Randomised, placebo-controlled trial of nebulised furosemide for breathlessness in patients with cancer

Andrew Wilcock 1, Abi Walton 1, Cathann Manderson 1, Luke Feathers 1, Bisharat El Khoury 1, Mary Lewis 1, Alpna Chauhan 1, Paul Howard 1, Sarah Bell 1, Jacky Frisby 1, Anne Tattersfield 2

Departments of:
1 Palliative Medicine
2 Respiratory Medicine
Nottingham University Hospitals NHS Trust, City Hospital Campus, Nottingham NG5 1PB, UK.

Correspondence to:
Dr Andrew Wilcock, Hayward House Specialist Palliative Cancer Care Unit, Nottingham University Hospitals NHS Trust, City Hospital Campus, Nottingham, NG5 1PB.
Email: andrew.wilcock@nottingham.ac.uk
Tel: (+44) (0)115 962 7778
Fax: (+44) (0)115 962 7779

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Word Count: 2565
ABSTRACT

Background: Breathlessness is a common and difficult symptom to treat in patients with cancer. Case reports suggest that nebulised furosemide can relieve breathlessness in such patients but few data are available.

Method: Patients with primary or secondary lung cancer and a Dyspnoea Exertion Scale score of \( \geq 3 \) were recruited. Following familiarisation patients received either nebulised furosemide 40mg or nebulised 0.9% saline under double-blind conditions or no treatment, in random order on three consecutive days. Patients undertook number reading and arm exercise tests to assess breathlessness and its impact, and were asked to report subjective benefit and any preference between nebulised treatments.

Results: Fifteen patients took part. There were no differences between furosemide, saline and no treatment in the outcomes of the number reading test (e.g. mean number read per breath was 6.7, 6.4 and 6.7 respectively) or arm exercise test (e.g. mean Borg score at maximum equivalent workload was 2.3, 2.5 and 2.7 respectively). No adverse effects were reported, although there was a small fall in FEV\(_1\) and FVC following saline. Six patients considered that their breathlessness improved with nebulised treatment, three preferring saline, one furosemide and two reporting they were of equal benefit.

Conclusions: Our findings do not support a beneficial effect from nebulised furosemide in patients with cancer-related breathlessness.

Listed on the National Research Register (N0170118249) and the UK Clinical Research Network Portfolio Database (1428).
INTRODUCTION

Breathlessness is common in patients with incurable lung cancer. About half of such patients say that breathlessness interferes with physical activities and a quarter report that it affects mood, enjoyment of life and relationships with others.[1] A systematic review supports the use of opioids to relieve breathlessness but the benefit is fairly small and adverse effects can occur.[2] New approaches are required.

In healthy volunteers nebulised furosemide 20–40mg has improved air hunger and respiratory discomfort induced by hypercapnia during constrained ventilation[3] or in combination with an inspiratory resistive load.[4] Nebulised furosemide 40mg also reduced breathlessness during endurance but not incremental exercise testing in patients with chronic obstructive pulmonary disease[5] and there are reports of benefit from nebulised furosemide 20mg in a small number of patients with cancer.[6-8] However, the only reported controlled study, a randomised double-blind cross-over pilot study in seven patients with cancer, found no difference in the difficulty or distress of breathing assessed by visual analogue scale between nebulised furosemide 20mg and 0.9% saline. Five of the seven patients said their breathing deteriorated following furosemide.[9]

The objective of the current randomised, double-blind, placebo-controlled, cross-over study was to examine for benefit of nebulised furosemide on breathlessness in patients with cancer, assessed by the number reading and arm exercise tests. A no treatment day was included as a second control to help assess the magnitude of any placebo or adverse effects resulting from nebulised saline.
METHODS

Subjects
Patients with primary or secondary lung cancer or mesothelioma with breathlessness on low levels of exertion or at rest i.e. a score of ≥3 on the Dyspnoea Exertion Scale[10] were recruited from oncology or respiratory clinics, a respiratory ward and a specialist palliative care unit. All were experiencing breathlessness that had developed or increased since cancer had been diagnosed. Patients with COPD were eligible as long as they had stable disease and their breathlessness was deemed to be predominantly cancer-related. Patients were excluded if breathlessness could be relieved by treatment such as drainage of a pleural effusion or blood transfusion. Other reasons for exclusion included radio- or chemotherapy within four weeks, asthma, angina, heart failure, or any problem that might affect the patient’s ability to read aloud or undertake arm exercise. The dose of any drug that could potentially affect exercise or breathlessness had to have been stable for at least four days prior to the study. Patients gave written informed consent and the study was approved by Nottingham City Hospital research ethics committee, and by the National Cancer Research Network as a locally adopted study.

Measurements

Dyspnoea Exertion Scale
The Dyspnoea Exertion Scale, a modified version of the Medical Research Council Dyspnoea Scale, was used to select and categorise patients.[10] It ranges from 0 = ‘I am able to walk at my own pace on the level without getting breathless over any distance’ to 5 = ‘I am breathless at rest’.

Spirometric values
Forced expiratory volume in one second (FEV1) and forced vital capacity (FVC) were measured with the patients seated as the best of three recordings using a dry wedge spirometer (Vitalograph Type R Spirometer, Buckingham, UK) or hand-held spirometer (Micro Spirometer, Micro Medical Ltd, Rochester, Kent, UK), depending on whether the patient was studied in the palliative care unit or a hospital side room. Patients used the same device on each day.

Number reading test
This has been developed to measure the limiting effect of breathlessness on reading numbers aloud at rest.[11] While seated patients are given a page containing a grid of numbers and asked to read the numbers aloud and in order as quickly and clearly as they can. The number of breaths taken and the number of numbers read over 60 seconds is recorded. The procedure is repeated five times using the same grid of numbers. Patients are allowed to recover between readings and continue when they feel rested. The highest number obtained from the five readings and the number of numbers read per breath during this reading are noted.

Arm exercise test
This has been developed to assess breathlessness in patients with cancer who are breathless at low levels of exertion.[12] While seated patients are
asked to move an outstretched arm between two points 20cm above and
20cm below shoulder height in time to a regular audible beat (80 beats min\(^{-1}\))
with no verbal encouragement. At one-minute intervals patients are asked to
quantify their sensation of breathlessness by pointing to a modified Borg
scale\[13\] and then switch to exercising the other arm. The Borg scale is a
vertical scale labeled from 0 to 10 with corresponding expressions of
increasing sensation intensity from 'nothing at all' to 'maximal.'

Breathlessness was defined to the patients as “a feeling of an uncomfortable
need to breathe rather than other sensations associated with exercise, such
as fatigue or the awareness that ventilation had simply increased.” Patients
are asked to continue the exercise for as long as possible. Breathlessness
scores at the maximum duration of exercise attained by each patient in all
three tests were used in the analysis.

**Protocol**

Tests were carried out in a quiet room in the palliative care unit or a ward side
room. Patients were given written instructions to avoid caffeine for one hour,
large meals for two hours and excess alcohol from the night before the tests.
Medication and times of drug administration were unchanged during the
study, which ran over four days. The initial day was for familiarisation with the
equipment, surroundings and procedure. On the next three days, at the same
time of day, patients undertook the assessments following nebulised
furosemide 40mg or 0.9% saline under double-blind conditions, or no
treatment. An independent pharmacist prepared the nebulised treatments
following a treatment order generated using randomised permuted blocks
containing the six possible orders of treatment. Furosemide and saline (both
4ml) were given by jet nebuliser (HOT Top Plus, Intersurgical Ltd.,
Wokingham, UK; particle mass median diameter 3.25 microns) attached to a
facemask and driven with air at 8L/minute. The nebuliser was attached to a
facemask and patients were instructed to take slow deep breaths through
their mouth until one minute after the nebuliser started to splutter
(approximately 5 minutes). Spirometry was measured immediately before and
after the nebulised treatment ended. After 10 minutes rest patients underwent
the number reading test and, after a further 10 minutes rest, the arm exercise
test. On the no treatment day patients underwent spirometry followed by the
number reading and arm exercise tests as above. Patients emptied their
bladder immediately before each study and urine output was measured over
the next two hours. The patient was allowed one 125ml drink during this
period. At the end of the study patients were asked if they had perceived any
benefit from the nebulised treatment and if so, any difference between the two
treatments.

The primary outcome was the two endpoints of the number reading test,
namely the total number read and the number read per breath. Our previous
within subject, between day, data indicated that nine and 15 patients
respectively would be required to detect a change in these two endpoints
(90% power; \(p = 0.05\)) equivalent to 50% of that seen following drainage of a
pleural effusion.\[11\] We had hoped to recruit 30 patients but, due to slow
accrual, recruitment was discontinued after reaching the minimum number.
Statistical analysis
Repeated measures analysis of variance was used to test for carry over, period and for treatment effects to establish the within subject difference in the mean values of number of numbers read, the number of numbers read per breath, the modified Borg score at maximum equivalent work load, the duration of arm exercise and urine output between the three treatment groups. Change in spirometric values following nebulised furosemide and saline were compared by paired t-test. Calculations were performed using Statistical Package for the Social Sciences (SPSS) version 14.0. A p value of <0.05 was regarded as statistically significant.
RESULTS
Recruitment commenced in October 2002 and was stopped when 15 patients (8 female; mean (SD) age 66 (11) years) had completed the study (Figure). Their median (range) Eastern Co-operative Oncology Group (ECOG) performance status was 2 (1–3). All had thoracic cancer, either primary (non-small cell lung (7), mesothelioma (2)), or secondary (breast (2), one each from pancreas, thymus, uterus and unknown). Regular medication included opioids (13), benzodiazepines (4) and oxygen (3). Seven patients were receiving bronchodilators, although only one had a documented diagnosis of COPD. Mean (SD) FEV₁/FVC ratio was 75 (11)%. The Dyspnoea Exertion Scale categorised three, eight and four patients as experiencing breathlessness moving around in bed or getting out of bed (level 3), on talking (level 4) or at rest (level 5) respectively. To date, 13 patients have died with a median (range) survival of 51 (5–353) days.

All patients were able to complete all the number reading tests satisfactorily. Two patients were unable to complete the arm exercise tests, despite a satisfactory familiarisation test, and could not be included in this analysis.

No carry over or period effect was found for the number of numbers read per minute (p = 0.2 and p = 0.09 respectively) or per breath (p = 0.55 and p = 0.73 respectively). Across the three study days, patients read a mean (range) of 70 (43–103) numbers per minute, and 6.1 (2.3–35) numbers per breath. There were, however, no significant differences between the number of numbers read per minute (68, 70, 70; p = 0.91) or per breath (6.7, 6.4, 6.7; p = 0.64) between the three treatments (Tables 1 and 2).

All 13 patients who completed the arm exercise test experienced an increase in breathlessness during exercise apart from one who scored breathlessness as ‘very slight’ throughout. Across the three study days, the mean duration of arm exercise was 457 seconds (7.62 minutes), representing a mean of 609 arm movements. There were, however, no significant differences between the duration of arm exercise (452, 485, 467 seconds; p = 0.96) or Borg score at maximum equivalent workload (2.3, 2.5, 2.7; p = 0.83) between the three treatments (Tables 1 and 2).
Table 1. Number reading and arm exercise test outcomes, spirometric values and urine output following nebulised furosemide or nebulised saline and on a no treatment study day. Values are mean (SD).

<table>
<thead>
<tr>
<th></th>
<th>Furosemide</th>
<th>Saline</th>
<th>No treatment</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Number reading test (n = 15)</strong></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Number of numbers read</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>68 (16)</td>
<td>70 (15)</td>
<td>70 (16)</td>
<td>0.91</td>
</tr>
<tr>
<td>Per breath</td>
<td>6.7 (7.9)</td>
<td>6.4 (5.1)</td>
<td>6.7 (5.9)</td>
<td>0.64</td>
</tr>
<tr>
<td><strong>Arm exercise test (n = 13)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Duration of arm exercise (seconds)</td>
<td>452 (349)</td>
<td>485 (345)</td>
<td>467 (342)</td>
<td>0.96</td>
</tr>
<tr>
<td>Borg Score at maximum equivalent work load</td>
<td>2.3 (1.5)</td>
<td>2.5 (1.4)</td>
<td>2.7 (1.9)</td>
<td>0.83</td>
</tr>
<tr>
<td><strong>Change in spirometric values (n = 15)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FEV₁ (ml)</td>
<td>-10 (80)</td>
<td>-80 (90)</td>
<td>-</td>
<td>0.03</td>
</tr>
<tr>
<td>FVC (ml)</td>
<td>10 (100)</td>
<td>-70 (80)</td>
<td>-</td>
<td>0.02</td>
</tr>
<tr>
<td><strong>Urine output (ml; n = 10)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>202 (124)</td>
<td>203 (131)</td>
<td>199 (150)</td>
<td>0.99</td>
</tr>
</tbody>
</table>

Table 2. Between-treatment differences (95% confidence intervals) for the number reading and arm exercise test outcomes, spirometric values and urine output.

<table>
<thead>
<tr>
<th></th>
<th>Furosemide versus no treatment</th>
<th>Saline versus no treatment</th>
<th>Furosemide versus saline</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Number reading test (n = 15)</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of numbers read</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>1.1 (-1.4, 3.7)</td>
<td>-1.3 (-4.9, 2.2)</td>
<td>2.4 (-1.6, 6.5)</td>
</tr>
<tr>
<td>Per breath</td>
<td>0.1 (-0.9, 1.1)</td>
<td>0.5 (-0.4, 1.4)</td>
<td>-0.5 (-1.2, 0.3)</td>
</tr>
<tr>
<td><strong>Arm exercise test (n = 13)</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Duration of arm exercise (seconds)</td>
<td>27 (-19, 71)</td>
<td>13 (-54, 26)</td>
<td>40 (-17, 98)</td>
</tr>
<tr>
<td>Borg Score at maximum equivalent work load</td>
<td>0.4 (-0.2, 1)</td>
<td>0.2 (-0.3, 0.8)</td>
<td>0.2 (-0.2, 0.6)</td>
</tr>
<tr>
<td><strong>Change in spirometric values (n = 15)</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>FEV₁ (ml)</td>
<td>-</td>
<td>-</td>
<td>-68 (-122, -15)</td>
</tr>
<tr>
<td>FVC (ml)</td>
<td>-</td>
<td>-</td>
<td>-87 (-153, -20)</td>
</tr>
<tr>
<td><strong>Urine output (ml; n = 10)</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>-4 (-92, 84)</td>
<td>-5 (-83, 72)</td>
<td>-2 (-100, 97)</td>
</tr>
</tbody>
</table>

Following nebulised saline there was a fall in FEV₁ and FVC of 8% and 5% from baseline respectively. After nebulised furosemide the changes in FEV₁ and FVC were 1% or less. The differences between the two treatments (70ml and 80ml) were statistically significant (Tables 1 and 2). No difference was
found in urine output in the 10 patients with a reliable urine collection (Tables 1 and 2). No adverse effects were reported by the patients following nebulised furosemide or saline.

Six of the 15 patients considered that their breathlessness improved with nebulised treatment, with three preferring saline, one furosemide and two finding the nebulised treatments of equal benefit.
DISCUSSION
In this randomised controlled study in 15 patients with cancer-related breathlessness we found no evidence of benefit from nebulised furosemide 40mg on any outcome measure. Only one of six patients who considered their breathlessness to have improved with nebulised treatment chose nebulised furosemide over 0.9% saline. Our findings do not support a beneficial effect of nebulised furosemide in this patient group.

Breathlessness is a distressing symptom for many patients with thoracic cancer and few treatments are available. Despite this, there is a paucity of research into finding new treatments. Undertaking formal trials in these patients is challenging and recruitment is difficult. Assessing breathlessness is also difficult, particularly when it is experienced at such low levels of exertion as to make most forms of exercise testing impractical. The number reading test and the arm exercise test were developed to help assess breathlessness in patients with cancer who are breathless on minimum exertion. The former is a measure of the limiting effect of breathlessness on reading numbers aloud at rest and the latter allows breathlessness to be measured in patients who become breathless on low-level exertion. We have shown that both are acceptable to patients and repeatable.[11,12] The number reading test was used as the primary outcome measure and our sample size was sufficient to detect a change equivalent to 50% of that seen following a thoracocentesis (mean volume 1840ml).[11]

The negative findings in our study differ from some previous reports in patients with breathlessness related to cancer. Sixteen of the 19 patients described in three reports benefited from nebulised furosemide 20mg administered as a single dose (13 patients)[6,8] or 20mg four times a day for up to three weeks (three patients).[7] However, the only controlled study of nebulised furosemide in seven patients with cancer-related breathlessness showed no benefit, in keeping with our findings.[9] The benefit seen in the case reports may therefore be a placebo effect. Differences in the underlying lung pathology causing cancer-related breathlessness is another possible explanation, although this varied considerably in the case reports as it did in our patients. Nebulised furosemide has also reduced experimentally-induced breathlessness in healthy subjects[3,4] but the relevance of this to patients with cancer is uncertain.

In our study there was a small fall in FEV₁ and FVC after nebulised saline of 80ml and 70ml respectively. Bronchoconstriction following nebulised 0.9% saline has been seen in patients with asthma and COPD and attributed to a non-specific bronchoconstrictor response to airway cooling.[14,15] The fact that the change was less following nebulised furosemide is in keeping with furosemide’s ability to protect against such non-specific bronchoconstrictor stimuli.[16]

Our finding contrasts with the small increase in FEV₁ (50ml) seen following nebulised furosemide 40mg in 20 patients with COPD undergoing an incremental exercise test.[5] However, bronchodilatation in this study may have been due to the exercise[17] rather than nebulised furosemide, as the
authors suggest. Although nebulised furosemide is very effective in inhibiting bronchoconstriction due to antigen and non-specific stimuli in patients with asthma[16] it has not caused bronchodilatation in these studies.

Although larger doses or more frequent administration of nebulised furosemide may have been more effective, the dose we gave is larger than the 20mg dose associated with benefit in 12 of the 15 patients reported by Kohara et al,[6] and similar doses are sufficient to inhibit bronchoconstriction in patients with asthma.[18] We also considered whether our outcome measures were insufficiently sensitive to detect a beneficial effect, but the lack of patient preference for nebulised furosemide suggests we did not miss a clinically important difference.

Thus our findings do not support a beneficial effect from a single dose of 40mg nebulised furosemide in patients with cancer-related breathlessness.
ACKNOWLEDGEMENTS
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COMPETING INTERESTS
None of the authors have any conflict of interest with any aspect of submitting this manuscript for publication.

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FIGURE LEGEND
Figure. Study flow diagram
REFERENCES


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Figure. Study flow diagram

Eligible patients: 43

Not enrolled 26:
• declined, 23
• deteriorated, 3

Enrolled: 17

Withdrew after familiarisation 2:
• unable to exercise arm, 1
• deteriorated, 1

Randomised: 15

Treatment order:
Furosemide, saline, no treatment, 3
Furosemide, no treatment, saline, 2
No treatment, saline, furosemide, 3
No treatment, furosemide, saline, 2
Saline, no treatment, furosemide, 3
Saline, furosemide, no treatment, 2

Analysis:
• number reading and spirometry, 15
• arm exercise test, 13
  - unable to complete test, 2
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