50} is 0.65 μ m. After sampling, the petri dishes were removed from the sampler and incubated at appropriate times and temperatures for the test organism being used. Colony forming units (CFUs) were then enumerated and their identity confirmed.

The microbial viruses were collected in AGI-30s. The AGI-30 is a high velocity liquid impinger operating at a flow rate of 12.3 to 12.6 L/min. The d_{50} is approximately 0.3 μ m. The AGI-30 is the sampler against which the other commonly used bioaerosol samplers are often compared.

For the inert aerosol filtration efficiency measurements, the particle sizing measurements were made with two particle counting instruments: a Climet model 500 spectrometer/optical particle counter (OPC) covering the particle diameter size range from 0.3 to 10 μ m in 12 particle sizing channels and a TSI scanning mobility particle sizer (SMPS) to cover the range from 0.03 (or as low as 16 nm) to 0.5 μ m. In the test/QA plan there is a data control parameter for the SMPS that states the standard deviation on upstream counts be computed for each efficiency test based on the upstream particle counts and be less than 0.30 before the data is used. The lower size ranges for the SMPS are included only if they meet the data control parameter. (Table A2 of test/QA plan).

Quality Control (QC) procedures for running the test duct and the measuring equipment are defined in the test/QA plan.

The product tested was collected by the Intertek ETL SEMKO on July 14, 2003 following the NAFA *Product Certification Program Procedural Guide*³. RTI provided oversight into the selection of representative filters. For each filter type, a box or a minimum of four filters were procured and sent to RTI. The filters were used as shown in Table 1.

Full details of the test method can be found in RTI's test/QA plan¹.

Table 1. Numbers of Filters and Expected Utilization

Tests	Filter #			
	1	2	3	4
ASHRAE Standard 52.2 ² test	X			
Initial efficiency for an inert aerosol		X		
Initial efficiency for three bioaerosols		X		
Dust load to final pressure drop with ASHRAE dust		X		
Efficiency for inert aerosol after dust-loading		X		
Efficiency for three bioaerosols after dust-loading		X		
Reserve filter*			X	X

^{*}Filters # 3 and # 4 have been kept in reserve to be used if needed.

4.0 Test Results

The bioaerosol filtration efficiency results are found in Table 2.

Table 2. Bioaerosol Filtration Results for Filter # 2

Filter Condition	Pressure Drop Pa (in. H ₂ O)	Filtration Efficiency for Removal of B. atrophaeus, %	Filtration Efficiency for Removal of S. marcescens, %	Filtration Efficiency for Removal of MS2 phage, %
Clean	77 (0.31)	69	64	73
Dust-loaded	348 (1.40)	99	99.5	99

The ASHRAE filtration efficiencies and the MERV are shown in Table 3. The filtration efficiencies (average of the minimum composite efficiency) are presented by particle size groupings: E1, 0.3 to 1.0 μ m; E2, 1.0 to 3.0 μ m; and E3, 3.0 μ m to 10 μ m. The full ASHRAE 52.2 test results are provided in the Appendix.

Table 3. Summary of Removal Efficiency Using ASHRAE 52.2 Test for Filter # 1

Filter	E1 0.3 to 1.0 μm, %	E2 1.0 to 3.0 μm, %	E3 3.0 to 10 μm,	MERV
Aeolus Corporation Synthetic Minipleat V-Cell, SMV-M13-2424	57	85	97	12 at 0.93m ³ /sec (1970 cfm)

The filtration efficiency for inert particles is plotted so that the efficiencies for particles from about 0.03 to 10 μ m can be observed (Figure 3). Note that this is a logarithmic base 10 scale on

the X axis. Two instruments were used to obtain the measurements. The SMPS was used to measure particles up to 0.5 μ m and the OPC was used for particles from 0.3 to 10 μ m. There is good agreement in the size range covered by both instruments. These measurements were made on a filter when clean and then when dust-loaded.

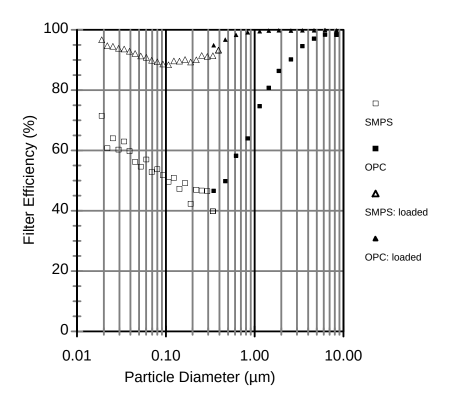


Figure 3. Summary of the Inert Aerosol Filtration Efficiency Data for the Clean and Dust-Loaded Filter, # 2.

The quality assurance officer has reviewed the test results and the quality control data and has concluded that the data quality objectives (DQOs) (Table 4) given in the approved test/QA plan have been attained.

Table 4. DQOs for Precision of Filtration Efficiency Measurements for Culturable Bioaerosol

	Test organism					
Data quality objective	Spore-forming bacteria	Vegetative bacteria	Bacterial virus			
	(B. atrophaeus)	(S. marcescens)	(MS2 phage)			
Precision of filtration	± 8 ^a	± 11 ^a	± 13 ^a			
efficiency, %						

^a Based on +/- one standard deviation of penetration computed from the coefficient of variance upstream and downstream culturable counts.

5.0 Limitations and Applications

This verification report addresses two performance measures of media air filters: filtration efficiency and pressure drop. Users may wish to consider other performance parameters such as service life and cost when selecting a general ventilation air filter for their application.

In accordance with the test/QA plan¹, this verification statement is applicable to filters manufactured from December 2003 through November 2006.

6.0 References

- 1. RTI. 2003. *Test/QA Plan for Biological Testing of General Ventilation Filters*. Research Triangle Institute, Research Triangle Park, NC.
- 2. ANSI/ASHRAE Standard 52.2-1999, *Method of Testing General Ventilation Air-Cleaning Devices*, American National Standards Institute/American Society of Heating, Refrigerating and Air-Conditioning Engineers, Atlanta, GA.
- 3. NAFA (National Air Filtration Association). 2001. *Product Certification Program Procedural Guide* Approved Version 1, Second Revision, February 2001. Virginia Beach, VA.

Appendix ASHRAE 52.2 Test Report

For Aeolus Corporation Synthetic Minipleat V-Cell, SMV-M13-2424

ASHRAE 52.2 TEST REPORT

Manufacturer: Aeolus Corp.

Product Name: Synthetic Minipleat V-Cell

SMV-M13-2424

ETV Filter ID: AEO3-A

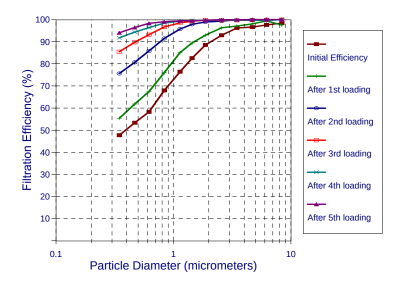
RTI Report No. AY09050301

Test Laboratory: RTI 919-541-6941

US EPA ARCHIVE DOCUMENT

ASHKAE Sta.	52.2 Air Cleaner This report app		-	•		
		31100 10 1				
Laboratory Data						
RTI Report No.	AY09050301			Date	9/5/03	
Test Laboratory	Research Tria	ngle Inst	itute			
Operator	Clayton			Supervisor	Owen/Hanley	
Particle Counter(s):	Brand C	Climet	_	Model	500	
Device Manufacturer's Data						
Manufacturer	Aeolus Corp.			_		
Product Name	Synthetic Mini	pleat V-0	Cell SMV-M13	-2424		
Product Model				_		
Test requested by	EPA			_		
Sample obtained from	NAFA			_		
Catalog rating:	Airflow rate		NA	Initial	dP (in. wg)	NA
Specified test conditions:	Airflow (cfm)		1970	Final	dP (in. wg)	1.40
	Face Velocity	(fpm)	493	-		
Device Description						
Nominal Dimensions (in.):	24 x 24 x 12		(height x wi	idth x depth)		
Generic name	V-Cell			Media color	white/red	
Amount and type of adhesive	NA			_		
Other attributes	minipleated me	edia in fo	our Vs			
Test Conditions						
Airflow (cfm)	1970	Temp	erature (F)	73	RH (%)	51
Face Velocity (fpm)	493	Final	Pressure Dro	p (in. wg)	1.40	
Test aerosol type:	KCI					
Remarks						
Resistance Test Results						
Initial resistance (in. wg)	0.31		Final resista	ance (in. wg)	1.40	
Minimum Efficiency Reporting D	Data					
Composite average efficiencies	E1_	57	E2	85	E3	97
Air cleaner average Arrestance pe	er Std 52.1:		NA	_		
Minimum efficiency reporting value	e (MERV) for the	device:		12	@ 1970	cfm

Report No. AY09050301 Research Triangle Institute



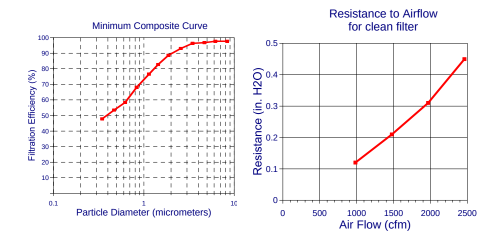


Figure A-1. Filtration Efficiency and Flow Resistance Curves for Aeolus Synthetic Minipleat V-cell SMV-M14-2424 Filter.

TABULATED DATA SUMMARY Report No. AY09050301 Research Triangle Institute

Summary of Test Conditions:

Product Manufacturer

Aeolus Corp. Synthetic Minipleat V-Cell SMV-M13-2424

Nominal Dimensions (in.)

24 x 24 x 12

Airflow (cfm)

Product Name

1970

Final Resistance (in. H2O)

1.40

Efficiency (%) per Indicated Size Range

OPC Channel Number Min. Diam. (µm) Max. Diam. (µm) Geo. Mean Diam. (µm)		1 0.3 0.4 0.35	2 0.4 0.55 0.47	3 0.55 0.7 0.62	4 0.7 1 0.84	5 1 1.3 1.14	6 1.3 1.6 1.44	7 1.6 2.2 1.88	8 2.2 3 2.57	9 3 4 3.46	10 4 5.5 4.69	11 5.5 7 6.20	12 7 10 8.37
	Run No.												
Initial efficiency	AY09040302	48	53	58	68	77	83	89	93	96	97	98	99
after first dust load	AY09050301	56	62	68	76	85	90	93	96	97	98	99	98
after second dust load	AY09050302	76	81	86	92	96	98	99	99	100	100	100	100
after third dust load	AY09050303	86	90	93	97	99	99	100	100	100	100	100	100
after fourth dust load	AY09050304	92	94	96	98	99	100	100	100	100	100	100	100
after fifth dust load	AY09060301	94	96	98	99	99	100	100	100	100	100	100	100
Minimum Composite E	fficiency	48	53	58	68	77	83	89	93	96	97	98	98

E1 = 57 (E1 is the average of the minimum composite efficiency values for particle diameters from 0.3 to 1 μ m.)

E2 = 85 (E2 is the average of the minimum composite efficiency values for particle diameters from 1 to 3 μ m.)

E3 = 97 (E3 is the average of the minimum composite efficiency values for particle diameters from 3 to 10 μ m.)

MERV: 12

Resistance to Airflow for Clean Filter:

Airflow (%)	Airflow (m3/s)	Airflow (cfm)	Air Velocity (fpm)	Air Velocity (m/s)	Resistance (in. H2O)	Resistance (Pa)
50	0.465	985	246	1.251	0.12	30
75	0.697	1478	369	1.876	0.21	52
100	0.930	1970	493	2.502	0.31	77
125	1.162	2463	616	3.127	0.45	112

Resistance to Airflow with Loading at 0.93 m3/s (1970 cfm)

	Resistance (in. H2O)	Resistance (Pa)
Initial	0.31	77
After first dust load	0.35	86
After second dust load	0.58	145
After third dust load	0.85	213
After fourth dust load	1.13	280
After fifth dust load	1.40	348

Weight gain of filter after completion of dust loading steps

252.9 g