

Effect of manually assisted cough and mechanical insufflation on cough flow of normal subjects, patients with chronic obstructive pulmonary disease (COPD), and patients with respiratory muscle weakness

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Abstract

Background—It has been suggested that cough effectiveness can be improved by assisted techniques. The effects of manually assisted cough and mechanical insufflation on cough flow physiology are reported in this study.

Methods—The physiological actions and patient self-assessment of manually assisted cough and mechanical insufflation were investigated in 29 subjects (nine normal subjects, eight patients with chronic obstructive pulmonary disease (COPD), four subjects with respiratory muscle weakness (RMW) with scoliosis, and eight subjects with RMW without scoliosis).

Results—The peak cough expiratory flow rate and cough expiratory volume were not improved by manually assisted cough and mechanical insufflation alone or in combination in normal subjects. The median increase in peak cough expiratory flow in subjects with RMW without scoliosis with manually assisted cough alone or in combination with mechanical insufflation of 84 l/min (95% confidence interval (CI) 19 to 122) and 144 l/min (95% CI 14 to 195), respectively, reflects improvement in the expulsive phase of coughing by these techniques. Manually assisted cough and mechanical insufflation in combination raised peak expiratory flow rate more than either technique alone in this group. The abnormal chest shape in scoliotic subjects and the fixed inspiratory pressure used made effective manually assisted cough and mechanical insufflation difficult in this group and no improvements were found. In patients with COPD manually assisted cough alone and in combination with mechanical insufflation decreased peak expiratory flow rate by 144 l/min (95% CI 25 to 259) and 135 l/min (95% CI 30 to 312), respectively.

Conclusions—Manually assisted cough and mechanical insufflation should be considered to assist expectoration of secretions in patients with RMW without scoliosis but not in those with scoliosis. (Thorax 2001;56:438–444)

Keywords: manually assisted cough; mechanical insufflation; cough flow physiology; chronic obstructive pulmonary disease; respiratory muscle weakness; scoliosis

Cough is an important part of the airway defence aiding the removal of secretions.¹ It has three components—an inspiratory phase consisting of inhalation to greater than resting end inspiratory volume; a compressive phase comprising glottic closure accompanied by an increased intrathoracic pressure as a result of expiratory muscle contraction; and an expulsive phase resulting from sudden glottic opening.² The high air flow velocities in the expulsive phase transfer kinetic energy from air to the secretions or foreign bodies, shearing them off the bronchial wall and transporting them to the pharynx or mouth.

Neuromuscular disorders such as amyotrophic lateral sclerosis,³ poliomyelitis,⁴ myasthenia gravis,⁵ and muscular dystrophies⁶ can impair any of these three phases. In pulmonary disorders such as chronic bronchitis and emphysema the reduced expiratory flow resulting from dynamic airway compression and increased viscosity of bronchial secretions is probably the main cause of cough ineffectiveness.⁷

Several methods of assisting the expulsive phase of coughing have been proposed.^{8,9} We have examined the efficacy of manually assisted cough and mechanical insufflation on cough dynamics in normal subjects, patients with chronic obstructive pulmonary disease (COPD), and those with respiratory muscle weakness with and without scoliosis.

Methods

SUBJECTS

We studied normal volunteers and patients suffering from respiratory muscle weakness (RMW) due to neuromuscular disorders and those with chronic obstructive pulmonary disease (COPD). All normal subjects were non-smokers without a history of respiratory, neuromuscular, or cardiovascular disease. Respiratory muscle weakness was defined as static inspiratory and expiratory maximal mouth pressures <70% predicted. Subjects with RMW were subdivided according to whether or not they had a thoracic scoliosis. Scoliosis was diagnosed on physical examination and from spinal radiographs with a Cobb angle of >70°. All cases of RMW had been diagnosed by a neurologist and had no other respiratory disease. All patients with COPD fulfilled the

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American Thoracic Society criteria.¹⁰ All subjects gave informed consent prior to participation in the study.

Nine normal subjects, eight with COPD, and 12 with RMW were studied. The RMW group without scoliosis comprised seven subjects with amyotrophic lateral sclerosis and one with Becker's muscular dystrophy. There were two subjects with previous poliomyelitis, one with Duchenne's muscular dystrophy, and one with spinal muscular atrophy who had scoliosis on spinal radiographs.

STUDY DESIGN

Forced expiratory volume in one second (FEV_1), forced vital capacity (FVC), and maximal voluntary ventilation (MVV) were measured using a pneumotachograph spirometer (Vitalograph, Buckingham, UK) and total lung capacity (TLC) was measured using body plethysmography (Masterlab, Jaeger, Wurzburg, Germany). Maximum mouth inspiratory (P_{imax}) and expiratory (P_{emax}) pressures were measured with a hand held myometer (P K Morgan, Gillingham, Kent, UK). P_{imax} was measured at residual volume and P_{emax} was measured at TLC with the cheeks held. Subjects performed these manoeuvres until a reproducible maximal effort was achieved. All values were expressed as raw data and percentage predicted values.^{11 12}

Cough evaluation was performed using a modification of a method previously described.¹³ A face mask (Hans Rudolf) was attached directly to a 4.5 cm pneumotachograph (P K Morgan) from which the peak cough expiratory flow and cough expiratory volume were derived using an electronic transducer and integrator. The pneumotachograph had a linear response up to 900 l/min and a response time of <5 ms. All data were recorded digitally with an analogue to digital converter onto an IBM compatible computer using the CARDAS data logging system (Oxford Medical Sciences, Oxford, UK). The cough expiratory volume and peak cough expiratory flow were defined as the maximal volume and flow recorded, respectively (fig 1A and B). The time from onset of expiratory flow to peak cough expiratory flow, the peak value time, was measured (fig 1B). The system was calibrated between subjects for flow using a rotameter, range 0–450 l/min (Si-plan Electronics Research, Stratford on Avon, UK) and for volume using a 1 litre calibration syringe (P K Morgan).

Oesophageal and gastric pressures were measured to estimate pleural and abdominal pressures during coughing using balloons (P K Morgan) connected to a differential pressure transducer (Si-plan Electronics Research). The data were recorded onto a paper chart recorder (Dash 3, Astromed, Warwick, RI, USA). The nasopharynx was anaesthetised with topical lidocaine before the balloons were inserted. Baseline cough evaluation was performed in subjects before and after passage of balloons. Subjects undergoing oesophageal manometry had fasted prior to the investigation.

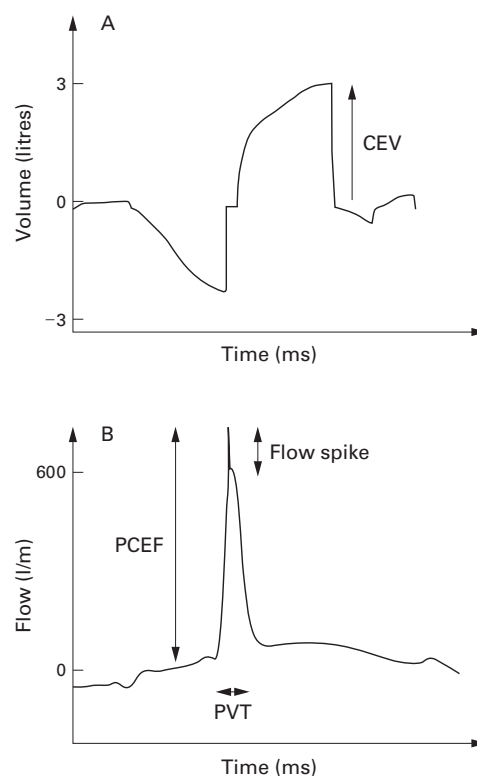


Figure 1 Normal recordings of (A) cough expiratory volume (CEV) and (B) cough expiratory flow. The peak cough expiratory flow rate (PCEF) was measured from the initiation of a positive flow rate to the maximum obtained value. This included the flow spike if present. The peak value time (PVT) was measured as the time taken from initiation of a positive flow rate to the PCEF. The cough expiratory flow volume was taken as the maximum expiratory volume recorded.

The cough manoeuvre was performed with the subjects in a semirecumbent position. It consisted of inspiration to near TLC followed by a maximal voluntary cough. All subjects practised coughing while the face mask was adjusted to minimise air leaks. An investigator held the subject's cheeks during the cough manoeuvres. Subjects were asked to perform three consistent coughs as baseline values prior to evaluation of assisted cough techniques. Consistent coughs were defined as peak cough expiratory flows that did not vary by more than 15%.

Manually assisted cough was performed by an experienced physiotherapist. It consisted of manual thoracoabdominal compression during the expulsive phase of a maximal voluntary cough. Hand positions during thoracoabdominal compression were optimised for patients with scoliosis by placing the hand used for thoracic compression on the hyperinflated hemithorax. Mechanical insufflation was performed with an In-exsufflator (J H Emerson Co, Cambridge, MA, USA) set to give 20 cm H_2O inspiratory and -20 cm H_2O expiratory pressure. Two In-exsufflator cycles were used to aid inflation and deflation of the thorax and after a third inspiration the subject was asked to make a maximal voluntary cough without the aid of negative pressure. All subjects practised with both manually assisted cough and mechanical

Table 1 Baseline comparison of lung function and respiratory muscle assessment

	Normal subjects (n=9)	COPD (n=8)	Neuromuscular weakness	
			Without scoliosis (n=8)	With scoliosis (n=4)
Age (years)	27 (17–71)	65 (52–74)	63 (27–73)	57 (44–66)
Sex	4M, 5F	3M, 5F	8M	3M, 1F
FEV ₁ (l)	3.8 (1.7–4.2)	0.95 (0.35–1.1)	0.73 (0.48–1.8)	0.8 (0.65–1.25)
FEV (% pred)	98 (88.3–120.1)	37 (13.4–44.2)	19 (13.6–43.9)	33 (28.4–40.4)
FVC (l)	4.6 (2.1–5.9)	1.8 (0.77–2.75)	0.83 (0.55–1.57)	1.5 (0.7–1.75)
FVC (% pred)	100 (78.3–120.8)	66 (24.5–84.9)	18 (13–42.0)	35 (15–42.9)
PEFR (l/min)	444 (410–633)	212 (110–270)	123 (68–150)	225 (220–240)
PEFR (% pred)	103 (86.9–140)	51 (24.9–66.7)	24 (13.1–31.9)	47 (45.7–66.7)
P _{imax} (cm H ₂ O)	99 (59–137)	37 (18–91)	15 (11–22)	37 (30–49)
P _{imax} (% pred)	115 (87–151)	44 (18.5–93.3)	14 (11.9–24.9)	41 (37.6–50.7)
P _{emax} (cm H ₂ O)	126 (104–239)	84 (52–167)	22 (16–35)	51 (17–62)
P _{emax} (% pred)	118 (83–228)	94 (74.7–194.3)	19 (11.7–27.6)	42 (36.4–67.3)
MVV (l/min)	128 (75–195)	33 (19–60)	26 (16–35)	33 (21–44)
MVV (% pred)	107 (71–115)	32 (16.7–66.1)	17 (11.1–27.8)	24 (17.0–33.8)

Data are expressed as median (range).

FEV₁ = forced expiratory volume in one second; FVC = forced vital capacity; PEFR = peak expiratory flow rate; P_{imax}, P_{emax} = maximum inspiratory and expiratory pressure; MVV = maximum voluntary ventilation.

insufflation prior to the investigations. Familiarisation with the techniques took no more than five practice coughs with each manoeuvre. At least 5 minutes elapsed between each cough manoeuvre and the best result of at least three attempts of each was analysed.

To exclude bias the order of the treatments was randomised for each subject. At the end of the treatment, subjects were asked whether each assisted cough technique had aided, impaired, or had no effect on their cough and to rank them in order of perceived effectiveness.

STATISTICAL ANALYSIS

Statistical analysis was performed using SPSS software for Windows (SPSS, Chicago, USA). The Mann-Whitney/Wilcoxon rank sum test was used to compare predicted pulmonary function test data with observed data for the normal subjects with those with COPD and

those with respiratory muscle weakness, with and without scoliosis. It was also used to compare the pulmonary function data between patients with RMW with and without scoliosis. The Wilcoxon matched pairs signed rank test was used to investigate the effect of different techniques (mechanical insufflation, manually assisted cough, and in combination) on peak cough expiratory flow, cough expiratory volume, and peak value time. Analyses of order of cough manoeuvres on results was made using the Friedman test. Statistical significance was taken as $p < 0.05$. The Bonferroni adjustment was used to correct for repeat pairwise comparisons.

Results

Data for age, sex, and pulmonary function of the four subject groups in this study are shown in table 1. The lung volumes of the normal subjects were not significantly different from predicted values. Those with COPD had a median FEV₁/FVC of only 44% (range 21.0–68.9) and a significantly increased TLC, a reduced P_{imax}, normal P_{emax}, and reduced MVV. Subjects with RMW with and without scoliosis had a significantly reduced FEV₁, FVC, P_{imax}, P_{emax}, and MVV compared with predicted values and those without scoliosis had a significantly lower FEV₁, peak expiratory flow rate (PEFR), P_{imax}, P_{emax}, and MVV than those who did not.

In normal subjects the digital recordings of flow during a cough revealed a sharp upstroke in flow with a transient flow spike followed by a gradual fall and finally a more rapid declining phase (fig 2A). Peak cough expiratory flow was higher than PEFR. The cough expiratory volume was lower than the FEV₁.

In the presence of RMW the upstroke of the cough flow rate was considerably reduced and

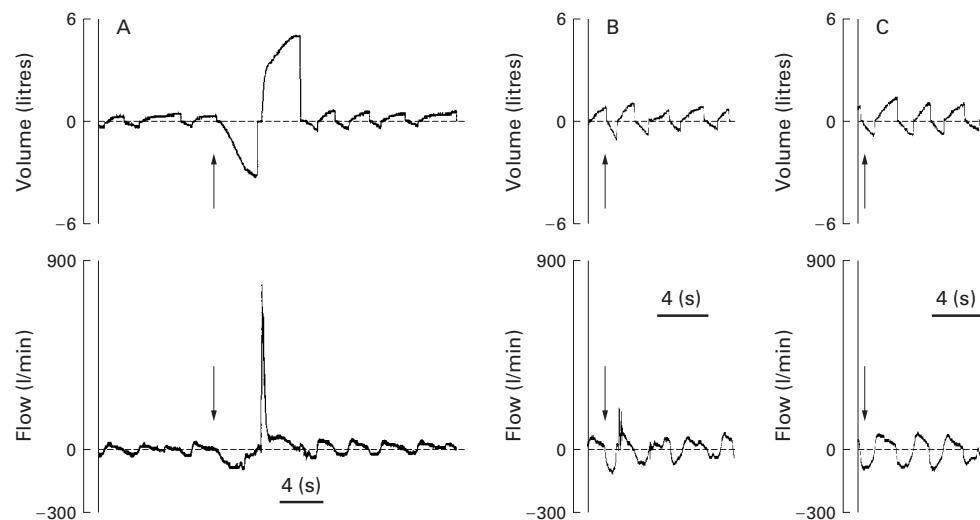


Figure 2 Typical flow and volume recordings before and during a maximum voluntary cough manoeuvre in (A) normal subjects, (B) patients with COPD, and (C) patients with respiratory muscle weakness without scoliosis. All recordings are shown on the same volume, flow, and time scale for comparison. The upper recording shows changes in inspiratory and expiratory volumes with time and the lower recording shows changes in flow with time. Arrows indicate the initiation of a maximal voluntary cough. Normal subjects were able to generate a large cough expiratory volume and peak cough expiratory flow with a large cough flow spike; patients with COPD were able to generate a small cough flow spike but were unable during a cough to generate cough expiratory volumes; patients with respiratory muscle weakness without scoliosis were typically unable to generate an expiratory flow spike or, if they were produced, they were greatly diminished in magnitude. The baseline flow and volume recordings of subjects with respiratory muscle weakness were similar whether or not scoliosis was present.

Table 2 Peak cough expiratory flow rate (PCEF), cough expiratory volume (CEV), and peak value time (PVT) in subjects during the various assisted cough techniques

	Normal subjects (n=9)	COPD (n=8)	Neuromuscular weakness	
			Without scoliosis (n=8)	With scoliosis (n=4)
Baseline				
PCEF (l/min)	668 (310–700)	370 (267–483)	104 (43–188)	288 (175–367)
CEV (l)	2.4 (1.31–4.91)	1.02 (0.4–2.51)	0.5 (0.3–0.8)	0.9 (0.50–1.1)
PVT (ms)	35 (30–45)	32 (25–40)	80 (40–220)	44 (40–50)
Manually assisted cough				
PCEF (l/min)	624 (326–700)	226 (120–315)*	185 (93–355)*	193 (185–287)
CEV (l)	2.91 (1.31–5.31)	0.8 (0.20–1.51)	0.7 (0.31–1.07)	0.5 (0.41–1.01)
PVT (ms)	50 (40–85)**	45 (30–60)*	118 (35–360)*	50 (35–55)
Mechanical insufflation				
PCEF (l/min)	676 (494–695)	288 (218–370)	156 (61–247)	231 (148–597)
CEV (l)	2.2 (0.8–5.91)	0.45 (0.2–0.91)*	0.6 (0.3–1.61)	0.7 (0.3–1.3)
PVT (ms)	35 (30–40)	33 (30–40)	85 (20–420)	45 (30–60)
In combination				
PCEF (l/min)	624 (288–695)	245 (218–370)*	248 (110–343)*	362 (218–440)
CEV (l)	2.2 (0.7–5.41)	0.8 (0.3–1.00)	0.6 (0.40–2.19)	0.6 (0.4–1.01)
PVT (ms)	55 (40–100)	40 (35–50)	75 (20–420)	50 (45–120)

Data are expressed as median (range).

*p<0.01.

the transient flow spike was reduced or absent (fig 2C). There was also a shorter declining phase. Peak cough expiratory flow was higher than PEF_R and the cough expiratory volume was lower than the FEV₁. The peak value time was significantly longer in the RMW group without scoliosis than in any of the other groups.

In those with COPD the transient flow spike had a rapidly declining phase similar to normal subjects (fig 2B) but the peak cough expiratory flow was significantly lower in COPD. No difference was found in the cough expiratory volume compared with the FEV₁ in patients with COPD.

The order in which the assisted cough manoeuvres were performed did not have a significant effect on the peak cough expiratory flow, cough expiratory volume, and peak value time. There were no discernible changes in baseline values of peak cough expiratory flow and cough expiratory volume with the various assisted cough manoeuvres in normal subjects (table 2) but there was a significant rise in the peak value time with manually assisted cough compared with baseline.

Those with RMW without scoliosis had a significant rise in peak cough expiratory flow with manually assisted cough alone and in combination with mechanical insufflation compared with baseline (fig 3). The peak value time with manually assisted cough was also significantly prolonged compared with baseline. No significant changes in peak cough expiratory flow, cough expiratory volume, or peak value time were noted with mechanical insufflation compared with baseline in this group.

In patients with RMW with scoliosis there were no significant changes in peak cough expiratory flow, cough expiratory volume, and peak value time compared with baseline with any of the three assisted cough manoeuvres. An example of the effects of the assisted cough manoeuvres on normal subjects, patients with COPD, and those with RMW without scoliosis on the flow recordings are shown in fig 3.

In patients with COPD there was a significant deterioration in peak cough expiratory flow compared with baseline with manually

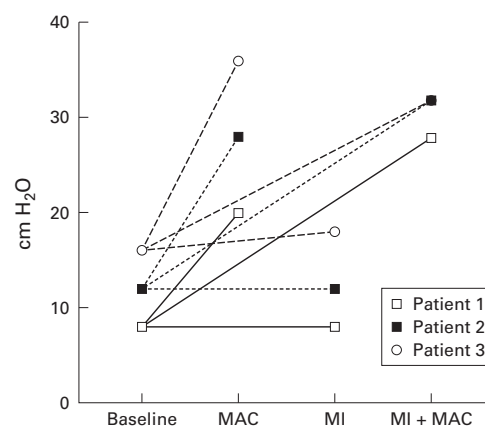


Figure 3 Effect of manually assisted cough (MAC), mechanical insufflation (MI), and both techniques in combination (MI+MAC) on oesophageal pressures during cough in comparison with baseline voluntary coughing.

assisted cough alone, mechanical insufflation alone, and in combination. The median cough expiratory volume fell with mechanical insufflation from 1.02 l to 0.45 l. There was also a significant increase in the peak value time with manually assisted cough alone over baseline values in subjects with COPD.

Oesophageal pressures during coughing in normal subjects and those with COPD exceeded 100 cm H₂O, at which pressure the balloons collapse. This made analysis of the values difficult to interpret. The three subjects with RMW all had oesophageal pressures of <50 cm H₂O. None had a significantly different peak cough expiratory flow, cough expiratory volume, or peak value time when baseline values were compared with values following balloon insertion. In these subjects with RMW manually assisted cough increased the oesophageal pressure (fig 3). No additional increase in the oesophageal pressure was seen when manually assisted cough was used in combination with mechanical insufflation.

Subjects with COPD reported no improvement with manually assisted cough and they felt mechanical insufflation alone and in combination with manually assisted cough made their cough less effective. In those with RMW without scoliosis manually assisted

Table 3 Comparisons of previous cough augmentation studies

Study	Year	Subjects	n	Apparatus	Cough adjunct	FEV ₁ (l or % predicted)	CEFR (l/min)	Poes
Present	2001	Normal	9	Pneumotachograph Poes and Pabd	MAC	3.80 (98%)	668	12***
		COPD	8		MI	0.95 (37%)	370	
		RMW with scoliosis	4		MAC + MI	0.73 (19%)	104	
		RMW without scoliosis	8			0.80 (33%)	288	
Barach et al ¹⁷	1953	Normal	15	Differential pressure*	ME	NA	465	NA
		COPD	19			251		
		Polio	1			<180		
Barach et al ²⁰	1954	COPD	53	Mask pressures	ME	NA	NA	NA
		Polio	12					
		Postoperative	8					
		Bronchiectasis	23					
Bickerman et al ²⁴	1958	Normal	18	Pneumotachograph	Isoproterenol	NA	409	NA
		Asthma	25			187		
Kirby et al ¹⁸	1966	Tetraplegics	12	Wedge spirometer	MI (50 cm H ₂ O) MAC	1.85 litres	218	NA
Leiner et al ¹⁴	1966	Normal	25	Wright peak flow meter	Isoproterenol	85.6%	597.2	NA
		Obstructive	40			56.7%	320.9	
		Restrictive	22			77.0%	387.3	
		Combined restrictive	50			47.8%	175.5	
Loudon et al ¹⁹	1967	Normal	9	Pneumotachograph Poes	Maximal voluntary cough	3.9 litres	672	167.3
		COPD	9			1.1 litres	288	153.2
Langlands et al ²⁰	1967	Normal	10	Pneumotachograph Poes and Pabd	30% citric acid	3.3 litres	487	147
		COPD	10			1.4 litres	165	180
Bach et al ²²	1993	Neuromuscular	46	Peak flow meter	MI-E	0.44	104	
Bach et al ¹⁶	1995	ALS	50 (27 successfully treated)	Peak flow meter	MI-E	0.73**	276	NA
				MAC		0.93**	150	

COPD = chronic obstructive pulmonary disease; ALS = amyotrophic lateral sclerosis; Poes = oesophageal pressure; Pabd = gastric pressure; MI = mechanical insufflation; MI-E = mechanical insufflation and exsufflation; MAC = manually assisted cough; ME = mechanical exsufflation; GPB = glossopharyngeal breathing.

*Unable to record less than 180 l/min.

**Sitting vital capacity within 3 months of initial ventilator use (second value is that of unsuccessfully treated subjects).

***Value for three subjects with respiratory muscle weakness (two without and one with scoliosis).

cough alone and in combination with mechanical insufflation improved cough clearance but mechanical insufflation alone had no effect. Subjects with RMW with scoliosis did not report a benefit from any of the assisted cough manoeuvres.

Discussion

In this study we have investigated the effect of manually assisted cough and mechanical insufflation alone and in combination in the presence of COPD, and RMW with and without scoliosis, looking for the first time at their effect on peak value time and cough expiratory volume in addition to peak cough expiratory flow. Cough flow and assisted cough techniques have been studied for over 40 years. During this period several assisted techniques have been evaluated in different subject groups (table 3) but in each study there has been a problem in the recording of peak cough expiratory flow.

Our study overcame difficulties encountered in other studies by measuring peak cough expiratory flow using a pneumotachograph linked to a digital recorder which was linear in response with regard to flow, volume, and time over the investigated data range. Other investigators have used an adapted Wright peak flow meter to measure peak cough expiratory flow but this only measures the peak expiratory flow maintained for 10 ms and fails to record the transient flow spikes so that recorded peak cough expiratory flow values are lower than actual values.¹⁴⁻¹⁶ Others have used differential pressure over a restricted orifice that was unable to record flows of less than 180 l/min which occurred in all polio patients,¹⁷ or wedge spirometry that failed to give information

about cough expiratory volume and peak value time,¹⁸ or pneumotachographs that failed to record peak cough expiratory flow accurately with an overshoot noted in testing¹⁹ or were not linear in response over 375 l/min but were still used to give a mean cough expiratory flow rate for normal subjects of 487 l/min.²⁰

The flow spikes seen on coughing in our study have been noted by other investigators using a pneumotachograph system¹⁹⁻²² or similar pressure transduction apparatus.¹⁷ The reduction or absence of flow spikes in subjects with RMW is consistent with findings when normal subjects were given d-tubocurarine to assess the effect of graded muscle weakness on peak cough expiratory flow.²¹ The transient cough expiratory flow spikes were diminished or abolished. Compression and subsequent expulsion of air from intrathoracic airways has been suggested as the cause of these transient cough spikes.²³ This intrathoracic airway compression is related to the pleural pressure which is diminished with curarisation.²¹ The cough expiratory volume was also reduced as curarisation progressed in this study.

The peak cough expiratory flow in our study was greater than PEF_R, which is in accordance with pneumotachographic studies where the same method was used to measure both¹⁸⁻²⁰ but in contrast to a study using the Wright peak flow meter in which no difference was noted.¹⁴ Cough expiratory volume was recorded as the maximum expiratory volume recorded during a cough and its duration of <1 second can explain its reduction compared with FEV₁ in normal subjects and those with RMW. Airflow obstruction in patients with COPD probably gave rise to their similar baseline cough expiratory volume and FEV₁.

In comparison with other studies the mean peak cough expiratory flow value obtained in our study was higher for our normal subjects and those with COPD and lower for our subjects with RMW without spinal abnormalities. The mean peak cough expiratory flow in normal, COPD, and RMW subjects in these pneumotachograph studies ranges from 465 l/min to 592 l/min for normal subjects,^{17 20 21} from 165 l/min to 321 l/min for subjects with COPD,^{17 20} and from <180 l/min to 218 l/min in subjects with RMW.^{17 18} The peak cough expiratory flow and cough expiratory volume have been shown to depend on lung function while the peak value time was independent of this.²⁴ The peak value time has been shown to reflect upper airway function. The prolonged peak value time in patients with unilateral vocal cord palsy was significantly reduced by vocal cord augmentation.²² The prolonged peak value time in our RMW group without scoliosis may reflect either delayed opening of the vocal cords or a delay in obtaining pressures to generate flow. All the groups increased their peak value time with manually assisted cough and, since this is independent of lung function, it is probably due to the assisted cough technique itself.²⁴ The peak value time has been shown also to be related to the timing of the maximum transpulmonary pressure (measured as the difference between oesophageal and mouth pressure) in normal subjects.²⁰ Incoordination of the thoracoabdominal thrust and the voluntary cough during manually assisted cough could have delayed increasing oesophageal pressure until after the glottis has opened, thus delaying the onset of peak cough expiratory flow. Improved coordination may have been achieved by commencing a voluntary cough once a thoracoabdominal thrust had been initiated, rather than vice versa as in our study, and this could have given rise to an even greater peak cough expiratory flow in subjects with RMW.

Subjects with COPD did not benefit with regard either to cough flow dynamics or subjectively with assisted cough techniques. They had no expiratory muscle weakness and it was not surprising that there was no improvement with manually assisted cough, but the decrease in peak cough expiratory flow and cough expiratory volume was unexpected. Manually assisted cough may cause premature airway closure in the peripheries causing the reduced peak cough expiratory flow. Mechanical insufflation may exacerbate hyperinflation of the lung in patients with COPD, and this may contribute to the reduced cough expiratory volume seen. We did not perform a PEFV after each manoeuvre and bronchoconstriction cannot be excluded as contributing to the decrease in flow.

In contrast to our study, improvements in FVC, sputum expectoration, and cough flow dynamics have been reported with other assisted cough techniques including voluntary coughing,²⁵ mechanical insufflation-exsufflation,²⁶ positive expiratory pressure,²⁷ and intrapulmonary percussive ventilator.²⁸ Voluntary coughing increases the clearance of

radiolabelled mucus from patients with airways obstruction compared with controls, and no significant improvement over voluntary coughing was noted when the forced expiration technique was used.²⁵ Mechanical insufflation-exsufflation differing from mechanical insufflation used in our study by the addition of negative pressure during coughing has been shown to improve radiolabelled suspension clearance from animal models,²⁹ increase peak cough expiratory flow in subjects with airways obstruction, and improve radiological and clinical signs in acutely ill patients with airways obstruction.²⁶

Our RMW subjects without scoliosis increased their peak cough expiratory flow with all three assisted cough manoeuvres. The rise in peak cough expiratory flow with manually assisted cough was probably due to the increased intrathoracic pressure generated by thoracic and abdominal compression. Mechanical insufflation probably raised the lung volume before coughing and thereby raised the peak cough expiratory flow. The cough expiratory volume with mechanical insufflation produced a non-significant increase. The tailoring of insufflation pressures to each patient rather than applying a standard pressure for all subjects as we used may give better results.

The use of assisted cough techniques by carers has reduced the frequency of pulmonary complications caused by retained airways secretions.¹⁵ Improvements in cough dynamics in patients with RMW have been reported using the same or alternative techniques of cough assistance using mechanical insufflation-exsufflation,^{8 26} manually assisted cough,^{8 18} trunk compression with abdominal binder,^{30 31} mechanical insufflation,¹⁸ and positive expiratory pressure.²⁷ They either aid the inspiratory phase of coughing (mechanical insufflation, glossopharyngeal breathing), or the expiratory phase of coughing (manually assisted cough, trunk compression with thoracoabdominal binder), or both (mechanical insufflation-exsufflation).

The peak cough expiratory flow was significantly greater when manually assisted cough and mechanical insufflation were combined than when mechanical insufflation alone was used. This is similar to other studies in which techniques in combination were used to increase lung volume (breath stacking, glossopharyngeal and mechanical insufflation) and the expulsive phase of coughing (mechanical exsufflation).^{15 16} In one study in patients with RMW requiring ventilatory support, manually assisted cough accompanied by either breath stacking or glossopharyngeal breathing was similar to the median increase of 144 l/min in our study with manually assisted cough and mechanical insufflation together. The same study also showed that mechanical insufflation-exsufflation improved peak cough expiratory flow to a greater extent than manually assisted cough with breath stacking or glossopharyngeal breathing.³²

Our subjects with RMW with scoliosis failed to benefit from any of the techniques used in

this study. The scoliosis made it difficult to locate the best position to place the hands for the generation of intrathoracic and intra-abdominal pressure in manually assisted cough. A higher insufflatory pressure may have shown mechanical insufflation to be beneficial in these patients. Although they suffered from alveolar hypoventilation and required non-invasive ventilatory support, their P_{imax} and P_{emax} were greater than those without scoliosis and this increased respiratory muscle strength probably contributed to the ineffectiveness of assisted cough techniques.

In conclusion, this prospective study confirms that manually assisted cough and mechanical insufflation alone or in combination improved the cough flow dynamics in those with RMW without scoliosis but not in those with scoliosis. In subjects with COPD all three assisted cough manoeuvres were detrimental to the cough flow dynamics and cannot be recommended for cough assistance. These techniques can be taught to carers and can reduce the frequency of pulmonary complications caused by retention of secretions.³¹ Mechanical insufflation should be tailored to the individual patient, taking into account the shape of the chest wall and adjusting the insufflation pressure to optimise its effectiveness.

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