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1 An investigation of the effects of procalcitonin testing on antimicrobial prescribing in respiratory

2 tract infections in an Irish University Hospital setting - a feasibility study.

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12 Synopsis

13 Diagnostic uncertainty and a high prevalence of viral infections present unique challenges for

- 14 antimicrobial prescribing for respiratory tract infections (RTIs). Procalcitonin (PCT) has been
- 15 shown to support prescribing decisions and reduce antimicrobial use safely in patients with RTIs but
- 16 recent study results have been variable.
- 17 Methods

We conducted a feasibility study of the introduction of PCT testing in patients admitted to hospital with a lower RTI to determine if PCT testing is an effective and worthwhile intervention to introduce to support the existing AMS programme and safely decrease antimicrobial prescribing in patients admitted with RTIs.

- 22 Results
- 23 A total of 79 patients were randomised to the intervention PCT guided treatment group and 40
- 24 patients to the standard care respiratory control group.
- 25 The addition of PCT testing led to a significant decrease in duration of antimicrobial
- 26 prescriptions (mean 6.8 versus 8.9 days p=0.012) and decreased length of hospital stay (median
- 27 7 versus 8 days, p=0.009) between the PCT and respiratory control group. PCT did not demonstrate
- a significant reduction in antimicrobial consumption when measured as DDDs and days of
- 29 therapy.
- 30 Conclusions

PCT testing had a positive effect on antimicrobial prescribing during this feasibility study. The successful implementation of PCT testing in a randomised controlled trial requires an ongoing comprehensive education programme, greater integration into the AMS programme and delivery of PCT results in a timely manner. This feasibility study has shown that a larger randomised controlled trial would be beneficial to further explore the positive aspects of these findings.

37 Introduction

Antimicrobial resistance (AMR) is a major risk to public health globally that leads to increasing healthcare costs, treatment failure, and increased morbidity and mortality.¹⁻³ There is a strong association between sub-optimal antimicrobial prescribing and AMR.⁴ To optimise prescribing, hospital antimicrobial stewardship (AMS) programmes should target areas of high antimicrobial prescribing. One such area is respiratory tract infections (RTIs). Shorter antimicrobial courses offer one potential solution to the overuse of antimicrobials for RTIs⁵ and there is evidence to support such strategies⁶⁷ even in severe hospital infections.⁸

Diagnostic uncertainty and a high prevalence of viral infections present unique challenges for 45 antimicrobial prescribing for RTIs.⁹⁻¹² This contributes to over-use and/or sub-optimal use of 46 antimicrobials¹³ ¹⁴ for RTIs such as community acquired pneumonia (CAP), including prolonged 47 48 treatment courses of up to 11 days,¹⁵ without a correlation between duration of treatment and infection severity.^{15 16} Physicians are often reluctant to shorten antimicrobial course durations due to 49 50 the fear of incomplete pathogen eradication which could potentially lead to relapse and associated morbidity and mortality.⁶ There is also a high rate of antimicrobial continuation where viral 51 infections,¹⁷ including influenza,¹⁸ are identified due to overriding concerns about secondary bacterial 52 53 infections. However, a recent study has shown a bacterial co-infection rate of only 40%.¹¹

54 To address these issues, there is a growing interest in the use of novel diagnostic techniques and biomarkers as an AMS tool.¹⁹ It is important that AMS programmes investigate the opportunity 55 afforded by these new techniques and the potential they offer to optimise antimicrobial treatment 56 more promptly²⁰ and change prescribing behaviour.²¹ Procalcitonin (PCT) testing is one such diagnostic 57 58 technique. PCT is a peptide precursor to the hormone calcitonin. It is usually undetected but is upregulated in response to a bacterial infection following stimulation of bacterial-induced cytokines.²² 59 60 Upregulation of PCT is blocked in viral infections due to the release of the cytokine interferon gamma, resulting in a higher specificity of PCT to distinguish between bacterial and viral infections when 61 compared to other inflammatory markers such as CRP.²³ PCT levels decrease rapidly when patients 62

are recovering from infection.²⁴ Hence it offers the potential to support clinical decision making for the initiation and discontinuation of antimicrobials in patients with a clinical suspicion of a bacterial infection when considered along with the clinical assessment of the patients. PCT has been shown to support prescribing decisions and reduce antimicrobial use safely in patients with RTIs²⁵⁻²⁸ but findings from recent studies have been variable,^{29 30} so it is unclear if it is an effective intervention as part of an AMS programme.

The purpose of this study was to conduct a feasibility study to determine if PCT testing is an effective
and worthwhile intervention to introduce in a University Teaching Hospital to support the existing
AMS programme and safely decrease antimicrobial prescribing in patients admitted with RTIs.

72

73 Methods

74 We conducted a single centre, randomised, open-label feasibility study of the introduction of PCT 75 testing in patients admitted to hospital with a lower RTI under the care of the respiratory medicine 76 team during on-call acute unselected general medical take to determine if PCT testing had an impact 77 on antimicrobial consumption and patient's length of stay (LOS) in hospital. The study was conducted 78 in a single 321 bed inner city, voluntary acute University Teaching Hospital, which is part of the South/South West Hospital Group³¹ in the Republic of Ireland. It is a Model 3 (smaller general) ³² 79 80 hospital with a 24-hour emergency department, ICU and admits undifferentiated acute medical and 81 surgical patients. The hospital has an established AMS programme and no significant changes were 82 made to the AMS policies or programme during this study.

83 Ethics

The study was approved by the local ethics committee (approval code ECM 4 (w) and ECM 3 (III)).

85 Written informed consent was obtained from all participants prior to study enrolment.

86 Education and training

The microbiology laboratory scientists received technical advice and training on the operation of the
PCT assay from the manufacturer prior to study commencement. They also received a presentation
on the introduction of PCT testing in the hospital.

90 The respiratory medicine team received presentations at the respiratory journal club meetings and 91 provision of written materials electronically. Presentations consisted of evidence supporting PCT use 92 in practice, limitations of PCT testing, PCT measurement, and interpretation using a PCT-based 93 antimicrobial prescribing algorithm (Supplementary material S1). Presentations were given prior to 94 the study commencement and following medical staff rotation changes. The study protocol 95 (Supplementary material S2), study flow chart and the PCT-based antimicrobial prescribing algorithm 96 was provided to all physicians electronically.

97 Recruitment and consent

98 Inclusion criteria

99 Adult patients ≥18 years of age, admitted to hospital under the care of the respiratory teams with an 100 initial diagnosis of an acute lower RTI (i.e. Community acquired pneumonia ³³ with severity defined by 101 CURB-65 score³⁴, Lower RTIs³⁵, exacerbation of asthma³⁶, COPD³⁷, bronchiectasis³⁸, interstitial lung 102 disease³⁹ and influenza³⁵) and commenced on antimicrobial therapy were identified from the daily 103 admission census or by the respiratory medicine teams.

The randomisation process stratified patients according to presence or absence of severe COPD GOLD Stage D criteria 2017³⁷ to ensure balanced treatment allocation. Patients were then randomly allocated in a 2:1 ratio to either the PCT guided treatment group or the standard care respiratory control group. Randomisation was carried out using sequentially numbered opaque sealed envelopes. A second general control group of patients admitted under general medicine teams with a diagnosed acute lower RTI and received standard care (no PCT measurement) was recruited to provide a comparison of antimicrobial prescribing in RTIs by non-respiratory specialist physicians in the hospital.

111 Exclusion Criteria

Exclusion criteria were: unable to give written informed consent due to language restrictions, 112 cognitive impairment or severe dementia; re-admission to hospital within 30 days of previous 113 114 admission; immunosuppression (neutropenic, chemotherapy, radiation therapy or 115 immunosuppressive therapy) other than corticosteroid use; life-threatening medical co-morbidities 116 leading to possible imminent death, Do Not Resuscitate (DNR) status; patients with concurrent chronic 117 infections necessitating prolonged antimicrobial treatment (cystic fibrosis, tuberculosis, infective 118 endocarditis, osteo-articular infections, hepatic or cerebral abscesses, chronic prostatitis); patients 119 with >24 hours of appropriate antimicrobial therapy prior to initial PCT level; active intravenous drug 120 users; pregnant women.

121 Intervention

PCT testing was commenced in the microbiology department following completion of staff training
 and instrument validation. It was available during routine working hours (Monday to Friday, 9am 5pm). PCT serum concentrations were measured using the VIDAS BRAHMS PCT (assay range 0.05-200
 µg/L) (Biomerieux, France).

126 PCT serum concentrations were interpreted using an evidence based algorithm (Supplementary material S1)⁴⁰ which has been validated in previous studies^{28 29} recommending antimicrobials strongly 127 128 discouraged for PCT levels < 0.1 μ g per litre, discouraged for levels 0.1 to 0.25 μ g/L, encouraged for 129 levels > 0.25 to 0.5 μ g/L and strongly encouraged for levels > 0.5 μ g/L. The algorithm also included specific overruling criteria where antimicrobials could be considered in the case of respiratory or 130 131 haemodynamic instability; life-threatening co-morbidity; need for ICU admission; PCT < 0.1 μ g /mL: CAP with CURB 65 > 3, COPD stage IV; PCT < 0.25 μ g/mL: CAP with CURB 65 > 2; localised infection 132 (abscess, empyema); immunocompromised (other than corticosteroids); concomitant infection in 133 134 need of antimicrobials.

135 The antimicrobial prescribing advice generated from the PCT algorithm was verbally communicated 136 to the respiratory medicine team and this advice was non-binding. The respiratory medicine team 137 retained prescribing autonomy regarding clinical decisions irrespective of the PCT level or algorithm 138 generated antimicrobial prescribing advice. The algorithm adherence for antimicrobial prescribing 139 recommendations was recorded at 24 hours following the PCT test for all patients along with the 140 rational for prescribing decisions. Algorithm adherence was defined as antimicrobial therapy that was continued or discontinued in accordance with the PCT cut-off ranges. Non-adherence was defined as 141 142 antimicrobial therapy that was not discontinued despite low PCT levels. Over-riding criteria were not 143 considered when measuring adherence but were recorded as reasons for non-adherence.

Patients were followed until their discharge. A further follow up of medical records took place at 30
days post admission to identify re-admitted patients and re-admitted patients with infection re-lapse.
Patient recruitment ran from June 1st 2017 to May 31st 2018. Figure 1 represents the patient hospital
journey with a respiratory tract infection.

148 Outcomes

149 The primary outcomes were to quantify the individual inpatient antimicrobial consumption, prescription duration and the inpatient LOS. Following a recent systematic review which 150 151 recommended that antimicrobial use should be expressed in at least two metrics simultaneously, ⁴¹ 152 antimicrobial consumption was measured using DDDs, days of therapy (DOTs) and prescription 153 duration. DDDs were calculated using the Anatomical Therapeutic Chemical/Defined Daily Dose (ATC/DDD) index of the WHO Collaborating Centre for Drug Statistics Methodology ⁴² but were not 154 155 adjusted for hospital activity. Days Of Therapy (DOT)⁴³ calculates individual patient-days of antimicrobial exposure and accounts for dosing and frequency of each drug. Antimicrobial prescription 156 duration was measured in days (defined as the number of days between the commencement and 157 158 discontinuation of antimicrobials). The LOS was defined as date of discharge less date of admission.

Secondary outcomes were number of infection and antimicrobial related adverse events during inpatient LOS including mortality, hospital re-admission, and infection re-lapse requiring re-admission both within 30 days. Algorithm adherence for antimicrobial prescribing recommendations was measured.

A qualitative process evaluation of the study was conducted in parallel with this feasibility study andwill be reported in a subsequent paper.

165 Statistical methods

166 A Microsoft Access database (version 1903) was developed to record the study data. Statistical

analysis was conducted using R (version 3.4.0) and was conducted on an intention to treat basis.

168 The primary outcome of antimicrobial consumption between the PCT and respiratory control arms

169 was evaluated using the non-parametric Wilcoxon Rank Sum test. A Kaplan-Meier curve was used to

analyse the median time to discharge between the PCT and respiratory control group.

171 Chi-square tests were used to evaluate differences between the PCT and respiratory control arms for

all secondary outcomes - number of adverse events, re-admission and infection re-lapse requiring re-

admission both within 30 days.

175 Results

The respiratory medical teams admitted 823 general medical patients of whom 313 patients were
 classified as a respiratory infection or respiratory disorder during the recruitment period of June 1st
 2017 to May 31st 2018. A CONSORT flow diagram of recruitment can be seen in figure 2.

A further 48 patients were recruited to the general control group. Three patients who were identified
as suitable to enter the general control group were not recruited due to confusion, isolation due to
infection and refused consent.

Demographic data and study overview are contained in Table 1. Clinical findings of patients onadmission to hospital are contained in Table 2.

There were several differences between the baseline characteristics of the PCT group and respiratory control group. The PCT group contained more male patients (60% versus 42%), active smokers (25% versus 12.5%) and patients with pre-existing COPD A-C (29% versus 17%).

There were a number of differences in final diagnosis between the PCT group and the respiratory control group with asthma (3.8% versus 15%), CAP (10% versus 7.5%), LRTI (30.4% versus 17.5%. CAP severity in the PCT group had CURB-65 scores ranging from 0 to 3 with a mean of 1.87 while the CAP severity in the respiratory control group had CURB-65 scores ranging from 0 to 1 with a mean of 0.66. The clinical findings on admission were similar between group with two exceptions where the PCT group had a higher percentage of patients who were productive of sputum on admission (49% versus

193 37%) and patients prescribed antibiotics prior to admission (35% versus 25%).

194 **Procalcitonin testing and results**

The 79 patients randomised to the PCT group had a total of 163 PCT levels taken (median of 2 tests per patient (range 1-6). Overall the PCT levels had a median of $0.075\mu g/L$ (IQR 0.05 - 0.26). The initial PCT level was $\leq 0.24 \mu g/L$ for 58 patients (including 38 patients with an initial PCT level of $\leq 0.05 \mu g/L$). Our primary outcome was to determine the inpatient antimicrobial consumption, duration of antimicrobial treatment and hospital LOS. The main outcomes can be seen in Table 3 and Figure 3. Statistical analysis was conducted on the PCT and respiratory control groups and does not include comparison with the general control group.

202 There was no significant difference in antimicrobial exposure or usage per patient when measured

203 as DDD (11.1 \pm 7.5 versus 13.1 \pm 10.7, p=0.218) (mean \pm SD) or DOT (8.9 \pm 6.3 versus 11 \pm 7.6,

p=0.077) of patients between the PCT and respiratory control group. Median values of both metrics

205 DDD (8.66 versus 9.57) and DOT (7.5 versus 8.25) showed a decrease of 9% in antimicrobial

206 consumption per patient.

207 There was a significant difference in the antimicrobial duration in days between the PCT and

respiratory control groups (median 7 versus 8 days, p=0.0125). There was also a significant

209 difference between the PCT and respiratory control groups in the median LOS (p=0.009) and this can

also be seen in the Kaplan–Meier curves in Figure 4.

In the analysis of secondary outcomes there was no significant differences between the PCT and respiratory control groups in the incidence of adverse events during in-patient hospital stay (p=0.9852), the rate of hospital re-admission (p=0.1507), and the rate of infection re-lapse requiring re-admission both within 30 days (p=0.0924).

Algorithm compliance is displayed in table 4.

216 Overall PCT algorithm compliance per patient was 35% within 24 hours of PCT level being taken. 25 217 patients had high PCT levels ($\geq 0.25 \ \mu g/L$) where the algorithm recommendation was to continue antimicrobial treatment and algorithm compliance was 100%. 67 patients had low PCT levels (< 0.25 μ g/L) where the algorithm recommendation was to discontinue antimicrobial treatment and algorithm compliance was low (10%). In these instances, the reasons for non-adherence were based on a clinical decision in 55/112 (49%) PCT levels with the remaining 57/112 (51%) PCT levels based on meeting various algorithm overriding criteria (respiratory or haemodynamic instability; life threatening co-morbidity; need for ICU admission; localised infection (abscess, empyema)).

Seven patients had their antimicrobial treatment discontinued in compliance with the algorithm when PCT levels were low (< 0.25 μ g/L). This resulted in shorter course lengths in five patients (< 7 days) 1 course length completion as planned at 7 days, and early antimicrobial discontinuation (day 2) in a patient with influenzae. There were no hospital readmissions among these patients.

In a further 9 patients where there was initial non-compliance with the algorithm recommendations when measured at 24 hours, their antimicrobial treatment was subsequently modified resulting in a shorter course length in 7 patients (< 7 days) and 2 further patients discontinued antimicrobials prior to discharge (1 patient re-admitted with infection).

Algorithm compliance by indication was as follows; CAP (80%), asthma (50%) LRTI (30%), COPD (12.5%) and influenza virus (42%). PCT levels and algorithm compliance were found to be low in patients with COPD stage D and structural lung conditions like bronchiectasis and interstitial lung disease. In these cases, the clinical judgement of physicians was to over-ride the algorithm recommendations and continue antimicrobials.

237 Microbiology positive specimens

38 patients (23%) had positive microbiology results : 13 influenzae virus, 10 bacterial isolates from
 respiratory specimens and 7 yeast isolates from respiratory specimens.

240 Adverse events

Infection and antimicrobial related adverse events included gastro-intestinal (antimicrobial related diarrhoea 1 patient) renal function (acute kidney injury secondary to antimicrobials 1 patient), liver function (increased liver function tests secondary to antimicrobials 1 patient), respiratory disorders (hospital acquired pneumonia, hospital acquired influenzae, respiratory deterioration, 3 patients) and other events 2 patients.

246 Mortality during the study

247 Five patients included in the study died during their hospital stay, four from the PCT group and one

from the respiratory control group (age range 75-94 years). All had multiple co-morbidities including

249 cardiac (congestive cardiac failure, atrial fibrillation), renal and new or existing cancer diagnosis.

250 Antimicrobial treatment decisions for these patients were based on clinical decisions.

251 Discussion

This feasibility study of the introduction of PCT testing has shown a positive effect on antimicrobial 252 prescribing resulting in a decrease in the duration of antimicrobial courses in patients with RTIs and a 253 254 decrease in LOS without an increase in adverse events or re-admission to hospital. The median 255 duration of antimicrobial treatment was reduced from 8 to 7 days and antimicrobial consumption fell by 9% when measured as DDD and DOT. This study confirms the findings of previous PCT trials²⁸⁴⁴ that 256 257 it is an effective and worthwhile intervention to safely reduce antimicrobial exposure in patients with 258 RTIs and supporting the AMS programme. However, there were several findings which may have 259 influenced the outcomes and these need to be considered when viewing the overall results and 260 considering progression to and design of a full randomised controlled trial (RCT).

261 Overall PCT algorithm compliance was 35%, and compliance with stopping recommendations was 10% 262 when PCT levels were low (<0.25 μ g/L). The reasons for non-compliance were clinical judgement (49%) 263 and meeting pre-determined over-riding criteria (51%). PCT was a new diagnostic test in the hospital 264 and physicians can require time to become familiar with and develop confidence in the use of PCT 265 testing.⁴⁵ Other studies have found algorithm compliance can be variable ranging from 35% to 80%.⁴⁴ 266 An international, multicentre study found that centres with experience of using PCT and ongoing reinforcement of PCT guided AMS had higher algorithm compliance than PCT naive centres.⁴⁴ Protocol 267 driven studies^{28 46} have also shown higher algorithm compliance and greater impact on antimicrobial 268 prescriptions than studies taking a quality improvement implementation approach.²⁹ 269

Algorithm compliance must improve significantly in a future trial to maximise the potential impact of PCT testing on antimicrobial prescribing decisions but also acknowledging the limitations of PCT and that physicians cannot rely on PCT alone to guide antibiotic therapy.²³ In a future trial this should be addressed by a more comprehensive educational programme and more effective incorporation into the AMS programme to re-enforce PCT recommendations. Such an approach has been shown to be effective^{30 46 47} and required for interventions such as PCT to realise their full benefit.¹⁹ The educational element of this study may not have been sufficient. A future trial should consider the inclusion of more
 frequent educational presentations prior to and during the intervention and include case reviews of
 PCT patients. Consideration should be given to the development of pocket cards, incorporation into
 local electronic antimicrobial prescribing guidelines and availability of results on the hospital
 electronic laboratory system.⁴⁶

281 Delays in availability of PCT results may have also decreased the impact of the intervention and 282 contributed to poor algorithm compliance with 38% of PCT serum results not available until the next 283 day (24 hours after the serum sample was taken). This included results which were delayed or 284 unavailable for 12 patients until after they were discharged. In a future trial prompt availability of PCT 285 levels is important. This would allow physicians to consider PCT along with routine biochemistry and 286 blood analysis, and the patients' clinical parameters at the point of care when making antimicrobial 287 prescribing decisions. Consideration should be given to measurement of algorithm adherence at 48 288 hours to account for unforeseen delays PCT result availability or delayed physician review of PCT results. 289

There were several factors involved in patient recruitment which may have influenced the primary outcomes of the study and should be addressed in a future trial design. These were the variation in infection severity between the PCT and respiratory control groups and the inclusion of patients who were already prescribed antimicrobials prior to hospital admission. These factors can be addressed in a suitably powered future RCT with the inclusion of illness severity scores, along with the use of multivariate and sub-group analysis.

A future RCT would include a broader range of physicians rather than respiratory specialists alone. Antimicrobial consumption in the general control group of patients in this study was higher than in either of the respiratory groups. The addition of a PCT testing to the existing AMS programme may have the potential to have a greater impact on this patient group.

301 Strengths and limitations

The study was conducted in a setting where PCT was a newly available test to physicians. A broad range of RTIs were recruited and the study took place over a calendar year and included seasonal variation in illness and prescribing. Patients were randomised to intervention or control, thus reducing selection bias. Serial PCT measurements were available to guide antimicrobial prescribing.

306 The study had some limitations. The study population had a clinical need for antimicrobial treatment 307 so the study was designed to examine the duration of therapy and LOS, rather than investigating the 308 potential to withhold antimicrobial therapy. The study results may have been influence by a study 309 effect. Both the PCT and respiratory control groups were treated by the same group of physicians who 310 all received education and as they were aware that their behaviour was being monitored which may have resulted in a Hawthorn effect.⁴⁸ The intervention was limited to one medical speciality which 311 312 may limit its generalisability to other medical specialties and settings. The need for consent and PCT results which were not available at the point of clinical decision making in a small number of cases. 313

314

315 Conclusion

PCT testing had a positive effect on antimicrobial prescribing during this feasibility study. Several factors were identified which may have influenced the outcomes and the intervention implementation. The successful implementation of PCT testing requires an ongoing comprehensive education programme, greater integration into the AMS programme and delivery of PCT results in a timely manner. This feasibility study has shown that a larger randomised controlled trial would be beneficial to further explore the positive aspects of these findings.

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- 332
- 333 Transparency declarations
- 334 Nothing to declare.

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482 Figure 1 Patient hospital journey with a respiratory tract infection





528 Table 1. Demographic data and study overview

Variable		Study Group				
		Overall	PCT arm	Respiratory Control arm	General Control arm	
Participants		167 (100%)	79 (47.3%)	40 (24.0%)	48 (28.7%)	
Gender	Female	79 (47.3%)	31 (39.2%)	23 (57.5%)	25 (52.1%)	
	Male	88 (52.7%)	48 (60.8%)	17 (42.5%)	23 (47.9%)	
Age (years)		68.7 ± 14	68.6 ± 13.6	68.4 ± 15.3	69.1 ± 13.9	
Co-existing conditions and risk factors						
Smoking Status	Non- smoker	50 (30.3%)	26 (32.9%)	13 (32.5%)	11 (23.9%)	
	Smoker	33 (20%)	20 (25.3%)	5 (12.5%)	8 (17.4%)	
	Ex-smoker	82 (49.7%)	33 (41.8%)	22 (55%)	27 (58.7%)	
Asthma		28 (16.8%)	13 (16.5%)	10 (25%)	5 (10.4%)	
COPD A-C		58 (34.7%)	23 (29.1%)	10 (25%)	25 (52%)	
COPD D		24 (14.4%)	10 (12.7%)	5 (12.5%)	9 (18.8%)	
Bronchiectasis		16 (9.6%)	9 (11.4%)	3 (7.5%)	4 (8.3%)	
Interstitial lung disease		7 (4.2%)	4 (5%)	2 (5%)	1 (2.1%)	
Final diagnosis						
Asthma		11 (6.6%)	3 (3.8%)	6 (15%)	2 (4.2%)	
Community acquired pneumonia		18 (10.8%)	8 (10%)	3 (7.5%)	7 (14.6%)	
COPD		62 (37.1%)	24 (30.4%)	13 (32.5%)	25 (52%)	
Lower respiratory tract infection		45 (27%)	28 (35.4%)	7 (17.5%)	10 (20.8%)	
Other lower respiratory tract infections		20 (12%)	10 (12.6%)	7 (17.5%)	3 (6.2%)	
Non-respiratory related		11 (6.6%)	6 (7.6%)	4 (10%)	1 (2.1%)	

531 Table 2. Clinical findings on admission to hospital*

	Total	РСТ	Respiratory	General control		
	(n = 167)	(n = 79)	Control	(n = 48)		
			(n = 40)			
Respiratory rate- breaths/minute	22.1 ± 5	22.1 ± 5.4	21.1 ± 3.7	22.7 ± 5.2		
Systolic blood pressure- mmHg	133 ± 23.1	130.9 ± 22.9	136 ± 20.9	134 ± 25		
Diastolic blood pressure- mmHg	75 ± 14.1	74.8 ± 12	78.6 ± 14.8	72.3 ± 16.1		
Heart rate- beats/minute	91.8 ± 20.1	93.4 ± 23.3	91.2 ± 16.7	89.8 ± 16.7		
Temperature- °C	36.8 ± 0.8	36.8 ± 0.8	36.9 ± 0.8	36.8 ± 0.9		
Rigors - no. (%)	24 (14.4%)	11 (13.9%)	6 (15%)	7 (14.6%)		
Fever - no. (%)	18 (10.8%)	8 (10.1%)	5 (12.5%)	5 (10.4%)		
Chills - no. (%)	15 (9%)	10 (12.7%)	1 (2.5%)	4 (8.3%)		
Number of clinical signs of infection	1.8 ± 1.3	1.9 ± 1.3	1.7 ± 1.2	1.8 ± 1.3		
Documented signs of respiratory illness						
Cough - no. (%)	132 (79%)	64 (81%)	31 (77.5%)	37 (77%)		
Shortness of breath - no. (%)	101 (60.5%)	45 (57%)	23 (57.5%)	33 (68.7%)		
Productive of sputum - no. (%)	81 (48.5%)	39 (49.4%)	15 (37.5%)	27 (56.2%)		
Dyspnoea - no. (%)	49 (29.3%)	22 (27.8%)	10 (25%)	17 (35.4%)		
Pleuritic pain - no. (%)	26 (15.6%)	10 (12.7%)	9 (22.5%)	7 (14.6%)		
Respiratory failure - no. (%)	19 (11.4%)	8 (10.1%)	5 (12.5%)	6 (12.5%)		
Abnormal chest exam - no. (%)	144 (86.2%)	70 (88.6%)	31 (77.5%)	43 (89.6%)		
Abnormal radiological findings - no. (%)	94 (56.3%)	42 (53.2%)	21 (52.5%)	28 (58.3%)		
CURB-65 score (CAP patients)	1.56 ± 1.05	1.87 ± 1.05	0.66 ± 0.47	1.57 ± 1.05		
Number of signs of acute respiratory	3.9 ± 1.4	3.8 ± 1.4	3.8 ± 1.4	4 ± 1.3		
illness						
Antimicrobials prescribed pre-	59 (35.3%)	28 (35.4%)	10 (25%)	21 (43.7%)		
admission - no. (%)						
Corticosteroids prescribed pre-	34 (20.4%)	14 (17.7%)	7 (17.5%)	13 (27%)		
admission - no. (%)						
Community - no. (%)	149 (89.2%)	70 (88.6%)	32 (80%)	47 (98%)		
Healthcare - no. (%)	13 (7.8%)	6 (7.6%)	6 (15%)	1 (2%)		
Hospital - no. (%)	5 (3%)	3 (3.8%)	2 (5%)	0 (0%)		

532 *plus minus values are means + SD

533 Table 3. Primary and secondary outcome data

Primary outcomes	РСТ	Respiratory Control	General control	n value		
Mean + SD	(n = 79)	(n = 40)	(n = 48)	p value		
(Median)	(11 - 75)	(11 – 40)	(11 – 40)			
		42.4 + 40.7	40 5 + 44	0.210		
Defined daily doses	11.1 ± 7.5	13.1 ± 10.7	18.5 ± 11	0.218		
per patient	(8.66)	(9.57)	(16.5)			
Days of therapy per	8.9 ± 6.3	11 ± 7.6	13.7 ± 11.1	0.077		
patient	(7.5)	(8.25)	(11.63)			
Total duration of	6.8 ± 2.8	8.9 ± 4	8.4 ± 3.6	0.0125*		
inpatient	(7)	(8)	(8)			
antimicrobials						
(days)						
Length of hospital	7.4 ± 4.3	10.5 ± 6.1	8.9 ± 3.8	0.009*		
stay (days)	(7)	(8)	(8)			
Secondary outcomes						
Hospital	7 (8.9%)	8 (20%)	7 (14.6%)	0.1507		
readmission within						
30 days						
Relapse of infection	6 (7.6%)	8 (20%)	6 (12.5%)	0.0924		
within 30 days						
Adverse events	6 (7.6%)	3 (7.5%)	4 (8.3%)	0.9852		

 *= statistical significance was set as <0.05, p-values relate to the comparison between the PCT and respiratory control groups



Figure 3. Main antimicrobial consumption outcomes.





543 Kaplan–Meier curves. Median probability of discharge is given by the horizontal dashed line.

Table 4. PCT algorithm compliance

PCT level	Algorithm	Number	Number of	Number of	Number of	Percentage
(µg/L)	recommendation	of	PCT test	patients	patients	of patients
		patients	results	compliant	non-	compliant
				with	compliant	with
				algorithm	with	algorithm
					algorithm	
≤0.05	Antimicrobial	67	119	7	60	10%
to <0.25	therapy					
	discouraged					
≥0.25	Antimicrobial	25	44	25	0	100%
	therapy					
	encouraged					