



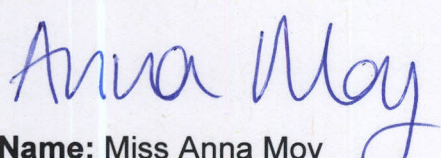
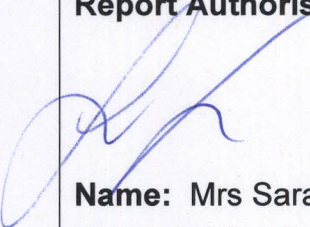
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# An Evaluation of Filtration Efficiencies Against Bacterial and Viral Aerosol Challenges

## Report No. 155/11

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<b>HPA Ref.</b>	Project No. 155/11
<b>Customer Ref.</b>	CAMR 112011
<b>Report Prepared For</b>	Medical Research and Development
<b>Operator</b>	Miss Anna Moy
<b>Issue Date</b>	11 <sup>th</sup> January 2012
<b>Copy Number</b>	1
<b>Distribution</b>	Mr. J. Hernandez (MRD) Mr. A. Bennett (HPA) HPA Central Records (Dr. P. Hammond)

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## SUMMARY

The efficiencies of eight MRD bacterial/viral filters for pulmonary function testing equipment (NeumoFilt® Ergo, Ref: 1121/100, Lot: 13211) supplied by MRD, were determined against aerosols containing micro-organisms. The filters were challenged with either bacterial spores of *Bacillus atrophaeus* or viral aerosols of MS-2 coliphage NCIMB 10108, fresh from the packaging. The filters were challenged at 55 or 750 litres min<sup>-1</sup>.

The results are summarised as follows:-

Filter N <sup>o</sup>	Aerosol Challenge	Flow Rate (l/min)	% Efficiency	Titre Reduction
1	<i>B. atrophaeus</i>	55	99.9976	4.25 x 10 <sup>4</sup>
2	<i>B. atrophaeus</i>	55	99.9972	3.54 x 10 <sup>4</sup>
3	<i>B. atrophaeus</i>	750	99.5024	2.01 x 10 <sup>2</sup>
4	<i>B. atrophaeus</i>	750	99.7506	4.01 x 10 <sup>2</sup>
5	MS-2	55	99.9987	7.49 x 10 <sup>4</sup>
6	MS-2	55	99.9969	3.25 x 10 <sup>4</sup>
7	MS-2	750	99.9873	7.90 x 10 <sup>3</sup>
8	MS-2	750	99.9825	5.70 x 10 <sup>3</sup>



## INTRODUCTION

Contamination of respiratory apparatus during expiration has been recognised since 1965 as a source of noscomial infections (1). Disposable filters placed between the patient and the apparatus are designed to prevent such contamination. There is a need for a standard method to test the effectiveness of these filters against bacteria and viruses. A system has been developed at the Health Protection Agency, Porton Down (HPA) to test the efficiencies of many types of microbiological filters including bacterial/viral filters for pulmonary function testing equipment.

The efficiencies of the filters were tested on our small rig facility. This rig is based on an apparatus developed originally by Henderson and Druett (2, 3) to study experimental airborne infection, where a suspension of micro-organisms in aqueous solution is nebulised by a Collison spray (4) forming a fine aerosol containing viable micro-organisms. The generated aerosols are injected into an air stream flowing into a long stainless steel tube. The efficiencies of the filters are calculated by determining the airborne concentration of viable micro-organisms upstream and downstream of the filter using suitable aerosol sampling techniques and microbial assay methods. Filters can be challenged with micro-organisms on the small test rig up to flow rates of approximately 2000 litres per minute. This system was used to test the filters at both 55 and 750 Litres  $\text{min}^{-1}$ .

The choice of bacterial strains to challenge and test these filters is based on a non-pathogenic model providing the highest possible challenge concentration of viable micro-organisms to allow a fully quantitative assessment of the filters to be made. To do this, spores of *Bacillus atrophaeus* were used as the bacterial model because they are known to survive the stresses caused by aerosolisation. The spores were washed thoroughly and finally suspended in distilled water before nebulisation. During nebulisation the water is rapidly evaporated from the droplets formed so that monodispersed aerosols of viable spores actually challenge the filter in this system (5).



Because of the health hazards involved, it is unrealistic to evaluate these filters using human viruses. Fortunately, RNA-phages are of a similar size as the smallest human viruses and the efficiencies of the filters for removing human viruses from air streams can be gauged by measuring the penetration of aerosolised coliphage through the filter. MS-2 phage is an unenveloped single stranded RNA coliphage, 23 nm in diameter with a molecular weight of  $3.6 \times 10^6$  Daltons. MS-2 coliphage sprayed from the supernatant of centrifuged spent bacterial lysate are known to remain infectious at the conditions tested here (6). By spraying this suspension from a Collison nebuliser, the airborne coliphage are carried in droplets, which are much larger than the infectious particles, consisting mostly of bacterial lysate and media constituents.

## MATERIALS AND METHODS

### Test micro-organism

#### *Bacillus atrophaeus* spores (NCTC 10073)

The *B. atrophaeus* spores ( $>10^9$  colony forming units (cfu) per ml) which had been thoroughly washed in distilled water were suspended in distilled water. The suspension was prepared from batches previously prepared by the HPA Production Division (7).

#### MS-2 phage (NCIMB 10108)

A vial of MS-2 phage (NCIMB 10108) was obtained from the National Collection of Industrial and Bacteria, Torry Research Station, Aberdeen. A stock suspension of coliphage was prepared by inoculating 0.1 ml of a  $10^{11}$  plaque forming unit (pfu) per ml coliphage suspension into 500 ml nutrient broth containing  $1 \times 10^8$  *Escherichia coli* (NCIMB 9481) during the logarithmic growth phase. The suspension was aerated by shaking at 37°C. The bacterial cells lysed within 30 minutes and the suspension was centrifuged to remove the cell debris. The supernatant was transferred to a fresh flask and 10 drops of chloroform were added to kill any contaminating bacteria. This was used as the stock suspension of MS-2. The concentration of phage was determined as described later.

A high-titre suspension of MS-2 for challenging the filters was prepared as follows:-

The *E. coli* 9481 host was inoculated on a fresh TSBA plate, which was incubated at  $37 \pm 2^\circ\text{C}$  for 19 - 20 hr. The *E. coli* was sub-cultured from this plate by a 10  $\mu\text{l}$  loop to 60 ml sterile Tryptone Soya broth (TSB) in a 500 ml flask. After mixing thoroughly the flask was placed in a shaking incubator (120 rpm) for 150 mins at  $37 \pm 2^\circ\text{C}$ . The suspension of coliphage was then prepared by inoculating a total of  $4 \times 10^{11}$  plaque forming unit (pfu) coliphage suspension into the 500 ml flask containing the 60 ml TSB. The suspension was then aerated by shaking at  $37 \pm 2^\circ\text{C}$  for a further 3 hours. The suspension was centrifuged twice at 2,000 g for 20 minutes each to remove the cell debris. The supernatant was transferred to a fresh flask. The concentration of phage was determined as described below.

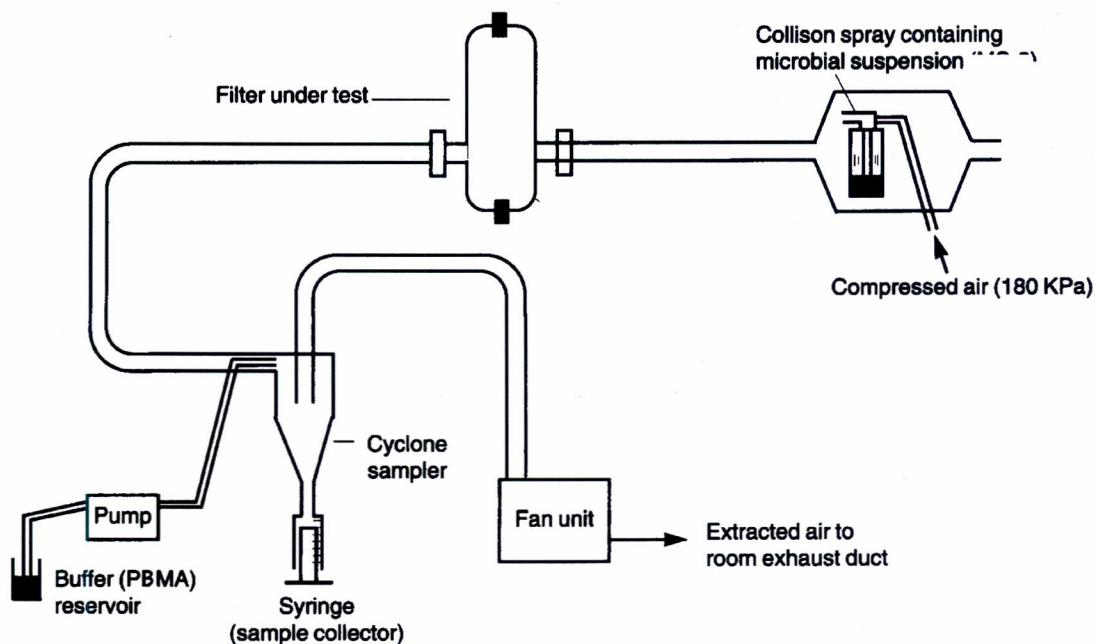
## Filter

Eight MRD bacterial/viral filters for pulmonary function testing equipment (NeumoFilt® Ergo, Ref: 1121/100, Lot: 13211) were provided for testing by MRD. Their efficiencies were determined against bacterial and viral aerosols containing *B. atrophaeus* spores and MS-2 coliphage at 55 and 750 litres min<sup>-1</sup>.

## Challenging filters on the small test rig

The small test rig (Figure 1) was designed to deliver a high titre challenge of *B. atrophaeus* spores and MS-2 coliphage in aerosols at either 55 or 750 litres per minute.

**FIGURE 1. SMALL TEST RIG FOR TESTING FILTERS WITH MICROBIAL AEROSOLS**



For the filters tested at 55 litres per minute a May Bottom Stage air sampler was used instead of the cyclone sampler.



The apparatus consisted of the following essential parts:-

- One pre-weighed 6-jet Collison spray containing 30 ml *B. atrophaeus* suspension (containing  $4.70 \times 10^9$  cfu per ml in distilled water) or one pre-weight 3-jet Collison spray containing 30 ml of MS-2 coliphage (containing  $2.85 \times 10^{12}$  pfu per ml in 50% (v/v) nutrient broth). The Collison spray was operated to nebulise its contents at a pressure of 180 KPa into the air stream in the spray tube.
- Stainless steel spray tube 90 cm length and 5 cm diameter to allow mixing and conditioning of the aerosols generated from the Collison.
- Suitable sterile tubing connectors and tapers to allow insertion of the filter to be tested in the system.
- For the filters tested at 55 litres per minute one May Bottom Stage air sampler (8) containing 15ml of sterile phosphate buffer containing manucol and antifoam (PBMA) operates the flow of sampled air via a vacuum pump. The air containing the microbial suspension is drawn through the system at a flow rate of 55 litres per minute. Each filter was inserted in turn in the apparatus and the Collison spray containing the microbial suspension was activated. The air was sampled for five minutes by the May Bottom Stage air sampler. The collecting fluid was removed from the May Bottom Stage air sampler and assayed for microbes as described below. The Collison spray was weighed after each test to determine the weight loss. The challenge concentration was determined by operating the system with the filters removed.
- One Cyclone sampler (9) (manufactured by The Hampshire Glass Company, Southampton) operates the flow of sampled air via a vacuum pump. The air containing the microbial aerosols is drawn through the system at a flow rate of 750 litres per minute. Sterile phosphate buffer containing manucol and antifoam (PBMA) was used as the collection fluid and was fed into the cyclone inlet at a rate of



approximately 1ml per minute by a peristaltic pump. The particles in the air stream were deposited by centrifugal force on the cyclone wall and were collected by the swirling liquid, which was withdrawn by a syringe at the end of the challenge period. The volume of collection fluid collected by the cyclone was measured for each filter. Each filter was inserted in turn in the apparatus and the pre-weighed Collison spray was activated. The air was sampled for 5 minutes by the Cyclone. The collecting fluid was removed from the samplers and assayed for spores or MS-2 as described below. The Collison spray was weighed after each test to determine the weight loss. The challenge concentration was determined by operating the system with the filters removed.

#### **Assay of *B. atrophaeus* in collecting fluids**

The collecting fluids from the samplers linked to the spray tube (i.e. without filter) were suitably diluted in PBMA and plated (0.1 ml) on duplicate Tryptone Soya agar (TSA) plates. The TSA plates were incubated at 37°C for 18 hours and any orange colonies were counted. Suitably diluted suspensions (0.1 ml) of the collecting fluid from each sampler placed behind the filter was also spread on duplicate TSA plates and these TSA plates were incubated at 37°C for 18 hours and any distinctive orange colonies were counted.

#### **Assay of MS-2 coliphage in collecting fluids**

A fresh TSA plate was inoculated with *Escherichia coli* NCIMB 9481 from a stock plate previously stored at  $4 \pm 2^\circ\text{C}$ . This plate was incubated at  $37 \pm 2^\circ\text{C}$  for 19 - 20 hrs. The *E. coli* 9481 was subcultured by transferring a 10  $\mu\text{l}$  loopful from the plate to 10 ml sterile nutrient broth in a glass universal bottle. After mixing, the universal bottle was incubated at  $37 \pm 2^\circ\text{C}$  for 260 minutes before use. Meanwhile, stoppered bottles containing 3 ml volumes of soft phage agar were heated for at least 90 minutes at 90 to 100°C and then stored at  $60 \pm 2^\circ\text{C}$  until required. These bottles were then cooled to 45°C before use. The suitably diluted MS-2 suspension in PBMA





(100 µl) was added to the soft agar followed immediately by 3 drops of the *E. coli* 9481 suspension using a 50 D (20 µl per drop) Pasteur pipette. After mixing, it was poured immediately on a TSBA (Tryptone Soya Broth agar) plate. Duplicate samples were carried out (the dilution selected should give 30 to 100 plaque forming units (pfu) per plate). The plates were incubated at 37 ± 2°C overnight. The clear plaques were counted.

### **Determination of effectiveness of the filter**

The effectiveness of the filter is expressed in the following ways:-

- Percentage efficiency. This is defined as follows:-

$$\frac{\text{cfu or pfu collected without filter in place} - \text{cfu or pfu with filter in place}}{\text{cfu or pfu collected without filter in place}} \times 100$$

- Titre Reduction. This is defined as follows:-

$$\text{Titre reduction} = \frac{\text{Total cfu or pfu collected without filter in place}}{\text{Total cfu or pfu with filter in place}}$$



## RESULTS

### Filter Integrity Tests Using Micro-Organisms

#### Test Conditions

Date	Dec 11/Jan 12	Challenge	<i>Bacillus atrophaeus</i>		
Operators	A. Moy	Micro-organisms			
Apparatus	Small Rig	Suspension Fluid	Sterile distilled water		
Spray	6-Jet Collison	Concentration	4.70 x 10 <sup>9</sup> cfu/ml		
Relative Humidity (RH):		Ambient	Temperature:		22 ± 3°C
Samplers	May Bottom Stage	Sampling Time	5	min at	55 Litres/min
Collecting Fluid		PBMA	Volume		15ml
Filters Tested:	2 x NeumoFilt® Ergo, Ref: 1121/100, Lot: 13211				

**Table 1.** Results from challenging 2 MRD Filters

Filter	Sample	Ave. Challenge (pfu)	Total Collected (pfu)	% Efficiency	Titre Reduction
F1	Spray off (background)	N/A	<1	N/A	N/A
	Spray on	1.34 x 10 <sup>9</sup>	3.15 x 10 <sup>4</sup>	99.9976	4.25 x 10 <sup>4</sup>
F2	Spray off (background)	N/A	<1	N/A	N/A
	Spray on	1.22 x 10 <sup>9</sup>	3.45 x 10 <sup>4</sup>	99.9972	3.54 x 10 <sup>4</sup>

\* cfu - colony forming units



### Test Conditions

Date	Dec 11/Jan 12	Challenge	<i>Bacillus atrophaeus</i>		
Operators	A. Moy	Micro-organisms			
Apparatus	Small Rig	Suspension Fluid	Sterile distilled water		
Spray	6-Jet Collison	Concentration	4.70 x 10 <sup>9</sup> cfu/ml		
Relative Humidity (RH):		Ambient	Temperature:	22 ± 3°C	
Samplers	Cyclone	Sampling Time	5	min at	750 Litres/min
Collecting Fluid	PBMA		Volume	Various	
Filters Tested:	2 x NeumoFilt® Ergo, Ref: 1121/100, Lot: 13211				

**Table 1.** Results from challenging 2 MRD Filters

Filter	Sample	Ave. Challenge (pfu)	Total Collected (pfu)	% Efficiency	Titre Reduction
F3	Spray off (background)	N/A	<1	N/A	N/A
	Spray on	8.20 x 10 <sup>8</sup>	4.08 x 10 <sup>6</sup>	99.5024	2.01 x 10 <sup>2</sup>
F4	Spray off (background)	N/A	<1	N/A	N/A
	Spray on	8.14 x 10 <sup>8</sup>	2.03 x 10 <sup>6</sup>	99.7506	4.01 x 10 <sup>2</sup>

\* cfu - colony forming units

### Test Conditions

Date	Dec 11/Jan 12	Challenge	MS-2 Coliphage
Operators	A. Moy	Micro-organisms	
Apparatus	Small Rig	Suspension Fluid	50% Nutrient Broth
Spray	3-Jet Collison	Concentration	$2.85 \times 10^{12}$
Relative Humidity (RH):		Ambient	Temperature: $22 \pm 3^{\circ}\text{C}$
Samplers	May Bottom Stage	Sampling Time	5 min at 55 Litres/min
Collecting Fluid	PBMA	Volume	15ml
Filters Tested:	2 x NeumoFilt® Ergo, Ref: 1121/100, Lot: 13211		

**Table 1.** Results from challenging a 2 MRD filters

Filter	Sample	Ave. Challenge (pfu)	Total Collected (pfu)	% Efficiency	Titre Reduction
F5	Spray off (background)	N/A	<1	N/A	N/A
	Spray on	$6.85 \times 10^{11}$	$9.15 \times 10^6$	99.9987	$7.49 \times 10^4$
F6	Spray off (background)	N/A	<1	N/A	N/A
	Spray on	$1.05 \times 10^{12}$	$3.23 \times 10^7$	99.9969	$3.25 \times 10^4$

\* pfu - plaque forming units



### Test Conditions

Date	Dec 11/Jan 12	Challenge	MS-2 Coliphage		
Operators	A. Moy	Micro-organisms			
Apparatus	Small Rig	Suspension Fluid	50% Nutrient Broth		
Spray	3-Jet Collison	Concentration	$2.85 \times 10^{12}$		
Relative Humidity (RH):		Ambient	Temperature:	$22 \pm 3^{\circ}\text{C}$	
Samplers	Cyclone	Sampling Time	5 min at	750	Litres/min
Collecting Fluid	PBMA	Volume	Various		
Filters Tested:	2 x NeumoFilt® Ergo, Ref: 1121/100, Lot: 13211				

**Table 1.** Results from challenging 2 MRD filters

Filter	Sample	Ave. Challenge (pfu)	Total Collected (pfu)	% Efficiency	Titre Reduction
F7	Spray off (background)	N/A	<1	N/A	N/A
	Spray on	$1.39 \times 10^{11}$	$1.76 \times 10^7$	99.9873	$7.90 \times 10^3$
F8	Spray off (background)	N/A	<1	N/A	N/A
	Spray on	$1.75 \times 10^{11}$	$3.07 \times 10^7$	99.9825	$5.70 \times 10^3$

\* pfu - plaque forming units



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