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1	Title: Malnutrition and poor food intake are associated with prolonged hospital
2	stay, frequent readmissions, and greater in-hospital mortality: Results from
3	the Nutrition Care Day Survey 2010
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14	
15	Short title: Malnutrition, poor food intake, poor outcomes. (50 characters)
16	
17	List of Abbreviations:
18	ANCDS- Australasian Nutrition Care Day Survey
19	ARDRG- Australian Refined Diagnosis Related Group
20	BMI- Body Mass Index
21	CI- Confidence Interval

- 22 DRG- Diagnosis Related Group
- 23 EQ-5Dvas- EQ-5D visual analogue scale
- 24 LOS- Length of stay
- 25 MDC- Major Diagnostic Category

26 MST- Malnutrition Screenin	g Tool
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- 27 PCCL- Patient Clinical Complexity Level
- 28 SGA- Subjective Global Assessment
- 29
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51 ABSTRACT

Background and Aims: The Australasian Nutrition Care Day Survey (ANCDS) 52 ascertained if malnutrition and poor food intake are independent risk factors for 53 health-related outcomes in Australian and New Zealand hospital patients. 54 Methods: Phase 1 recorded nutritional status (Subjective Global Assessment) and 55 24-hour food intake (0, 25, 50, 75, 100% intake). Outcomes data (Phase 2) were 56 collected 90-days post-Phase 1 and included length of hospital stay (LOS), 57 readmissions and in-hospital mortality. 58 **Results:** Of 3122 participants (47% females, 65 ± 18 years) from 56 hospitals, 32% 59 were malnourished and 23% consumed ≤ 25% of the offered food. Malnourished 60 patients had greater median LOS (15 days vs. 10 days, p<0.0001) and readmissions 61 rates (36% vs. 30%, p=0.001). Median LOS for patients consuming $\leq 25\%$ of the food 62 was higher than those consuming \geq 50% (13 vs. 11 days, p<0.0001). The odds of 90-63 day in-hospital mortality were twice greater for malnourished patients (CI: 1.09-3.34, 64 p = 0.023) and those consuming $\leq 25\%$ of the offered food (CI: 1.13-3.51, p = 0.017) 65 respectively. 66 **Conclusion:** The ANCDS establishes that malnutrition and poor food intake are 67 independently associated with in-hospital mortality in the Australian and New 68 Zealand acute care setting. 69 (196 words) 70 71 72 **Keywords:** malnutrition, poor food intake; disease type and severity; length of stay; 73 readmissions; in-hospital mortality 74 75

76 **INTRODUCTION**

77 The Australasian Nutrition Care Day Survey (ANCDS) is the largest multicentre study in the Australasian region, reporting the prevalence of malnutrition and poor food 78 intake in 3122 patients across 56 Australian and New Zealand hospitals [1]. With 79 one-in-three patients malnourished; and two-in-three patients not consuming all of 80 the offered hospital food, it was evident that malnutrition and poor food intake are a 81 common occurrence in Australian and New Zealand hospitals [1]. 82 Numerous studies have suggested that in comparison to well-nourished patients, 83 malnourished patients experience worse outcomes such as prolonged length of 84 85 hospital stay (LOS), increased readmissions, and mortality [2-6]. There is 86 documented evidence to suggest that malnourished patients incur greater hospitalisation costs [7], related to longer LOS, readmissions, and greater utilisation 87 of hospital resources [2, 5]. 88 The ANCDS found that one-in-three malnourished patients (n= 305, 30%), and one-89 in-five well-nourished patients (n= 371, 18%) consumed nothing or up to 25% of the 90 food offered during the 24-hour data collection period [1]. Since continued sub-91 optimal food intake can eventually lead to deterioration of nutritional status, it is 92 93 important to evaluate the effect of poor food intake on health-related outcomes. Two studies have reported the link between poor food intake during hospitalisation and 94 mortality [6, 8], however there is no published evidence regarding the association 95 96 between poor food intake and readmissions and/or LOS. 97

Although previous studies have investigated associations between malnutrition and
patient outcomes, issues such as heterogeneity in patient populations; study design;
methods of evaluating nutritional status, food intake and/or outcomes; prevent the

101	results from these studies being generalised throughout the acute care population.
102	Factors such as type and severity of disease are major causes of malnutrition [9],
103	poor food intake [10], and patient outcomes, and yet they have rarely been controlled
104	for. Without accounting for the confounding effect of disease type and severity most
105	studies fail to distinguish the association between the effect of disease, nutritional
106	issues, and other factors (such as age, gender), and patient outcomes. Therefore,
107	there is a risk of underestimating the independent effects of disease, and
108	overestimating the independent effects of nutritional issues. The aim of this study
109	was to take into account disease type and severity and explore associations
110	between: (1) nutritional status; (2) food intake; and health-related outcomes (LOS,
111	mortality, and readmissions) in participants from the ANCDS.
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126 METHODS

The ANCDS was conducted in two phases. Participants were recruited in Phase 1 of 127 the study and the episode of admission was referred to as "index hospitalisation". 128 In Phase 1 data were collected by dietitians from participating hospitals [1]. Data 129 included demographic, nutritional status, and 24-hour food intake information for 130 each participant [1]. Participants' body mass index (BMI) were calculated based on 131 their recorded weight and height [1]. To evaluate nutritional status, each participant 132 was screened using the Malnutrition Screening Tool (MST) [11] and those deemed at 133 risk of malnutrition underwent comprehensive nutritional assessment using 134 135 Subjective Global Assessment (SGA) [12]. Based on the International Classification of Disease and Related Health Problems (ICD-10-AM) [13], malnutrition was defined 136 as BMI <18.5 kg/m² and an SGA rating of moderately malnourished (SGA-B) or 137 severely malnourished (SGA-C). Over a 24-hour period, each participants' 138 percentage food intake was observed and recorded by meal and snack on a five-139 point scale (0%, 25%, 50%, 75%, and 100%) [1]. Information on the prescribed diet 140 on the day of the survey was also recorded [11]. 141 The present study (Phase 2) is a prospective cohort study and includes participants 142 143 from Phase 1. Data were collected 90 days post Phase 1 and includes: Admission-related data: Nature of admission, type and severity of disease, 144 discharge status (Appendix 1); 145 • Outcomes-related data: Length of stay, readmissions, date of death (Appendix 146 147 1).

Quality of life data: Participants' self-perceived quality of life was assessed
 using EQ-5D [14], a non-disease specific two part questionnaire (Appendix 1).

- 150 Ethical approval for the present study was provided by the Medical and Research
- 151 Ethics Committee of The University of Queensland and local Human Research Ethics
- 152 Committees of participating hospitals. Data were collected in accordance with the
- 153 ethical standards of the ethics committees.
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155 Statistical Analysis

- Data were analysed using PASW Statistics 18. The following variables weredichotomised:
- Age- < 65 years, ≥ 65 years;
- PCCL scores- not severe/catastrophic PCCL (i.e. PCCL score of 0, 1 or 2),
- severe/catastrophic PCCL (i.e. PCCL score of 3 or 4);
- EQ-5Dprofile (i.e. each of the five dimensions (mobility, self-care, activity,

pain/discomfort, anxiety/depression)- no problem, some problem (included

- 163 moderate/severe problem)[14];
- Nutritional status: Malnourished (included SGA-B[12], SGA-C[12], and patients

with BMI <18.5 kg/m² [13]), well-nourished (included MST < 2 [11]and SGA-A[12]);

- Food Intake- Since food intake of ≤25% (i.e. nil-by-mouth (NBM), 0%, 25% food
- 167 consumption during Phase 1 of the survey) was significantly associated with the
- outcomes at the bivariate level, food intake was dichotomised as $\leq 25\%$ and $\geq 50\%$
- (i.e. 50%, 75%, and 100% food consumption during Phase 1 of the survey).
- 170 Appendix 2 describes the steps undertaken to clean the dataset for outcomes
- 171 variables.
- 172 All categorical variables were reported as frequency and percentage. The distribution
- of LOS, as a continuous variable, was analysed. Length of stay remained skewed
- after trimming, and is therefore reported using median (range). LOS was transformed

by using the square root for analysis. Bivariate analyses were undertaken using chi-175 square tests for categorical variables and independent sample t-tests or equivalent 176 non-parametric t-tests for continuous variables, to identify significance between 177 confounders and outcome variables. Variables considered as risk factors from the 178 literature (confounding variables) and those demonstrating a significant association 179 with each outcome variable at a bivariate level (evaluable confounding variables) 180 were entered into regression models (Appendix 3). Preliminary assumption testing 181 were conducted to ensure no violation of the assumptions, including multicollinearity. 182 High intercorrelations were observed between diet type and nutritional status, and 183 184 therefore diet type was excluded from the regression models. A *p*-value < 0.05 was 185 considered statistically significant.

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188 **RESULTS**

Outcomes data were available for 3017 of the total 3122 participants (97%). After data cleaning (as previously outlined), data analyses for LOS and mortality included 2982 participants (95%), and readmissions data were analysed for 2942 participants (94%).

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Table 1 depicts admission-related characteristics of the participants. Malnutrition was significantly associated with age \geq 65 years, emergency admissions, admissions other than surgical or medical, certain MDCs, severe/catastrophic PCCL scores, discharge status (excluding those who left against medical advice), EQ-5D_{profile} and EQ-5D_{vas} scores, and pre-survey LOS (Table 1). Consumption of \leq 25% of the offered hospital food was significantly associated with age \geq 65 years, certain MDCs, surgical and medical admissions, severe/catastrophic PCCL scores, EQ-5D_{profile} and EQ-5D_{vas} scores (Table 1). Participants who consumed ≥ 50% of the offered food were more likely to be discharged to their home/place of usual residence (Table 1). Percentage food intake was not associated with pre-survey LOS (Table 1).

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LOS: The median LOS for all patients was 11 days (Table 2) with 67 patients (2%)

having a LOS of ≥90 days. Malnourished participants had longer median LOS (15

days, range: 2 – 119 days) compared to well-nourished participants (median LOS: 10

days, range: 2 – 158 days) (p < 0.0001) (Table 2). Severely malnourished

209 participants (SGA-C) had a significantly longer median LOS (21 days, range: 2 – 259

210 days) versus well-nourished participants (12 days, range: 2 – 291 days) and

211 moderately malnourished (SGA-B) participants (15 days, range: 2 – 467 days) (p <

212 0.0001). The median LOS of participants who consumed $\leq 25\%$ of the offered food

was longer (13 days, range: 2 - 158 days) than those who consumed $\ge 50\%$ of the

food (11 days, range: 2 – 119 days) (p < 0.0001) (Table 2).

The multiple regression analysis model explained 32% of the variance in LOS (R^2 =

216 0.329, adjusted R^2 = 0.319, F (34, 2290) = 32.95, p < 0.0001). PCCL scores were the

217 largest unique contribution (beta: 0.353, CI: 0.417 – 0.513, p-value < 0.0001).

Nutritional status made a statistically significant contribution (beta: 0.084, CI: 0.167 –

219 0.414, *p*-value < 0.0001). Percentage food intake was not significant.

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Readmissions: The overall readmission rate was 30% (n= 882) (Table 2) within 90days from post-index hospitalisation. While malnourished patients had a significantly
higher readmission rate (35%) in comparison to well-nourished patients (27%), no
association was found between percentage food intake and readmissions (Table 2).

An ordinal regression model did not find malnutrition to be a significant risk factor for readmissions. Neoplastic disease and discharge to other healthcare facilities were the highest risk factors for significantly increasing the odds of readmissions within 90 days of index hospitalisation (Table 3).

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Mortality: The 30-day and 90-day in-hospital mortality rate were 1.5% (n= 46) and 2.4% (n= 72) respectively (Table 2). Malnourished patients and those who ate \leq 25% of the offered food had significantly higher mortality rates than others (Table 2). Risk factors for in-hospital mortality have been included in Tables 4a and 4b. Logistic regression analysis revealed:

- Although malnutrition was not an independent risk factor for 30-day in-hospital
 mortality (Table 4a) it increased the odds of 90-day in-hospital mortality by
 almost two times (OR: 1.91, CI: 1.09-3.34, p= 0.023) (Table 4b).
- Eating ≤ 25% of the offered food increased the risk of 30- and 90-day in-
- 239 hospital mortality by > 2.5 times (OR: 2.69, CI: 1.31 5.52, p= 0.007) (Table

4a) and 2 times (CI: 1.13 - 3.51, p = 0.017) respectively (Table 4b).

• Severe/catastrophic PCCL score and age \geq 65 years were independent risk

factors common for both, 30- and 90-day in-hospital mortality (Tables 4a, 4b).

- 243 The hazard ratio of 90-day in-hospital mortality for malnourished patients who
- consumed \leq 25% of the offered food was 2.3 times greater (CI: 1.39-3.76, *p*= 0.001)
- than well-nourished patients (Table 5; Figure 1).
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250 **DISCUSSION**

The ANCDS is the first multicentre study in acute care hospitals across Australia and 251 New Zealand to report the association between patients' nutritional status, food 252 intake and health-related outcomes. The study found that patients who were 253 malnourished or consumed $\leq 25\%$ of the hospital offered food had significantly longer 254 LOS and higher in-hospital mortality rates. Malnourished patients also had 255 significantly higher readmissions rates than well-nourished patients. 256 Considering there are several non-nutritional factors that can influence LOS [15], 257 readmissions [16], and in-hospital mortality, it is important to account for these 258 259 factors. Although three studies have previously used multivariate regression analyses to control for the effect of confounders in a general, adult acute care population [2, 5, 260 6], they have limited comparability as they did not control for disease severity. They 261 262 also did not evaluate readmissions as an outcome [2, 6], participants' food intake [2, 5] or participants' nutritional status using validated and reliable methods [6]. To the 263 best of our knowledge, the ANCDS is the only study to control for disease severity 264 (using PCCL scores) and other non-nutritional factors (age, gender, disease type, 265 QoL indicators) in multivariate regression models to report the independent 266 267 association between malnutrition and poor food intake and LOS, readmissions, and mortality in a general, adult acute care population. Multivariate regression analyses 268 confirmed that non-nutritional factors associated with all three outcomes were 269 270 severe/catastrophic disease severity and age \geq 65 years. Respiratory disease was a common risk factor for readmissions and 90-day in-hospital mortality. 271 LOS: Three other studies have used regression analyses to report associations 272 between malnutrition and LOS [2, 5, 17]. Pirlich et al used number of prescriptions 273 per day as a surrogate marker for disease severity, although they acknowledged the 274

limitation of this method [17]. Lim et al did not control for disease severity per se, 275 276 however, they used the DRG-matching technique and controlled for diagnosis, investigations, and treatment costs. Their study demonstrated that malnutrition was 277 an independent risk factor for longer LOS [5]. Other nutrition studies have not 278 controlled for disease severity [2] while establishing associations between 279 malnutrition and LOS. Results from the ANCDS establish that malnutrition is a 280 contributor to prolonged LOS, independent of the disease status. 281 Studies evaluating the association between food intake and LOS in hospitals are 282 extremely limited and conflicting. Kandiah et al reported a positive association 283 284 between extended LOS and greater plate waste [18]. Conversely, Dupertuis et al found that patients with a hospital LOS of more than eight days were less likely to "be 285 underfed" and speculated that the extended duration of hospital stay helped with 286 287 adapting to the taste of hospital food, and mealtimes [19]. The present study could not find a significant difference in the median pre-survey LOS of patients consuming 288 \leq 25% of the hospital offered food versus those consuming \geq 50% of the food. Given 289 that the present study demonstrated a significant association between malnutrition 290 291 and LOS, and poor food intake during hospitalisation is a risk factor for malnutrition, it 292 is important to recognise and provide timely nutrition support to patients with poor food intake during hospital admission. 293

Readmissions: The ANCDS reported that one-in-three patients (30%) are
readmitted within three months of index hospitalisation. The readmission rate at three
months in this study is substantially higher than the 19 – 24% rate previously
reported [20].

Although analyses found that the readmission rate of malnourished patients was 1.3 times higher than that of well-nourished patients, this effect was lost during ordinal

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regression analysis. Five previous studies have reported a positive association 300 301 between malnutrition and readmissions [3-5, 21, 22]. The findings from three of these studies cannot be compared to the present study as they were conducted in small 302 cohorts of participants \geq 50 years of age, and used anthropometric and/or 303 biochemical measures to define malnutrition [3, 21, 22]. The findings by Planas et al 304 have limited application as despite having a larger cohort and using a validated 305 method to define malnutrition (i.e. SGA), they did not control for the effect of 306 confounding variables [4]. The study by Lim et al is comparable as they included a 307 large cohort (n: >800 participants, age: >18 years), used the validated SGA to define 308 309 malnutrition, and controlled for various confounders (age, gender, ethnicity, DRG) [5]. Similar to the ANCDS, their study could not find an association between malnutrition 310 and readmissions within 90-days of index hospitalisation [5]. However, they found 311 312 that malnourished patients had a 60% higher readmission risk within 15-days posthospital discharge [5]. It was beyond the scope of this study to record the nutritional 313 status of the participants at each episode of readmission. Further research evaluating 314 the effectiveness of hospital- and/or community-based nutrition interventions in 315 preventing readmissions will be valuable in filling this gap in the literature. 316 317 The ANCDS found that neoplastic disease, discharge destinations, severe/catastrophic disease severity, and age \geq 65 years were associated with 318 increased readmissions. Several studies, as summarised in one meta-analysis [16] 319 320 and two systematic reviews [23, 24], have previously reported these associations. *Mortality:* The ANCDS also found that malnourished patients consuming $\leq 25\%$ of 321 the offered food had more than a two-fold risk of 90-day in-hospital death compared 322 to well-nourished patients who consumed at least half the offered food. This effect 323 was not significant for 30-day in-hospital mortality. Our results contrast with the 324

nutritionDay Survey by Hiesmayr et al, which was also a one-day multicentre study 325 326 (involving >16000 patients from >250 hospitals in 25 European countries), reported an adjusted hazard ratio of 2.10 (CI: 1.53 – 2.89) for 30-day in-hospital mortality in 327 patients who consumed a guarter of the offered meal [6]. More detailed analysis of 328 disease severity and nutritional status characteristics of the sub-group of patients in 329 the ANCDS who experienced 30-day in-hospital mortality indicated that there was no 330 significant difference in the number of well-nourished (n= 20, 45%) and malnourished 331 patients (n= 24, 55%) (p > 0.05) and that a majority of these patients (n= 44, 96%) 332 had a severe/catastrophic PCCL score during index hospitalisation. Since disease 333 334 severity is associated with increased mortality, and highly correlated with 335 malnutrition, this could explain why malnutrition was not a significant independent risk factor for 30-day in-hospital mortality. 336

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LIMITATIONS: The ANCDS could record readmissions only within participating
 hospitals. Even though the readmission rate was higher than that reported by other
 studies, considering that readmissions to other hospitals can account for
 approximately 25% of all readmissions [25], this study may have underreported
 readmission rates.

The ANCDS has provided in-hospital mortality data only. Mortality data for those that may have occurred post-discharge in a different setting were not recorded making it likely that mortality rates may have also been underreported in this study.

Participating hospitals represent at least 20% of acute care hospitals in Australia [26] and 40% of acute care hospitals in New Zealand [27] (that have more than 60 beds) limiting the generalisability of the results across the acute care population in Australia and New Zealand. Nevertheless, the ANCDS is the first and largest multicentre study to provide a snapshot of the association between malnutrition, poor food intake andpatient outcomes in this region.

The ANCDS reported point prevalence malnutrition for a majority of the patients and 352 food intake was recorded for a 24-hour period only. In addition, being a cross-353 sectional observational study it cannot determine if poor food intake caused in-354 hospital mortality within 30-days of hospital admission. It is noteworthy that 355 regardless of the type and severity of disease, age, nutritional status, and other 356 potential confounders, consuming $\leq 25\%$ the offered food (during Phase I) 357 independently increased the odds for 30- and 90-day in-hospital mortality. It was 358 359 beyond the scope of this study to calculate the nutritional intake for participants who consumed ≤25% the offered food; however, it can be speculated that consumption of 360 ≤25% of the offered food would be unlikely to meet participants' nutritional 361 362 requirements.

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STRENGTHS: The ANCDS is the first study to highlight the independent association 364 of malnutrition and poor food intake during hospitalisation on health-related outcomes 365 in Australian and New Zealand acute care patients, after controlling for various 366 367 confounders including disease type and severity. Evidence regarding the association between poor food intake and negative outcomes is scarce. Even though previous 368 studies have reported the association between nutritional status and negative 369 370 outcomes, they have seldom controlled for disease severity and other confounding factors, thus providing an incomplete analysis of association. It is possible that 371 controlling for disease severity was a challenge for previous studies, particularly 372 when there is no universally accepted measure for disease severity [6]. There are a 373 variety of generally accepted comorbidity indices [28] that can reduce all the 374

coexisting diseases and their severities to a single score to allow comparisons with 375 376 other patients with the same score [28]. However, they measure comorbidity at a given time and are either designed for a specific patient group or consist of a limited 377 number of disease categories [28]. The ANCDS cohort was anticipated to include 378 patients with a vast variety of acute care condition/s, limiting the application of any 379 particular comorbidity index. In addition, comorbidity indices require data abstraction 380 by reviewing patients' medical charts [28]. Given the large cohort, it would not only be 381 time-consuming and impractical to review individual hand-written medical charts to 382 record each participants' comorbidities, missing data would also be a risk [28]. 383 384 Therefore, the ANCDS used a novel approach to overcome the challenge of 385 controlling for disease type and severity- by using diagnostic codes (AR-DRG) and PCCL scores respectively. Moreover, PCCL scores are reflective of the cumulative 386 effect of patients' complications and comorbidities for the entire episode of 387 admission, and thus a more accurate measure of patients' disease severity. 388 389

PRACTICAL IMPLICATIONS: The ANCDS is the first study that we know of which 390 demonstrates that poor food intake, independent of disease type and severity, and 391 392 malnutrition is associated with in-hospital mortality in acute care patients. While onein-three malnourished patients consumed $\leq 25\%$ of the offered food, one-in-five well-393 nourished patients also consumed ≤ 25% of the offered food [1]. These results call 394 395 for more consistent monitoring of hospitalised patients' food intake levels. Perhaps protocols for recording patients' food intake after each meal need to be implemented 396 397 akin to those for authorising medication charts soon after medications are administered. In light of our results, and those from the European NutritionDay 398

Survey, perhaps consumption of ≤25% of the offered food should be used as a
screening (and rescreening) tool to commence appropriate medical nutrition therapy.

CONCLUSION: The ANCDS is the first multicentre study in acute care patients 402 across Australia and New Zealand to examine the association between nutritional 403 status and food intake, and health-related outcomes (LOS, readmissions, and in-404 hospital mortality), after controlling for a range of confounding factors (including 405 disease type and severity). The ANCDS confirms that malnutrition and poor food 406 intake have independent associations with health-related outcomes in acute care 407 408 patients. Both these risk factors are modifiable, in contrast to other risk factors such as age and disease. Findings from the ANCDS accentuate the importance of 409 implementing every step of the nutrition care process (nutrition screening and 410 411 assessment, nutrition support, nutrition monitoring and evaluation of nutrition support) as standardised practice across acute care hospitals. 412

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424 **Authors' contributions to manuscript:**

EA designed and coordinated the study; acquired, analysed and interpreted the data;

and wrote the manuscript. MF, MB and EI provided significant advice on the study

- 427 design. MBatterham provided statistical advice. All authors participated in editing and
- 428 final revisions of the manuscript. All authors have read and approved the final
- 429 manuscript.
- 430
- 431 Conflict of Interest: EA, MBatterham, JB, and SC have no conflict of interest to
 432 declare. MF, MB and EI are employed by Queensland Health, Australia.
- 433

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 Table 1: Admission-related characteristics of the participants:

Characteristics	Overall Results	<u>Results as</u>	s per Nutritional S	<u>Status</u>		s as per % Fe	bod
		Well- nourished ^a	Malnourished ^b	<i>p</i> - value	≥50% intake ^c	≤ 25% intake ^d	<i>p</i> -value
Age group ^e				٦			
< 65 years	1314 (44%)	911 (46%)	382 (40%)	0.003	981 (43%)	316 (48%)	<u>၂</u> 0.028
≥ 65 years	1650 (56%)	1064 (54%)	564 (60%)	J	1294 (57%)	343 (52%)	ſ
Admission Status (n (%)) ^e							
Emergency	2173 (73%)	1426 (72%)	719 (76%)	0.027	1669 (73%) ^g	483 (73%) ^g	0.935
Elective	623 (21%)	433 (22%) ^g	180 (19%) ^g	0.075	468 (21%) ^g	148 (22%) ^g	0.302
Other ^h	183 (6%)	127 (6%) ^g	51 (5%) ^g	0.321	149 (7%) ^g	32 (5%) ^g	0.110
Main Diagnostic Categories (MDC) (n (%)) ^e	• •		• •		· ·		
Digestive, Hepatobilliary	562 (19%)	335 (17%)	222 (23%)	0.000	351 (15%)	206 (31%)	0.000
Musculoskeletal	445 (15%)	326 (16%)	108 (11%)	0.000	348 (15%) ^g	86 (13%) ^g	0.150
Circulatory	388 (13%)	295 (15%)	87 (9%)	0.000	329 (14%)	55 (8%)	0.000
Respiratory	372 (13%)	231 (12%)	135 (14%)	0.045	296 (13%) ^g	76 (12%) ^g	0.311
Nervous	277 (9%)	192 (10%) ^g	80 (8%)g	0.285	220 (10%) ^g	56 (9%) ^g	0.360
Skin, Subcutaneous Tissue, Burns, Breast	124 (4%)	100 (5%)	24 (2%)	0.002	113 (5%)	10 (2%)	0.000
Kidney, Urinary Tract	109 (4%)	64 (3%) ^g	43 (5%) ^g	0.075	90 (4%) ^g	17 (3%) ^g	0.096
Others	110 (4%)	64 (3%)	45 (5%)	0.041	96 (4%) ^g	14 (2%) ^g	0.013
Pre-MDC	100 (3%)	54 (3%)	46 (5%)	0.003	58 (3%)	40 (6%)	0.000
Infectious & Parasitic	99 (3%)	56 (3%)	42 (4%)	0.023	77 (3%) ^g	22 (3%) ^g	0.950
Neoplastic	82 (3%)	46 (2%)	36 (4%)	0.023	67 (3%) ^g	15 (2%) ^g	0.357
Endocrine, Metabolic and Nutritional	82 (3%)	56 (3%) ^g	25 (3%) ^g	0.779	64 (3%) ^g	16 (2%) ^g	0.590
Injuries, Poisoning, Drug and Alcohol abuse	80 (3%)	56 (3%) ^g	20 (2%) ^g	0.258	62 (3%) ^g	17 (3%) ^g	0.836
Male & Female Reproductive System	72 (2%)	56 (3%) ^g	16 (2%) ^g	0.064	55 (2%) ^g	17 (3%) ^g	0.816
Eye, Ear-Nose-Throat, Mouth	42 (1%)	35 (2%)	6 (1%)	0.015	32 (1%) ^g	10 (2%) ^g	0.835
Blood & Blood-forming Organs	31 (1%)	19 (1%) ^g	12 (1%́) ^g	0.443	25 (1%) ^g	5 (1%) ^g	0.443
Partition (Admission type) (n (%)) ^e							
Surgical	1270 (43%)	847 (43%) ^g	403 (43%) ^g	0.953	886 (39%)	369 (56%)	0.000
Medical	1547 (52%)	1044 (53%) ^g	482 (51%) ^g	0.390	1279 (56%)	255 (39%)	0.000
Other	158 (5%)	94 (5%)	62 (7%)	0.041	118 (5%) ^g	38 (6%) ^g	0.563

Patient Clinical Complexity Level Scores							
(PCCL) (n (%)) ^e Not severe	1145 (39%)	887 (45%)	244 (26%)	0.000	933 (41%)	200 (30%)	0.000 ر
Severe/Catastrophic	1821 (61%)	1096 (55%)	696 (74%)	5.000	1344 (59%)	459 (70%)	5 0.000
Discharge Status (n (%)) ^e	- (/						
Usual Residence	2129 (74%)	1521 (79%)	576 (65%)	0.000	1667 (75%)	440 (71%)	0.024
Other Hospital	303 (11%)	177 (9%)	123 (14%)	0.000	224 (10%) ^g	74 (11%) ^g	0.198
Other Facility ⁱ	423 (14.5%)	231 (12%)	185 (21%)	0.000	317 (14%) ^g	103 (17%) ^g	0.161
Left Against Medical Advice	9 (0.5%)	5 (0.3%) ^g	4 (0.4%) ^g	0.401	5 (0.2%) ^g	4 (0.6%) ^g	0.102
EQ-5D _{profile} (n (%)) ^e : Some/Major Problem with:							
Mobility	1870 (64%)	1181 (60%)	667 (72%)	0.000	1422 (63%)	432 (68%)	0.014
Pain	1846 (63%)	1189 (61%)	634 (68%)	0.000	1376 (61%)	451 (71%)	0.000
Self-Care	1296 (45%)	772 (40%)	510 (55%)	0.000	934 (42%)	349 (55%)	0.000
Anxiety/Depression	1246 (43%)	727 (38%)	507 (55%)	0.000	919 (41%)	316 (51%)	0.000
Activity	1893 (65%)	1171 (60%)	699 (75%)	0.000	1412 (63%)	460 (73%)	0.000
EQ-5D _{vas} (median (range)) ^f	51 (0 - 100)	60 (0 - 100)	50 (0 - 100)	0.000	58 (0 - 100)	50 (0-100)	0.000
Pre-survey Length of Stay (median (range)) ^f	6 (0 - 449)	5 (0 - 364)	9 (0 - 449)	0.000	6 (0 - 449)	6 (0 - 364)	0.459

^a Well-nourished participants [1]: included those not at risk of malnutrition (MST[28]) and SGA-A[26]

^b Malnourished participants [1]: included moderately (SGA-B)[26] and severely (SGA-C)[26] malnourished participants, and patients with

BMI < 18.5 kg/m2 [27]

 $^{c} \ge 50\%$ intake includes 50%, 75% and 100% food intake

 $^{\rm d}$ \leq 25% intake includes nil-by-mouth, 0%, and 25% intake

^e Categorical variables represented as n (%)

^f Continuous Variable presented as Median (Range) for data that is not normally distributed

^g non-significant (*p*-value >0.05)

^h includes waitlists and non-assigned

ⁱ includes residential aged care facility, rehabilitation, episode change within same hospital, other health facility

NOTE: Admission status data were missing for 3 participants; MDC data were missing for 9 participants; Partition data were missing for 7 participants; PCCL data were missing for 16 participants; Discharge Status data were missing for 78 participants; EQ-5Dprofile: Mobility data were missing for 62 participants, Pain data were missing for 64 participants, Self-care data were missing for 69 participants, Anxiety/Depression data were missing for 83 participants, Activity data were missing for 82 participants, EQ-5D_{vas} data were missing for 249 participants, Pre-survey Length of Stay data were missing for 17 participants.

Table 2: Comparison of outcomes by participants' nutritional status and 24-hour % food intake bivariate level

	Overall <u>As per Nutritional Status</u>			As per % food intake			
Variables	Results	Well- nourished ^a	Malnourished ^b	<i>p</i> -value	≥50% intake ^c	≤ 25% intake ^d	<i>p</i> -value
Length of Stay (LOS) (days) ^e	11 (2 – 158)	10 (2 – 158)	15 (2 – 119)	0.000	11 (2 – 119)	13 (2 – 158)	0.000
Readmissions ^f (n (%)):			· · ·			· · ·	
1 readmission (n (%))	564 (19%)	349 (18%)	206 (23%)	J	435	122	J
2 readmissions (n (%))	198 (7%)	127 (6%)	66 (7%)	ര.000	161	35	0.378 ⁹
\geq 3 readmissions (n (%))	120 (4%)	68 (3%)	49 (5%)	J	88	31	J
Mortality ^f :	· · ·	• •	• •				
90 day in-hospital mortality (n (%)) ^h	72 (2.4%)	28 (1%)	43 (5%)	0.000	40 (2%)	32 (5%)	0.000
30-day in-hospital mortality (n (%))	46 (1.5%)	22 (1%)	23 (2.5%)	0.010	25 (1%)	21 (3%)	0.001

^a Well-nourished participants [1]: included those not at risk of malnutrition (MST[28]) and SGA-A[26]

^b Malnourished participants [1]: included moderately (SGA-B)[26] and severely (SGA-C)[26] malnourished participants, and patients with

BMI < 18.5 kg/m2 [27]

- $^{c} \ge 50\%$ intake includes 50%, 75% and 100% food intake
- $d \le 25\%$ intake includes nil-by-mouth, 0%, and 25% intake
- ^e Continuous Variable presented as Median (Range) for data that is not normally distributed
- ^f Categorical variables represented as n (%)
- ^g non-significant (*p*-value >0.05)
- ^h Includes 30-day in-hospital mortality results

Risk Factors	Bivariate Analyses			Ordinal Regression Analyses		
	Readmissions n (%)	No readmissions n (%)	<i>p</i> - value	Odds Ratio	Cl (<i>p</i> - value)	
MDC: Neoplastic	35 (43%)	47 (57%)	0.032	1.55	1.20 - 1.99 (0.001)	
Discharge to Other Facility ^a	210 (50%)	209 (50%)	<0.001	1.43	1.16 – 1.51 (0.000)	
Discharge to Usual Residence	633 (30%)	1465 (70%)	<0.001	1.33	1.16 – 1.51 (0.000)	
Severe/Catastrophic PCCL score	650 (36%)	1171 (64%)	<0.001	1.30	1.18 – 1.43 (0.000)	
Medical Partition	571 (37%)	976 (63%)	<0.001	1.22	1.00 - 1.48 (0.049)	
MDC: Respiratory	145 (39%)	227 (61%)	0.005	1.15	1.00 - 1.31 (0.048)	
Age ≥ 65 years	587 (36%)	1063 (64%)	<0.001	1.11	1.02 – 1.22 (0.021)	
EQ _{vas} score ^b	50 (0 - 100)	55 (0 - 100)	<0.001	1.00	1.00 - 1.004 (0.044)	
Malnutrition ^c	346 (36%)	605 (64%)	0.001	1.06 ^d	1.04 – 1.17 (0.235) ^d	

Table 3: Bivariate and Ordinal Regression results for readmissions within 90-days of index hospitalisation (N= 3017)

CI: Confidence Intervals; MDC: Major Diagnostic Category; PCCL: Patient Clinical Complexity Level

^a includes residential aged care facility, rehabilitation, episode change within same hospital, other health facility

^b Represented as median (range)

^cMalnutrition[1]: included moderately (SGA-B)[26] and severely (SGA-C)[26] malnourished participants, and patients with BMI < 18.5

kg/m2 [27]

^d non-significant (*p*-value >0.05)

Table 4a: Bivariate and Logistic Regression results for 30-day in-hospital mortality (N=3017)

Risk Factors		Bivariate Analyses	Logistic Regression Analyses		
-	In-hospital mortality n (%)	No in-hospital mortality n (%)	<i>p</i> - value	Odds Ratio	CI (p- value)
Severe/Catastrophic PCCL score	44 (3%)	1745 (97%)	<0.001	8.18	1.93 - 34.73 (0.004)
MDC: Respiratory	13 (4%)	348 (96%)	0.03	1.78	0.81 – 3.93 (0.151) ^a
≤ 25% Food Intake	21 (3%)	629 (97%)	0.001	2.69	1.31 - 5.52 (0.007)
Malnutrition ^b	23 (3%)	906 (97%)	0.01	1.27	0.63 – 2.59 (0.504)
Age ≥ 65 years	40 (3%)	1573 (97%)	<0.001	2.74	1.11 - 6.79 (0.03)
EQ _{vas} score ^c	50 (0 - 85)	51 (0 - 100)	0.03	0.99	0.98 – 1.01 (0.215)

CI: Confidence Intervals; PCCL: Patient Clinical Complexity Level; MDC: Major Diagnostic Category

^a non-significant (*p*-value >0.05)

^b Malnutrition [1]: included moderately (SGA-B) [26] and severely (SGA-C) [26] malnourished participants, and participants with BMI <

18.5 kg/m²[27].

^c Represented as median (range)

Table 4b: Bivariate and Logistic Regression results for 90-day in-hospital mortality (N=3017)

Risk Factors		Bivariate Analyses	Logistic Regression Analyses		
-	In-hospital mortality n (%)	No in-hospital mortality n (%)	<i>p</i> - value	Odds Ratio	CI (p- value)
Severe/Catastrophic PCCL score	68 (4%)	1721 (96%)	<0.001	6.01	2.14 - 16.89 (0.001)
MDC: Respiratory	19 (5%)	342 (95%)	0.001	1.91	1.01 - 3.61 (0.047)
≤ 25% Food Intake	32 (5%)	618 (95%)	<0.001	1.99	1.13 – 3.51 (0.017)
Malnutrition ^b	43 (5%)	886 (95%)	<0.001	1.91	1.09 - 3.34 (0.023)
Age ≥ 65 years	58 (4%)	1555 (96%)	<0.001	2.23	1.15 - 4.34 (0.018)
EQ _{vas} score ^c	43 (0 - 99)	51 (0 - 100)	<0.001	0.98	0.97 - 0.99 (0.015)

CI: Confidence Intervals; PCCL: Patient Clinical Complexity Level; MDC: Major Diagnostic Category

^a non-significant (*p*-value >0.05)

^b Malnutrition [1]: included moderately (SGA-B) [26] and severely (SGA-C) [26] malnourished participants, and participants with BMI <

18.5 kg/m²[27].

^c Represented as median (range)

Risk Factor	Hazard Ratio	CI (p- Value)
Surgical Partition	3.03	1.06 - 8.69 (0.039)
Medical Partition	3.71	2.01 - 6.85 (0.000)
Age ≥ 65 years	2.84	1.53 - 5.29 (0.001)
Severe/Catastrophic	3.55	1.27 – 9.92 (0.016)
PCCL		
≤ 25% Food Intake	2.29	1.39 – 3.76 (0.001)
Confidence Interval: PCC	I · Dationt Clinica	Complexity Level

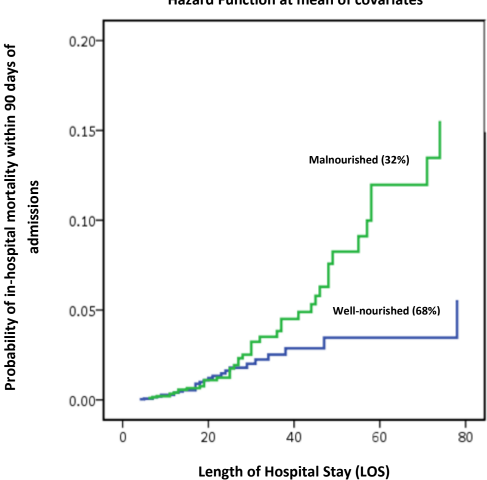
Table 5: Hazard Analysis of risk factors and 90-day in-hospital mortality (N= 3017)

CI: Confidence Interval; PCCL: Patient Clinical Complexity Level

Note: Other risk factors such as Main Diagnostic Categories, Admission Status, and

gender were not significant

Figure 1: Cumulative incidence of 90-day in-hospital mortality in well-nourished and malnourished patients (N= 3017)



Hazard Function at mean of covariates

Admission-	Admission	Whether it was an emergency, elective or other
related:	Status	admission
	Australian Refined Diagnosis Related Group (AR-DRG)	Refers to Australia's national diagnosis related care (DRG) classification scheme that provides a clinically meaningful way for relating the number and types of patients treated in hospitals to the resources required by the hospitals [29]. AR- DRGs are assigned based on Principal Diagnosis [29]. While New Zealand used version 5.0 of the AR-DRGs, hospitals in Australia used a range of versions (4.2, 5, 5.1, 5.2, and 6). Since the study cohort represented a large number of AR-DRGs (n= 685) it was necessary to simplify the categorisation of participants by disease type. Major Diagnostic Categories (MDCs), which are based on a single body system or aetiology that is associated with a medical speciality and therefore include AR-DRGs and principal diagnoses [29], were used for this purpose. Since MDCs are uniform across various AR-DRG versions, categorising the type of disease into MDCs maintained consistency across the AR-DRG
		versions.
	Partition	MDCs are sub-divided into a maximum of three separate partitions or type of admissions: surgical, medical, and other. The presence or absence of operating room and non-operating room procedures is generally responsible for the assignment of the episode of admission to one or other of these partitions [29].
	Patient Clinical Complexity Level (PCCL) scores	refers to the cumulative effect of a patient's complications and comorbidities [29]. The calculation of these scores is a complex process and is designed to prevent similar conditions from being counted more than once [29]. PCCL scores are calculated for each episode of admission and range from $0 - 4$ (for surgical episodes) and from 0 - 3 (for medical episodes) and are defined as follows [29]: 0 = not a complication or comorbidity 1 = a minor complication or comorbidity 2 = a moderate complication or comorbidity 3 = a severe complication or comorbidity 4 = a catastrophic complication or comorbidity.
	Discharge Status	refers to the discharge destinations of the participants after index hospitalisation. The following categories were used: Home/Usual residence Other hospital

Appendix 1: Data collected for each participant:

		 Other healthcare facility- included residential aged care facility, rehabilitation, episode change within same hospital, other health facility Left against medical advice Death 		
Outcomes- related	Pre-survey LOS	Was computed as the difference between the date of the survey and date of admission. This was done to evaluate if length of hospital stay impacts food intake.		
	Index LOS	Refers to the LOS for the index hospital admission (i.e. hospital admission during which participants were enrolled in Phase 1 of the study). It was computed as the difference between date of discharge and date of index hospital admission.		
	Date of Death	Was used to compute the number of days between date of admission and date of in-hospital death.		
	Readmissions	Were recorded, along with the frequency of readmissions, for up to 90 days from the date of index hospitalisation.		
Quality of life				

Appendix 2: Steps undertaken to clean the dataset for outcomes variables:

Outcome	Steps undertaken to clean the dataset	
Length of Stay (LOS)	Since LOS was positively skewed and varied across the Major Diagnostic Categories (MDC); trimming (deleting) LOS methodology was used to prevent outliers from having a significant and unrepresentative impact on the average LOS. The following steps were followed to trim the LOS data [30]: Step 1: Patients were excluded based on the following criteria [30]: • Death during index hospitalisation;	
	 Death during index hospitalisation, Missing data values for: LOS, age, discharge status, MDC, admission source, admission status, PCCL; Discharge against medical advice. Step 2: Upper and lower trim points were calculated for each 	
	MDC as per the following equations [30]: Lower Trim Point= Q1 – (3*IQR); Upper Trim Point= Q3 + (3*IQR) where:	
	 Q1: the first quartile of all patients records from the LOS dataset 	
	 Q3: the third quartile of all patients records from the LOS dataset IQR: Q3 – Q1 	
	Step 3: Since the lower trim points for MDCs were in negative values, participants with LOS > upper trim points for each MDC were excluded [30]. Participants with LOS= 1 day were also excluded as their admissions were more likely to be associated with clinical investigations or tests.	
Readmissions	Participants who died during index hospitalisation were excluded from the analyses related to readmissions data.	
In-hospital mortality	Participants who were not discharged within 90 days of index hospital admission were included in the analyses.	

Appendix 3: Regression Models used for evaluating the association between

Outcome variables	Regression Model used	Confounding variables ^a	Evaluable confounding variables ^b
LOS (square root)	Linear regression model	Partition, MDCs[31], age group[31], admission status, disease severity[31] (dichotomised PCCL score), nutritional status[31]	Dichotomised EQ-5D _{profile} , EQ-5D _{vas} score, dichotomised percentage food intake (i.e. ≤25% and ≥50%)
Readmission	Ordinal regression model	LOS [21], surgical admission, medical admission, MDCs (respiratory, neoplastic), age group[31], disease severity [31](dichotomised PCCL score), discharge status (home/usual residence, other healthcare facility)	Nutritional status, dichotomised EQ-5D _{profile} (excluding pain), EQ-5D _{vas} score
Mortality	Logistic regression model	Emergency admissions, surgical admissions, respiratory disease, disease severity[31] (dichotomised PCCL score), age group[31]	Nutritional status, dichotomised % food intake, dichotomised EQ-5D _{profile} (mobility, self-care), EQ- 5D _{vas} scores.
Hazard Analysis	Cox Regression model	Surgical and medical admission, MDCs, age group[31], gender[31], admission status, disease severity [31] (dichotomised PCCL score)	Nutritional status, dichotomised % food intake

confounding and outcome variables

EQ-5Dvas: EQ-5D visual analogue scale; LOS: length of hospital stay, MDC: major

diagnostic category; PCCL: Patient Clinical Complexity Level

^a Confounding variables: Variables that are considered risk factors as per the literature.

^b Evaluable Confounding variables: Variables that demonstrated a significant association

with the outcomes variable at a bivariate level requiring an evaluation of their significance

at a multivariate level.