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- 1 The impact of outpatient vs inpatient management on health-related
- 2 quality of life outcomes for patients with malignant pleural effusion -
- 3 the OPTIMUM randomized clinical trial.
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# 75 Abstract

76

### 77 Background

78 The principal aim of malignant pleural effusion (MPE) management is to improve health related

79 quality of life (HRQoL) and symptoms.

### 80 Methods

- 81 In this open-label randomised controlled trial, patients with symptomatic MPE were randomly
- 82 assigned to either IPC insertion with the option of talc pleurodesis or chest drain and talc
- 83 pleurodesis. The primary endpoint was global health status, measured with the EORTC QLQ-C30
- 84 questionnaire at 30 days post-intervention. 142 participants were enrolled from July 2015 to
- 85 December 2019.

### 86 *Results*

87 Of participants randomly assigned to IPC (n=70) and chest drain (n=72), primary outcome data were 88 available in 58 and 56 patients, respectively. Global health status improved in both groups at day 30 89 compared to baseline: IPC (mean difference 13.11 p=0.001) and chest drain (mean difference 10.11 90 p=0.001). However, there was no significant between-group difference at day 30 (mean inter-group difference in baseline-adjusted global health status of 2.06 ([95% CI -5.86 to 9.99]; p = 0.61), day 60 91 92 or day 90. No significant differences were identified between groups in breathlessness and chest 93 pain scores. All chest drain arm patients were admitted (median length of stay 4 days); 7 in the IPC 94 arm required intervention-related hospitalization.

### 95 Conclusion

- While HRQoL significantly improved in both groups, there were no differences in patient reported
  global health status at 30 days. The outpatient pathway using an IPC was not superior to inpatient
  treatment with a chest drain.
- 99 Trial Registration
- 100 ISRCTN registration:15503522.

### 102 Introduction

Malignant pleural effusions (MPE) result in breathlessness, reduced function and impaired health related quality-of-life (HRQoL), often representing an advanced terminal illness with a median

survival of 3-12 months[1].

106 Two definitive management options for MPE include hospital admission for a chest drain insertion

107 with talc slurry pleurodesis or outpatient ambulatory management with an indwelling pleural

108 catheter (IPC). Over the last decade, IPCs have increasingly become a first-line intervention for MPE

109 with recent advances incorporating talc pleurodesis[2] or a daily drainage strategy[3]. These

approaches may stop fluid production via pleural symphysis. However, these two options may exert

distinct influences on patients' quality of life. For example, an IPC shifts the burden of care to the

112 community, patient and their carers, and involves a period of repeated drainage and frequent

healthcare visits, whilst a chest drain and talc pleurodesis entails a median hospital stay of 4 days[4].

114 Although improving HRQoL is a central treatment goal, a systematic review has identified limited

115 comparative data on HRQoL outcomes to guide best practice[5]. This trial tests the hypothesis that

116 outpatient management of MPE utilising an IPC with the option of talc pleurodesis improves HRQoL

117 compared to usual inpatient management with a chest drain and talc pleurodesis.

118 This is the first randomised controlled study to evaluate HRQoL as a primary outcome measure in

119 MPE intervention.

120

- 122 Methods
- 123 Trial Design

The Out Patient Talc Slurry via Indwelling Pleural Catheter for Malignant Pleural Effusion Vs Usual
Inpatient Management (OPTIMUM) trial is a randomised, two-arm, open-label superiority trial
conducted at 11 hospitals in the United Kingdom and one hospital in Australia. UK ethics approval
was obtained from the National Research Ethics Service (NRES) Committee South East Coast,
Brighton and Sussex (15/LO/1018). For Australia, approval was obtained from the Sir Charles
Gairdner Group Human Research Ethics Committee. The trial protocol has been published[6]
(Supplement 1).

#### 131 Participants

132 All participants were adults diagnosed with MPE made either by histocytological confirmation or

- clinical and radiological features of metastatic pleural disease in patients with histologically proven
- primary cancers. Participants were required to have a WHO performance status of two or less,
- unless a performance status of three was likely to improve with pleural drainage. Participants also

136 needed an expected survival of greater than three months. The exclusion criteria were age less than

- 137 18 years old; pregnant or lactating; known allergy to talc or lidocaine; lack of symptomatic relief
- 138 from effusion drainage; district nurse/carers/hospital team unable to carry out at least twice weekly
- 139 IPC drainage; underlying lymphoma or small cell carcinoma except if chemotherapy had failed or the
- 140 patient was to be referred for palliative management; non-malignant effusions; loculated pleural
- 141 effusion; and patients unable to provide written informed consent to trial participation. Participants
- 142 were screened from both the outpatient and inpatient setting.

#### 143 Randomization

- 144 Participants were randomised 1:1 to either IPC insertion or chest drain and talc slurry pleurodesis.
- 145 Permuted block randomisation (block sizes 4, 6, 8) was performed with allocation concealment
- 146 maintained using a web-based randomisation service (<u>www.sealedenvelope.com</u>). Treatment
- allocation was unblinded and stratified to the following factors: age (<65 years,  $\geq$ 65 years),
- 148 malignancy subtype (lung, mesothelioma, breast, other) and WHO performance status (0, 1, 2, 3).
- 149 Patient blinding was not practical due to inherent differences between the interventions.
- 150
- 151 Procedures

152 Supplementary figures s1-s2 (section 3 in Supplement 2) provides further detail of the trial

153 interventions.

154 Participants randomised to the IPC group underwent catheter insertion as a day case under local 155 anaesthesia using a percutaneous Seldinger technique. Following insertion, an attempt was made to 156 evacuate the fluid completely, using pleural manometry where available, to enable safe large-157 volume drainage and reduce the risk of re-expansion pulmonary oedema. Patients then returned 158 after three days for review and a further attempt at maximal pleural fluid drainage. If the fluid 159 removed at this visit averaged less than 150mls/day since IPC insertion and non-expandable lung 160 was ruled out by ultrasound or X-ray, talc pleurodesis was attempted through the IPC. The protocol 161 was amended in October 2016 to remove the <150mls/day drainage criteria as feedback from 162 recruiting centres suggested incomplete drainage at insertion made fluid estimation at the day 4 visit 163 difficult. Four grams of sterile talc was then administered as a slurry (section 1.1.1 in Supplement 2) 164 and participants observed for 1 hour following instillation. Patients or district nurses were advised to 165 perform daily IPC drainages as an 'aggressive' drainage strategy using 1 litre bottles and return at 166 day 7 for review, chest X-ray and repeat drainage. If satisfactory pleural apposition with absence of 167 pleural sliding was seen on ultrasound in 5 of 6 areas (Figure s3 in Supplement 2), repeated drainage 168 was halted. Participants returned on day 14 for review and the IPC removed if pleural apposition was 169 maintained with evidence of minimal fluid on ultrasound.

170 In cases where talc instillation was either not attempted or in participants that did not meet the

171 criteria for IPC removal at day 14, regular IPC drainage was continued throughout the study follow

172 up period. The frequency of drainage was at the discretion of the treating physician. For these

173 participants, the recommended approach was early assessment in the clinic if they experienced

three consecutive drainages of less than 50mls of fluid. If either a chest x-ray or ultrasound showed

175 no significant residual pleural collection, then the IPC could be removed.

Participants randomised to chest drain and talc pleurodesis underwent management as per the 2010
British Thoracic Society (BTS) pleural disease guidelines[7]. A 12F-14F chest drain was inserted using
ultrasound guidance under local anaesthesia, and the patient was admitted to hospital. After 24hrs,
if the chest X-ray ruled out an non-expandable lung, 4 grams of talc slurry was instilled. The drain
was then removed when the pleural fluid output dropped below 250mls/day. Patients with nonexpandable lung or any other contraindication to talc slurry pleurodesis (such as an air leak) could be

182 managed using an alternative strategy and continued their follow up in the study.

### 183 Follow-up

184 All participants underwent follow up until 90 days after intervention or death, whichever occurred

185 first. Trial visits were conducted at the outpatient clinic on day 4 for the IPC group and 7, 14, 30, 60

and 90 days after the intervention for both groups. If fluid recurrence or complications developed,

the trial clinicians were permitted to perform further tests and procedures (e.g. IPC insertion orreferral for thoracic surgery) as part of usual clinical care.

#### 189 *Outcomes*

#### 190 Primary Outcome

The primary outcome was global health status, at 30 days post-intervention measured with the European Organisation for Research and Treatment of Cancer Quality of Life Questionnaire–Cancer 30 (EORTC QLQ C-30). This is a validated, cancer specific, multidimensional instrument that asks participants to report on aspects of their health-related quality of life over the previous week. It is suitable for all cancer diagnoses[8] and can be repeated at frequent intervals to monitor quality of life over time[9].

#### **197** Secondary Outcomes

198 Secondary outcomes included global health status at day 60 and day 90 post-intervention, adverse

- 199 event rates, breathlessness, chest pain scores and pleurodesis failure rate. Pleurodesis failure was
- 200 defined as chest X-ray opacification greater than 25% on the side of intervention or the need for
- subsequent pleural intervention on the same side as pleurodesis at 30, 60 and 90 days post-
- 202 intervention. Breathlessness and chest pain scores at 30, 60 and 90 days post-intervention were
- 203 measured using the 0-100mm visual analogue scale.

#### 204 Statistical analysis

205 Using an analysis of covariance model (ANCOVA), adjusting for baseline global health status, a

sample size of 142 (71 vs 71) would detect a clinically significant difference of 8 points in global

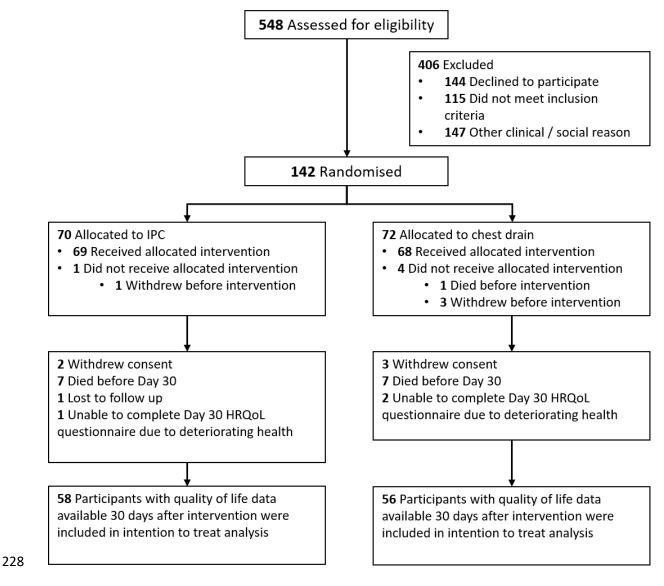
207 health status with 80% power and a 5% significance level. This assumes a common standard

- 208 deviation of 23.6 in Stage III-IV cancer[8]. The covariate has an R-squared of 0.49. An interim analysis
- 209 was planned once 50% of patients were enrolled to determine if the recruitment target needed210 amendment.
- 211 At this interim analysis, a blinded assessment of data demonstrated a 14% loss to follow-up at 30
- 212 days. The trial steering committee agreed that given funding, recruitment rates and the nature of
- the patient population, increasing the sample size would not be feasible and agreed to continue
- 214 recruitment until December 2019 to achieve the original trial objective.
- 215 Data were analysed on an intention-to-treat basis in all patients in whom outcome data were
- available. The primary efficacy analysis was based on ANCOVA. Regression analyses were adopted to
- 217 compare estimates for secondary efficacy analyses.

- 218 Data attrition for the complete case analysis prompted a post-hoc sensitivity analysis. Multiple
- 219 imputation by chained equations was used to impute the missing data[10]. Using mice (Multivariate
- 220 Imputation by Chained Equations) R package[11], 300 simulated datasets were generated, and an
- ANCOVA analysis was performed on all of the imputed datasets with the results pooled. All analyses
- were conducted using R version 4.0. The trial statistical analysis plan is outlined in Supplement 3.
- 223 The trial was registered on the ISRCTN registry (Identifier: 15503522).

# 224 Results

- 225 After screening a total of 548 patients for eligibility, 142 participants were recruited between July
- 226 2015 and November 2019. Four patients withdrew, and one died before undergoing the randomised
- intervention. They were excluded from the analyses (Figure 1).



- 229 Figure 1: Consort diagram for OPTIMUM Study.
- 230 Abbreviations: HRQoL health related quality of life; IPC indwelling pleural catheter.

- 232 Baseline characteristics
- 233 Of the 142 randomised patients, two patients that withdrew following randomisation did not
- consent to baseline data collection. Baseline data for both groups are presented in Table 1.
- 235

		IPC (n=70)	Chest Drain (n=70)
Age, mean (SD), yr	6	69.0 (12.5)	66.5 (12.7)
Female : Male	3	38 : 32	42 : 28
Tobacco smoking status (%) Current smoker Former smoker Never smoked Unknown	3	5 (8.6%) 36 (51.4%) 28 (40%)	5 (7.1%) 45 (64.2%) 19 (27.1%) 1 (1.4%)
Side of intervention (%) Right Left		48 (68.6%) 22 (31.4%)	39 (55.7%) 31 (44.3%)
Bilateral pleural effusion (%)	1	12 (17.1%)	13 (18.6%)
WHO performance status 0 1 2 3	3	9 (12.8%) 32 (45.8%) 23 (32.9%) 5 (8.6%)	5 (7.1%) 32 (45.8%) 26 (37.1%) 7 (10%)
Malignancy Lung Breast Mesothelioma Renal Ovarian Unknown Primary Colorectal Upper gastrointestinal Uterine Other		22 (31.4%) 18 (25.7%) 9 (12.9%) 5 (7.1%) 5 (7.1%) 4 (5.7%) 2 (2.9%) 1 (1.4%) 1 (1.4%) 3 (4.3%)	21 (30%) 15 (21.4%) 12 (17.1%) 4 (5.7%) 3 (4.3%) 4 (5.7%) 2 (2.9%) 2 (2.9%) 3 (4.3%) 4 (5.7%)
Duration of cancer diagnosis at the time of recruitment, mean (SD), months	2	26.1 (40.8)	15.6 (28.3)
Treatment at enrolment Chemotherapy Targeted (e.g. HER2, ALK, EGFR, multikinase) Hormonal Immunotherapy		n=29 13 (18.6%) 9 (12.9%) 4 (5.7%) 3 (4.3%)	n=27 13 (18.6%) 6 (8.6%) 5 (7.1%) 3 (4.3%)
Steroid therapy at baseline	ç	9 (12.9%)	5 (7.1%)

# 237 Table 1: Summary of baseline characteristics

Size of effusion on chest radiograph		
<25% hemithorax	7 (10%)	12 (17.1%)
25-50% hemithorax	25 (35.7%)	29 (41.4%)
>50% hemithorax	34 (48.6%)	25 (35.7%)
EORTC QLQ-C30 global health	n=70	n=69
status at baseline, mean (SD)	37.3 (25.4)	37.8 (25.4)
100mm VAS breathlessness score	n=68	n=67
at baseline, mean (SD)	60.8 (26.0)	50.3 (28.4)

Abbreviations: WHO – World Health Organization; HER2 - human epidermal growth factor receptor 2, ALK - anaplastic
 lymphoma kinase; EGFR - epidermal growth factor receptor; EORTC QLQ-C30 - European Organisation for Research and
 Treatment of Cancer Quality-of-life Questionnaire Core 30; VAS – Visual Analogue Scale.

242

## 243 *Primary Outcome*

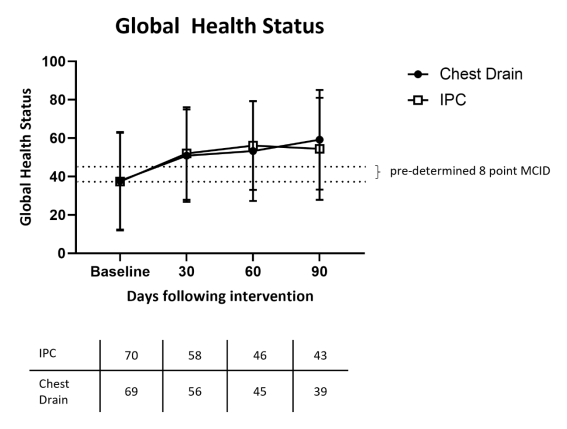
Data were available in 58 and 56 patients in the IPC and chest drain groups, respectively. Global

- health status improved significantly at day 30 post-intervention (compared with the baseline) in
- both the IPC group (mean difference 13.11 [95% CI 5.6 to 21.1]; *p* = 0.001) and the chest drain group
- 247 (mean difference 10.11 [95% CI 4.5 to 15.7]; *p* = 0.001). 57% (33 of 58) of patients in the IPC group
- and 54% (30 of 56) in the chest drain group experienced a greater than 8 point improvement in
- 249 global health status. Mean global health status at day 30 (primary endpoint) was 52.0 (SD 24.1) in
- the IPC group and 50.9 (SD 24.1) in the chest drain group with an observed mean difference at 30
- days of 2.06 ([95% CI -5.86 to 9.99]; p = 0.61) when adjusted for baseline global health status as a
- 252 covariate.
- 253 Findings from the sensitivity analysis remained consistent with the primary analysis; at 30 days, a
- mean difference (IPC vs drain) in global health status of 2.18 ([95% Cl -5.62 to 9.99]; p = 0.59) was
- 255 observed.
- 256 Secondary Outcomes
- 257 Secondary outcome data are summarised in Table 2, Figures 2, and 3.
- 258 Quality of Life at 60 and 90 days
- 259 At 60 days after the intervention, global health status data were available in 46 patients in the IPC
- 260 group and in 45 patients in the chest drain group. The mean change in global health status from
- 261 baseline at day 60 was 15.6 (SD 26.4) in the IPC arm (*n* = 46), and 7.96 (SD 26.9) in the chest drain
- arm (*n* = 45). The ANCOVA, (adjusted for baseline global health status) indicated an observed mean
- 263 difference of 4.82 ([95% CI -4.59 to 14.23]; *p* = 0.31).

- At day 90 after the intervention, 43 patients completed follow up in the IPC arm versus 39 in the
- chest drain group. The mean change in global health status from baseline at day 90 was 13.4 (SD
- 266 30.6) in the IPC arm, and 14.93 (SD 25.1) in the chest drain arm. The ANCOVA (adjusted for baseline
- 267 global health status) indicated an observed mean difference of -3.12 ([95% CI -13.76 to 7.51]; p =
- 268 0.56).

269 Pleurodesis Failure

- Figure 3 summarises outcomes related to talc pleurodesis and subsequent pleural intervention inboth treatment arms.
- Twenty-nine of 65 patients (44.6%) in the IPC arm received talc slurry vs 49 of 67 patients (73.1%) in
- the chest drain arm. The incidence of non-expandable lung (defined as <50% pleural apposition
- following drainage) was similar in both groups (IPC 15 (23.0%) vs chest drain 16 (23.9%)).
- 275 Thirteen participants in the IPC and talc instillation subgroup were eligible for IPC removal at Day 14,
- 276 however 10 participants underwent IPC removal with the remaining three electing not to have the
- 277 IPC removed. The rate of pleurodesis failure at 30 days in the IPC group (*defined as IPC remaining in*
- situ, need for subsequent pleural intervention or chest x-ray opacification of >25% hemithorax) was
- 279 64.3% (18 of 28), which includes the 3 participants that declined catheter removal at day 14. 4 IPCs
- 280 were removed after day 30 due to pleurodesis: 1 IPC was removed after day 30 and 3 IPCs were
- removed after day 60. The pleurodesis failure rate at day 60 was 64.3% (18 of 28). At day 90, this
- 282 was 57.1% (16 of 28).
- 283 Pleurodesis failure in the chest drain group (defined as a need for subsequent pleural intervention or
- 284 chest x-ray opacification of >25% hemithorax) at 30 days was 18.4% (9 of 49). At day 60 this was
- 285 24.5% (12 of 49). At day 90 this was 26.5% (13 of 49).
- A summary of patients that underwent an additional pleural intervention is provided in section 2.1
- of Supplement 2.



289 Figure 2: Global health status scores measured using the EORTC QLQ C-30 over the 90-day follow-up

290 period. Points represent the mean, and bars represent standard deviation. Higher values indicate

291 better global health status. Table below graph denotes the number of patients with global health

292 status data at each time point. IPC – indwelling pleural catheter.

## 294 Table 2: Secondary outcome results

	Chest drain arm ( <i>n</i> = 72) <sup>ª</sup>	IPC arm ( <i>n</i> = 70) <sup>a</sup>	Treatment effect estimate (95% Cl)	<i>p</i> -value
Change in global health status from baseline, mean (SD)	( <i>n</i> = 69 with ≥ 1 measurement)	( <i>n</i> = 70 with ≥1 measurement)	Absolute difference <sup>b</sup>	
Baseline	37.8 (25.4)	37.3 (25.4)		
Day 60	8.0 (26.9)	15.6 (26.4)	4.82 (-4.59 to 14.23)	0.31
Day 90	14.9 (25.1)	13.4 (30.6)	-3.12 (-13.76 to 7.51)	0.56
Change in VAS breathlessness score from baseline, mean (SD), mm	(n = 67 with ≥ 1 measurement)	( <i>n</i> = 68 with ≥1 measurement)	Absolute difference <sup>c</sup>	
Baseline	50.3 (28.4)	60.8 (26.0)		
Day 30	-19.9 (30.4)	-34.3 (28.4)	-6.8 (-15.97 to 2.41)	0.15
Day 60	-14.4 (32.3)	-30.3 (30.3)	-5.3 (-16.67 to 6.15)	0.37
Day 90	-17.3 (33.0)	-23.1 (41.4)	9.6 (-3.88 to 23.11)	0.17
Change in VAS chest pain score from baseline, mean (SD), mm	( <i>n</i> = 67 with ≥ 1 measurement)	( <i>n</i> = 68 with ≥1 measurement)	Absolute difference <sup>c</sup>	
Baseline	20.7 (27.1)	22.1 (27.1)		
Day 30	-1.9 (24.1)	-7.5 (24.7)	-2.84 (-9.69 to 4.02)	0.42
Day 60	-0.6 (22.0)	0.6 (30.0)	3.7 (-5.01 to 12.44)	0.41
Day 90	-4.27 (25.3)	2.8 (34.1)	8.7 (-2.79 to 20.17)	0.14
Pleurodesis failure, number (%)	( <i>n</i> = 49)	( <i>n</i> = 28)	Unadjusted odds ratio	
Day 30	9 (18.4%)	18 (64.3%)	8.0 (2.77 to 23.1)	<0.001
Day 60	12(24.5%)	18 (64.3%)	5.55 (2.02 to 15.25)	<0.001
Day 90	13 (26.5%)	16 (57.1%)	3.7 (1.38 to 9.8)	0.01

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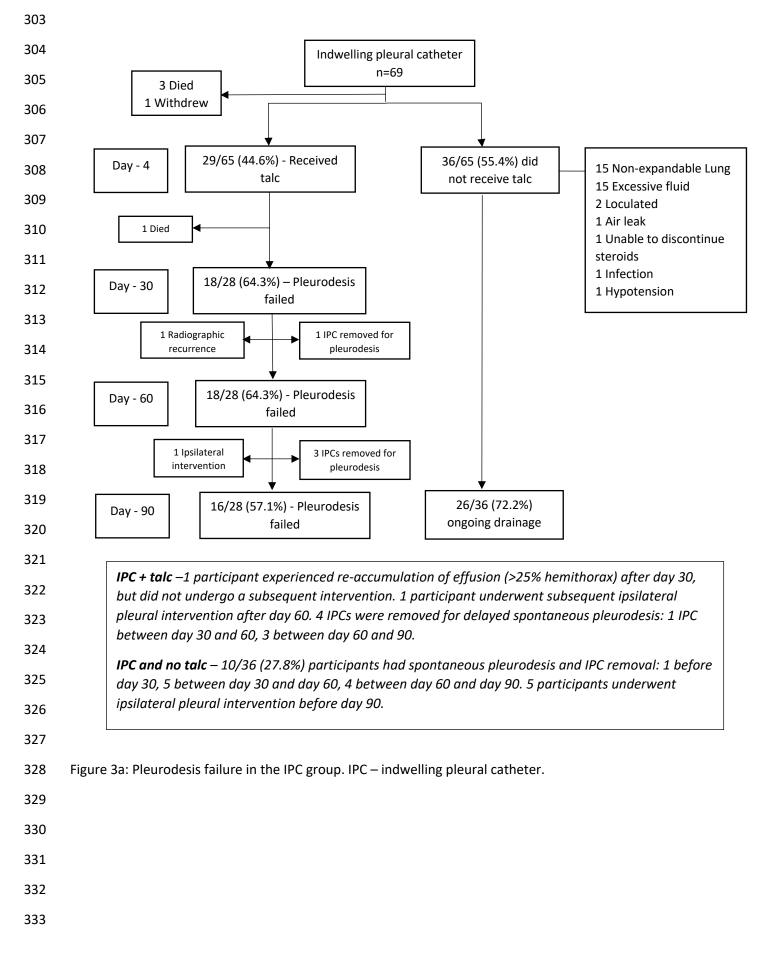
296 VAS – visual analogue scale.

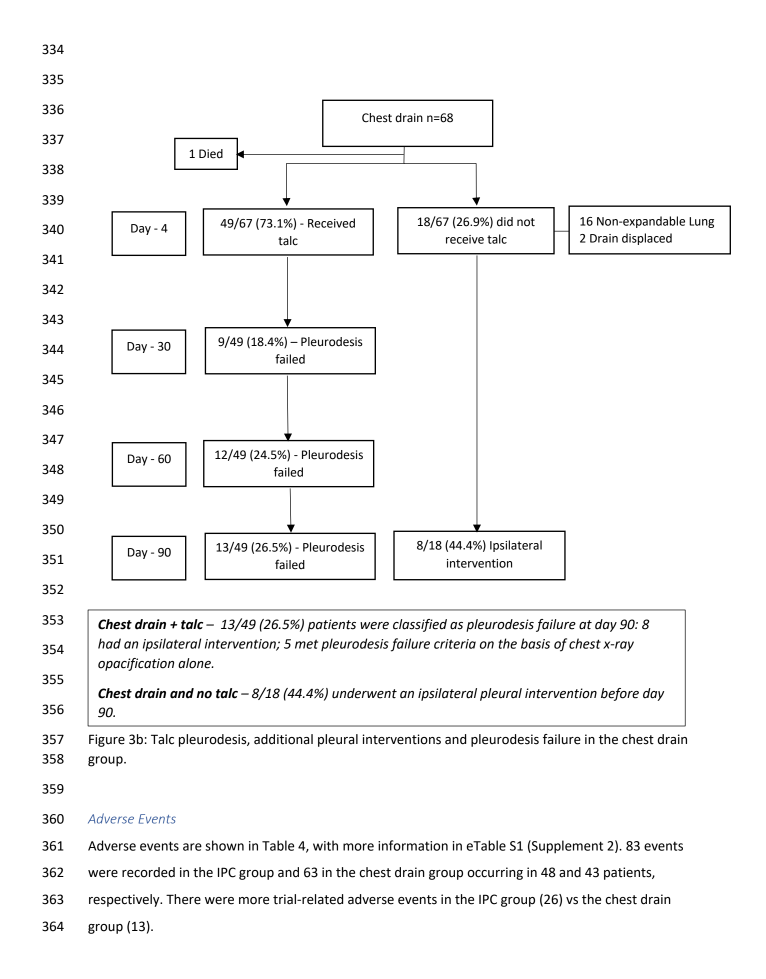
a: Unless otherwise stated.

b: ANCOVA adjusted for baseline global health status as a covariate.

c: Adjusted regression model for the stratification factors: (age [≤65years, >65years], WHO performance status [0,1,2 or 3],
 underlying malignancy [mesothelioma, breast cancer, lung cancer or other]).

301





- 365 Median per protocol hospitalization for chest drain patients was 4 days. There were 26 additional
- 366 hospitalizations in the chest drain arm; none of these hospitalizations was related to the trial
- 367 intervention. There were an additional 40 hospitalizations in the IPC arm; 7 of 40 were deemed
- 368 intervention related. Two patients in the IPC arm were admitted primarily due to anxiety related to
- their indwelling catheter.
- Thirty-six patients died (20 in the chest drain arm and 16 in the IPC arm). One patient died in the IPC
- arm due to pleural infection on a background of advanced malignancy. The odds ratio for death was
- 372 0.71 (95% Cl, 0.33 to 1.52, p=0.39).
- 373 Table 4: Reported adverse events

	No. of events		
Adverse Event	Chest Drain	IPC	
	(n=63)	(n=83)	
Intervention related serious adverse event	0	7	
Hospital admission for drain related anxiety	-	2	
Hospital admission for drain related pain	-	1	
Hospital admission with pleural infection	-	1	
Hospital admission with pleurodesis related	-	1	
pain			
Pre-pleurodesis steroid withdrawal	-	1	
Post insertion oxygen requirement	-	1	
Intervention related adverse event	13	19	
Pleurodesis related pain	1	1	
Drain related pain	-	4	
Hydropneumothorax with air leak	1	2	
Pleurodesis related fever	1	-	
Cutaneous infection	2	5	
Pleural infection	1	1	
Tube displacement	1	2	
Drain blockage	5	2	
Tract metastasis	-	1	
Vasovagal syncope during insertion	1	-	
Failed drain insertion	-	1	
Events not related to intervention	50	57	
Death	20	16	
Admission for symptom control / cancer	9	10	
progression			
(non-intervention related)			
Other (see eTable S1)	21	31	

#### **376** *Post-hoc analysis*

Data were examined on physical, social, emotional, cognitive and role functional domains. Theseresults are summarised in Supplement 2 (Figure S4 and eTable S4).

379

#### 380 Discussion

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To the best of our knowledge, this is the first multicentre, open-label randomised controlled trial to investigate the impact of management of MPE on HRQoL as a primary outcome.

Within the limits of the study, our findings suggest that the outpatient IPC pathway was not superior

to inpatient treatment with a chest drain and talc slurry in improving global health status, as

measured by the EORTC QLQ C-30 questionnaire at 30 days post-intervention. However, both

treatment approaches delivered sustained improvements in global health status at 60 and 90-days.

388 These findings align closely with HRQoL outcomes observed in the TIME2 trial, where both

approaches yielded similar EORTC QLQ C-30 outcomes in a smaller patient cohort[12]. Additionally,

another interventional study assessing HRQoL as a secondary measure, did not identify any

391 significant differences between the outpatient and inpatient treatment pathways over 1 year of

follow up when utilising the EQ 5D-5L and the 100mm VAS tools[4].

393 IPCs are increasingly being used as a first line treatment in the management of symptomatic, recurrent 394 malignant pleural effusion in some parts of the world, despite a lack of trial data to support its 395 superiority over chest drain and talc pleurodesis in symptom control and hospital length of stay in the 396 year after intervention[4]. A key goal of treatment is to improve health-related quality of life, and 397 given our findings, we advocate that patients select a treatment pathway best suited for their overall 398 health, psychological and social circumstances. However, the outpatient pathway is a multifaceted 399 intervention, requiring a secure framework of community healthcare support. In regions where this 400 may not be viable, we recommend opting for a chest drain and talc pleurodesis.

401 Our study also examined pleurodesis failure as a secondary outcome. This was significantly higher in 402 the IPC arm than in the chest drain arm (69% vs 26.5% at 90 days, respectively). Several reasons may 403 explain this finding. The definition of talc pleurodesis failure in the outpatient arm included patients 404 in whom the IPC remained in situ despite successful pleurodesis; chosen because a key aim of offering 405 pleurodesis via an IPC is to promote expeditious catheter removal. Three of 13 (23%) eligible patients 406 chose not to have their IPCs removed following successful pleurodesis, highlighting the anxiety surrounding fluid recurrence. Patients in the IPC group had larger effusions, and more were receiving 407 408 systemic steroid therapy at the time of trial recruitment; which may have had an effect.[13, 14]

409 The IPC-PLUS trial, which primarily examined pleurodesis outcomes via an IPC, reported a 43%

410 pleurodesis success rate at 35 days post-talc insufflation,[2] which is slightly higher than the rate of

411 pleurodesis success at Day 30 (35%) in this study. However, differences in drainage protocols

412 (affecting the degree of pleural apposition) and patient selection may account for this disparity.

Of note, there were significantly more intervention related adverse events in the outpatient treatment arm, particularly hospitalization due to an intervention-related adverse event (7 vs 0). Given the lack of superiority of outpatient management for the primary outcome measure and similar outcomes, these findings may suggest that chest drain and talc pleurodesis may be the preferred treatment option in some patients.. Therefore, we advocate careful consideration and counselling of the potential risks before offering management with an IPC.

419 The main limitation is the data attrition for the primary outcome. Despite this, the findings still 420 contribute valuable insights into HRQoL outcomes. Significant improvements in global health status 421 were seen in both treatment arms with the observed difference well below clinically significant levels, 422 and these findings are also supported by the sensitivity analysis. This study reflects the considerable 423 practical challenges faced in conducting studies in this population, particularly when implementing 424 complex interventions in a real-world context. Strategies to mitigate attrition in future studies in this 425 area may include utility of a scoring tool for prognostication or delivery of the study follow-up within 426 the patient's home setting.

427 The EORTC QLQ-C30 questionnaire used to assess overall HRQoL was selected due to its validation in 428 the cancer population. However, the minimal important difference (MID) for the EORTC QLQ-C30 429 global health status scale varies in different cancer populations. The MID chosen for the primary 430 outcome analysis was selected based on a combination of previously published data in the lung and 431 breast cancer population. More importantly, this has not been specifically defined in the MPE 432 population and to the best of our knowledge, there is no HRQoL questionnaire that is specifically 433 designed and validated for patients with MPE. A bespoke questionnaire focused on a combination of 434 physical, emotional and social functions in patients within the last year of life may be useful in 435 understanding outcomes in these other domains. Only patients with a performance status of less than 436 3 and an expected prognosis of 3 months were included, as assessed by the treating clinician. This was 437 to ensure patients committed to the outpatient treatment pathway would be ambulant and well 438 enough to undertake the requirements of the study. Therefore these results may not be applicable to 439 a key group of patients with poorer performance status and shorter life expectancy.

The outpatient pathway is resource intensive, especially concerning community nursing assistance,
 expenses linked to drainage bottles, and the hospital resources required for administering outpatient

talc treatment. Despite the promotion of ambulatory management approaches by policymakers, thesedemands might not be achievable in every setting.

To summarise, this study suggests that treatment using an IPC and managing as outpatient is not superior to inpatient chest drain and pleurodesis at improving HRQoL. However, the IPC pathway is associated with an increased rate of adverse events. An informed treatment choice should be made based on patient preferences, acceptability of risk, social circumstances, values, affordability and treatment accessibility.

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### 451 Contributors

- 452 Sivakumar, Douiri and Ahmed had full access to all of the data in the study and take responsibility for
- 453 the integrity of the data and the accuracy of the data analysis.
- 454 Concept and design: Ahmed, Sivakumar, Rao, Douiri
- 455 Acquisition, analysis, or interpretation of data: All authors
- 456 Drafting of the manuscript: Sivakumar, Ahmed
- 457 Critical revision of the manuscript: All authors
- 458 Statistical analysis: Douiri, Sun
- 459 Obtained funding: Ahmed
- 460 Administrative, technical, or material support: Wallace, Noorzad, West, Simpson, Sivakumar, Ahmed
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### 462 Declaration of interests

- 463 Dr. Ahmed reported receiving grants from GE outside the submitted work.
- 464 Dr. Sivakumar reported receiving grants from Rocket Medical outside the submitted work.
- 465 Dr. Fitzgerald reports grants from European Respiratory Society outside the submitted work.
- 466 Professor. Maskell reports grants and personal fees from Beckton Dickinson and Company outside of467 the submitted work.
- 468 Professor. Lee reports non-financial support from Rocket Med Plc, grants from Rocket Med PLC,
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- 470 No other disclosures were reported.
- 471

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## 477 *Role of the Funder/Sponsor*

- 478 Neither the funder nor the sponsor had a role in the design and conduct of the study; collection,
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