

Journal Pre-proof

Nutritional risk screening (NRS 2002) is a strong and modifiable predictor risk score for short-term and long-term clinical outcomes: *Secondary analysis of a prospective randomised trial*

Lara Hersberger, MD, Laura Bargetzi, MD, Annika Bargetzi, MD, Pascal Tribolet, RD, Rebecca Fehr, RD, Valerie Baechli, RD, Martina Geiser, RD, Manuela Deiss, RD, Filomena Gomes, PhD, Alexander Kutz, MD, Nina Kägi-Braun, MD, Claus Hoess, MD, Vojtech Pavlicek, MD, Sarah Schmid, RD, Stefan Bilz, MD, Sarah Sigrist, MD, Michael Brändle, Prof., Carmen Benz, RD, Christoph Henzen, Prof., Melina Nigg, RD, Robert Thomann, MD, Claudia Brand, RN, Jonas Rutishauser, Prof., Drahomir Aujesky, Prof., Nicolas Rodondi, Prof., Jacques Donzé, Prof., Zeno Stanga, Prof., Beat Mueller, Prof., Philipp Schuetz, Prof.

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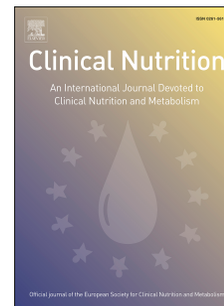
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1 **Nutritional risk screening (NRS 2002) is a strong and**
2 **modifiable predictor risk score for short-term and long-**
3 **term clinical outcomes:**

4 ***Secondary analysis of a prospective randomised trial***

5
6 ^{1,9}Lara Hersberger, MD*, ^{1,9}Laura Bargetzi, MD*, ^{1,9}Annika Bargetzi, MD*, ²Pascal
7 Tribolet, RD, ¹Rebecca Fehr, RD, ¹Valerie Baechli, RD, ¹Martina Geiser, RD,
8 ¹Manuela Deiss, RD, ^{1,13}Filomena Gomes, PhD, ^{1,9}Alexander Kutz, MD, ³Nina Kägi-
9 Braun, MD, ³Claus Hoess, MD, ³Vojtech Pavlicek, MD ³Sarah Schmid, RD, ⁴Stefan
10 Bilz, MD, ⁴Sarah Sigrist, MD ⁴Michael Brändle, Prof., ⁴Carmen Benz, RD, ⁵Christoph
11 Henzen, Prof., Melina Nigg, RD, ⁶Robert Thomann, MD, ⁶Claudia Brand, RN, ⁷Jonas
12 Rutishauser, Prof., ⁸Drahomir Aujesky, Prof., ^{8,11}Nicolas Rodondi, Prof., ^{8,12}Jacques
13 Donzé, Prof., ¹⁰Zeno Stanga, Prof., ^{1,9}Beat Mueller, Prof. ^{1,9}Philipp Schuetz, Prof.

14

15 **equally contributing first authors*

16

17 ¹ Medical University Department, Division of General Internal and Emergency

18 Medicine, Kantonsspital Aarau, Aarau, Switzerland;

19 ² Internal Medicine, Spital Lachen, Switzerland;

20 ³ Internal Medicine, Kantonsspital Muensterlingen, Switzerland;

21 ⁴ Internal Medicine & Endocrinology/Diabetes, Kantonsspital St.Gallen, Switzerland;

22 ⁵ Internal Medicine, Kantonsspital Luzern, Switzerland;

23 ⁶ Internal Medicine, Buergerspital Solothurn, Switzerland;

24 ⁷ Internal Medicine, Kantonsspital Baselland, Switzerland;

25 ⁸ Department of General Internal Medicine, Inselspital, Bern University Hospital,
26 University of Bern, Switzerland;

27 ⁹ Medical Faculty of the University of Basel, Switzerland

28 ¹⁰ Division of Diabetology, Endocrinology, Nutritional Medicine & Metabolism,
29 Inselspital, Bern University Hospital, University of Bern, Switzerland

30 ¹¹Institute of Primary Health Care (BIHAM), University of Bern, Switzerland;

31 ¹²Division of General Internal Medicine, Brigham and Women's Hospital, Boston, MA,
32 USA.

33 ¹³The New York Academy of Sciences, New York City, NY, USA

34

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36

37 **Correspondence and reprint requests:**

38 Prof. Dr. med. Philipp Schuetz, MD, MPH

39 University Department of Medicine

40 Kantonsspital Aarau

41 Tellstrasse

42 CH-5001 Aarau, Switzerland

43 Tel: +41 62 838 4141(phone)

44 Fax: +41 62 838 4100

45 E-mail: schuetzph@gmail.com

46

47

48 **Abstract**

49 **Introduction:** The Nutritional Risk Screening 2002 (NRS 2002) identifies patients at
50 risk of malnutrition. We studied the prognostic implications of this score with regard to
51 short-term and long-term clinical outcomes in a well-characterised cohort of medical
52 inpatients from a previous trial.

53 **Methods:** This is a secondary analysis of an investigator-initiated, prospective
54 randomised controlled multicenter trial in Switzerland (EFFORT) that compared the
55 effects of an individualised nutritional support intervention with standard of care. We
56 investigated associations between admission NRS and several short-term and long-
57 term outcomes using multivariable regression analyses.

58 **Results:** Of the 2,028 patients, 31% had an NRS of 3, 38% of 4 and 31% of ≥ 5
59 points, and 477 (24%) died during the 180 days of follow-up. For each point increase
60 in NRS, we found a stepwise increase in risk of 30-day mortality (adjusted Hazard
61 Ratio (HR) 1.22 (95% CI 1.00 to 1.48), $p=0.048$) and 180-day mortality (adjusted HR
62 1.37 (95% CI 1.22 to 1.55), $p<0.001$). NRS was associated with length of hospital
63 stay (adjusted difference of 0.60 days per NRS point increase, 95%CI 0.23 to 0.97,
64 $p=0.002$) and functional outcomes at 180 days (adjusted decrease in Barthel index of
65 -4.49 points per NRS point increase, 95%CI -6.54 to -2.45, $p<0.001$). In a subgroup
66 analysis, associations of NRS and short-term adverse outcomes were less
67 pronounced in patients receiving nutritional support (intervention group) compared to
68 control group patients (adjusted HR for 30-day mortality 1.12 [95%CI 0.83 to 1.52,
69 $p=0.454$] vs. 1.33 [95%CI 1.02 to 1.72, $p=0.032$]).

70 **Conclusion:** The NRS is a strong and independent risk score for malnutrition-
71 associated mortality and adverse outcomes over 180 days. Our data provide strong
72 evidence that the nutritional risk, however, is modifiable and can be reduced by the
73 provision of adequate nutritional support.

74 **Introduction**

75 Malnutrition is a common condition in medical inpatients affecting approximately 30-
76 50% in the western patient population [1-3]. Patients with poor nutritional status are
77 more likely to suffer from adverse outcomes, have an elevated risk of mortality and
78 morbidity, as well as experience significant socioeconomic implications [4-7].
79 Importantly, recent studies have found that malnutrition risk factors in medical
80 inpatient populations are at least partly modifiable [8-10]. More specifically, two trials
81 reported positive outcomes on mortality associated with a nutritional intervention [11,
82 12]. The placebo-controlled NOURISH (Nutrition effect On Unplanned Readmissions
83 and Survival in Hospitalized patients) trial found a significant reduction in mortality
84 over 90 days in medical inpatients treated with a high protein oral nutrition
85 supplement [13]. Similarly, the recent EFFORT (*Effect of Early Nutritional Support on*
86 *Frailty, Functional Outcomes and Recovery of Malnourished Medical Inpatients*) trial
87 found a reduction in the risk for severe complications and mortality associated with
88 the use of nutritional support compared to a control group not receiving additional
89 nutritional support [11]. These findings have provided conclusive evidence to support
90 current guideline recommendations regarding early screening of patients for
91 malnutrition upon hospital admission and the use of nutritional support intervention
92 for at-risk patients [14-16].

93 For this purpose, several screening tools for malnutrition have been proposed and
94 validated in different patient populations [17, 18]. Of these, the Nutritional Risk
95 Screening (NRS 2002) has become particularly well established for the medical
96 inpatient population [19, 20]. NRS includes assessment of the patient's nutritional
97 status (based on weight loss, Body Mass Index (BMI) and general condition or food
98 intake) and disease severity (stress metabolism due to the degree of disease), and is
99 associated with higher risk for adverse outcomes. Each section is scored from 0 to 3

100 points, and patients receive an extra point if they are 70 years or older [21-23].
101 Earlier observational retrospective studies also found that the NRS has prognostic
102 implications and is associated with short-term and long-term mortality [24, 25]. It
103 remains unclear, however, if the association can be explained by other disease-
104 related factors, or whether the type of nutritional support may influence the
105 connection between NRS and outcome.
106 Herein, we hypothesized that an elevated risk for malnutrition, as assessed by the
107 NRS, is associated with an increased long-term risk for mortality and that this risk is
108 modifiable through the provision of individual nutritional support. To test this
109 hypothesis, we performed a secondary analysis of a prospective, multicentre,
110 randomised trial [11] to investigate the association of NRS with different clinical
111 health outcomes at short-term and long-term follow-up, and studied the differences
112 according to the nutritional support provided to patients.

113 **Methods**

114 ***Study design and setting***

115 This study is a secondary analysis of the overall EFFORT study population, an
116 investigator-initiated, non-commercial, prospective and open-label randomised trial
117 that compared the effects of individualised nutritional support intervention versus no
118 nutritional support on medical outcomes in patients at nutritional risk (as assessed by
119 the NRS). The trial protocol and the main results have been published elsewhere [26].
120 The ethics committee of northwest / central Switzerland (EKNZ) approved the study
121 protocol in January 2014 (EKNZ; 2014_001). The eight participating sites were
122 secondary and tertiary care hospitals in Switzerland and included the University
123 Clinic in Aarau, the University Hospital in Bern, the Cantonal hospitals in Lucerne,
124 Solothurn, St. Gallen, Muensterlingen and Baselland, and the hospital in Lachen.
125 Patients were enrolled between April 2014 and February 2018.

127 ***Patient population***

128 Adult patients with a NRS total score ≥ 3 points, an expected length of hospital stay
129 (LOS) > 4 days and willingness to provide informed consent were eligible. Exclusion
130 criteria were defined as initial admission to an intensive care unit or surgical unit; the
131 inability to tolerate oral nutrition intake; nutritional support received at time of
132 admission; patients with a terminal condition; admission to hospital due to anorexia
133 nervosa, acute pancreatitis, acute liver failure, cystic fibrosis, stem cell
134 transplantation or gastric bypass surgery; contraindications for nutritional support;
135 and previous inclusion in the trial.

136

137

138

139 Outcomes

140 The primary endpoint of this study was all-cause mortality from inclusion in the trial
141 up to day 30 and day 180.

142 Secondary endpoints included the composite endpoint adverse events (all-cause
143 mortality, admission to the intensive care , readmission and major complications) as
144 well as major complications (nosocomial infection or abscess requiring antibiotic
145 treatment, major cardiovascular events, acute renal failure); economic outcome
146 including total LOS, non-elective hospital readmission (defined as non-scheduled
147 hospital readmission after discharge), and admission to the intensive care unit from
148 the medical ward. Functional outcomes included functional impairment (assessed
149 with the Barthel scale), quality of life (European Quality of Life 5 Dimensions Index
150 (assessed with the European Quality of Life 5 Dimensions Index (EQ-5D)) and
151 visual-analogue scale [EQ-5D VAS]), fractures, and accidental fall events. All
152 outcomes were defined and assessed as short-term (30 days) and long-term (180
153 days) outcomes. To assess primary and secondary endpoints, all patients were
154 contacted by blinded study nurses for a structured telephone interview after 30 days
155 and 180 days. The survival status of all patients during follow-up was confirmed
156 either by family members or the patient`s family physician.

157 The Barthel scale was used to assess the performance of activities of daily living.

158 Functional impairment was defined as a decline of 10% or more in functional status.

159 The EuroQol Group 5- Dimension Self-Report Questionnaire, which ranges from 0 to
160 1, with higher scores indicating better life quality and EQ-5D VAS, which scores from
161 0 to 100, with higher scores indicating better health status, were used to rate quality
162 of life [11].

163 Nutritional status and procedures

164 Nutritional status was assessed as recommended by nursing staff within 24-48 hours
165 after hospital admission using the NRS score[18, 27]. We scored for each predictor
166 of the NRS (i.e. patient's nutritional status (based on weight loss, Body Mass Index
167 [BMI] and general condition or food intake) and disease severity) between 0 to 3
168 points, and added an extra point for patients aged 70 years or older. A NRS total
169 score of ≥ 3 points was considered "at risk" for malnutrition. We then divided the
170 study population into three groups (i.e., moderate risk, high risk, very high risk)
171 according to NRS (3 points; 4 points; ≥ 5 points).

172

173 ***Nutritional support provided during the trial***

174 Nutritional support during the trial differed according to randomisation of patients, and
175 details of the intervention have been published [26]. In summary, in the intervention
176 group, nutritional support was initiated as soon as possible after trial inclusion.
177 Patients received individualised nutritional support to reach protein and energy
178 requirements according to a previously published consensus protocol and under the
179 guidance of a registered dietician [15]. Energy requirements were predicted using the
180 weight-adjusted Harris-Benedict equation [28]. Daily protein intake was set at 1.2–1.5
181 g/kg body weight, [29] with lower targets for patients with acute renal failure but
182 without need of renal replacement therapy (0.8 g per kg of body weight). An
183 individual nutritional plan was developed for each patient that was initially based on
184 oral nutrition provided by the hospital kitchen and further increased to enteral tube
185 feeding or parenteral feeding if at least 75% of energy and protein targets could not
186 be reached within 5 days by oral (or enteral) feeding. In total 8, respectively 12
187 patients received enteral or parenteral nutrition. Nutritional intake was reassessed
188 every 24–48 h throughout the hospital stay and compliance to the nutrition care plan
189 was reinforced. Upon discharge from hospital, patients received dietary counselling

190 and, if indicated, a prescription for oral nutritional supplements to be taken in the
191 outpatient setting.

192 Control group patients received standard hospital food according to their ability and
193 desire to eat, with no additional nutritional consultation and no recommendation for
194 supplementary nutritional support.

195

196 ***Study aims***

197 The overall aim of this analysis was to investigate the prognostic implications of NRS
198 in connection with short-term and long-term clinical outcomes in a well-characterised
199 cohort of patients from the EFFORT intervention trial, as well as to compare
200 differences when stratifying patients based on nutritional support received.

201

202 ***Sample size and statistical analyses***

203 For this secondary analysis looking at associations of NRS and long-term mortality
204 within 180 days, we used patients previously included in a randomized trial and the
205 sample size was therefore based on the available number of patients included in the
206 initial trial. Still, with 477 patients reaching the primary endpoint, this sample provides
207 adequate power to support over 47 degrees of freedom in the models. We thus
208 assume that inclusion of up to 47 covariates is possible in the regression models.
209 Categorical variables are expressed as counts (percentages, standard deviations
210 (SD)) and continuous variables as medians (interquartile ranges [IQR], 25th and 75th
211 percentiles).

212 We calculated regression models adjusted for important confounders (sex,
213 comorbidities, admission diagnosis, study centre and randomisation) to explore the
214 association between the NRS and several short-term and long-term outcomes.

215 Models were not additionally adjusted for age as this variable is already a part of

216 NRS. We used Cox regression models for time-to-event data with recorded hazard
217 ratios (HRs), logistic regression for binary outcomes with recorded odds ratios (ORs)
218 and linear regression for continuous outcomes with recorded coefficients. We also
219 calculated Kaplan-Meier survival curves to present the results visually.
220 Finally, we conducted different analyses according to the pre-specified subgroups,
221 stratifying patients based on age, sex, and main admission diagnosis, as well as
222 those receiving individual nutrition support for different short-term outcomes.
223 All statistical analyses were performed with STATA 15.1 (Stata Corp, College
224 Station, TX, USA). A *P* value <0.05 (for a 2-sided test) was considered to indicate
225 statistical significance.
226

227 **Results**

228 We included all 2,028 patients who were enrolled in the EFFORT trial. A total of 624
229 (31%) patients had a NRS score of 3 points, 775 (38%) a NRS score of 4 points and
230 629 (31%) a NRS score of ≥ 5 points. Overall, the median age of the patients was
231 72.6 years and 1,064 (52%) were male. When comparing patients with NRS of 3, 4
232 and ≥ 5 points, we found significant differences in regard to age, weight, admission
233 diagnosis, and comorbidities. More detailed patient baseline characteristics, stratified
234 by NRS and by mortality at 180 days, are shown in **Table 1**.

235

236 ***Association of NRS with short-term and long-term mortality (primary endpoint)***

237 At 30-day and 180-day follow-up, a total of 173 patients (9%) and 477 patients (24%)
238 respectively had died. Mortality showed a stepwise increase consistent with higher
239 NRS scores at short term and long term follow-up. This was also confirmed in a
240 multivariable regression analysis with an adjusted HR of 1.22 (1.00 to 1.48, $p=0.048$)
241 for mortality at 30 days and an adjusted HR of 1.37 (1.22 to 1.55, $p<0.001$) for 180-
242 day mortality (**Table 2**).

243 These results were also confirmed in Kaplan-Meier survival estimates showing a
244 higher likelihood for mortality with increasing NRS scores (**Figure 1**).

245

246 ***Associations of NRS with secondary endpoints***

247 We also investigated associations between NRS and different secondary endpoints
248 (**Table 2**). We observed a stepwise increase in the incidence of adverse outcomes
249 within 30 days - from 22.6% (3 points) to 24.0% (4 points) to 28.1% (5 points and
250 more) with an unadjusted OR of 1.16 (95% CI 1.02 to 1.32, $p=0.023$) but without
251 remaining significant after multivariate adjustment ($p=0.130$). There was also a
252 significant increase in mean LOS (from 8.8 to 9.8 to 9.9 days, respectively) with an

253 (adjusted) increase of 0.6 days (95% CI 0.23 to 0.97) $p=0.002$) per increase in NRS
254 point. In addition, there was an increase in the risk for impairment of activities of daily
255 living as defined by Barthel scale at days 30 and 180 (coefficient of -0.65 points (95%
256 CI -1.18 to -0.11, $p=0.018$) for day 30 and -7.52 points (95% CI -9.63 to -5.39,
257 $p<0.001$) for day 180. Similar results were found for impairment in quality of life within
258 180 days, as measured by EQ-5D and the EQ-5D VAS.

259

260 ***Subgroup analysis for the primary endpoint***

261 We also performed several pre-planned subgroup analyses to investigate whether
262 the association between NRS and mortality was dependent on age, sex and main
263 admission diagnosis. **Figure 2** shows associations of the NRS and 180-day mortality
264 within these different subgroups. Overall, results were similar, with little difference
265 between groups.

266

267 ***Subgroup analysis regarding effects of nutritional support***

268 Finally, to understand whether the nutritional risk is modifiable through the provision
269 of nutritional support, we performed a subgroup analysis comparing associations of
270 NRS and outcomes stratified by nutritional support received during the trial
271 (nutritional support group vs. control group) (**Table 3, Figure 3**). We found a stronger
272 association of NRS and mortality within 30 days for patients not receiving nutritional
273 support (i.e. control group patients) compared to patients receiving nutritional support
274 (HR of 1.43 (95% CI 1.11 to 1.85) vs. 1.20 (95% CI 0.89 to 1.61). Results were
275 similar for other endpoints including overall adverse outcomes, non-elective hospital
276 readmission, and admission to an intensive care unit.

277

278

279 **Discussion**

280 The main findings of this secondary analysis from a recent multicentre trial are
281 twofold. First, we found associations of NRS with different adverse clinical outcomes
282 at short-term and long-term follow-up, which proved to be independent of important
283 confounders in multivariate analysis and showed robust results in different subgroup
284 analyses. This demonstrates that NRS has strong prognostic implications regarding
285 malnutrition-associated adverse clinical outcomes. Secondly, the association
286 between NRS and adverse outcomes were less pronounced in patients receiving
287 nutritional support compared to patients not receiving nutritional support, suggesting
288 that the risk for adverse outcomes for patients with malnutrition is at least partly
289 modifiable through provision of nutritional support.

290

291 There are several findings of this study worth mentioning. Firstly, the association
292 between malnutrition and mortality has been known for some time [1, 30, 31]. A
293 previous retrospective observational study performed in Italy, including 5,698 patients
294 hospitalized between from October 2015 and July 2016, showed that nutritional risk
295 identified by NRS at time of hospital admission was a good predictor of short-term (1-,
296 3-, 6-month) and long-term (1 year) mortality, with a doubling in mortality comparing
297 patients scoring $NRS \leq 3$ with those $NRS \geq 3$ [24]. These findings are in line with our
298 results, which also show an increase of 5% within 30 days and 17% within 180 days
299 between patients with an NRS of 3 and those with ≥ 5 points. Importantly, we were
300 also able to adjust our analysis for important confounders such as socio-
301 demographic factors, main admission diagnosis and comorbidities, suggesting that
302 malnutrition has an independent negative effect on health outcomes, which is not
303 explained by the heavier burden of disease seen in the malnourished population. Our
304 prospective sample of patients with detailed clinical information thereby confirms

305 results of other observational and retrospective studies with less rigorous statistical
306 adjustment[24].

307

308 Secondly, our findings regarding secondary endpoints are also partly in line with
309 multiple previous studies, which report associations between nutritional risk and
310 various economic outcomes such as increased LOS [32-38], hospital readmission [4,
311 39] and admission to an intensive care unit. The economic burden of malnutrition
312 derives mostly from extended LOS, which leads to higher use of hospital resources
313 and thus increased costs. A prospective cohort study of 818 patients in Singapore
314 found an increased LOS by two days when comparing well nourished with
315 malnourished and severely malnourished patients (using the Subjective Global
316 Assessment SGA) [39]. In our study, we were able to adjust all analysis for
317 confounders showing that NRS might be indeed independently associated with these
318 economic outcomes.

319

320 Thirdly, we were able to look at the association of malnutrition risk as assessed by
321 NRS within different subgroups with different underlying main diagnosis- asking the
322 question whether the individual situation of a patient with regard to socio-
323 demographics, admission diagnosis, and comorbidities may influence the strength of
324 association.[40] Overall, we found little variation within these groups, suggesting that
325 malnutrition is a risk factor across the entire medical inpatient population and the
326 consequence of different illnesses, rather than caused by specific conditions.[41]
327 Screening and treatment of malnutrition should, therefore, not be limited to certain
328 patient populations, but rather include all medical inpatients.[42] This is also in line
329 with the EFFORT trial, which demonstrates the benefits of nutritional support
330 independent of the medical condition.[11]

331

332 Fourthly, most studies looking at malnutrition and risk of impaired functional
333 outcomes (such as quality of life or performance of activities of daily living) were
334 carried out on a geriatric population [43, 44]. Functional impairments have an
335 important impact on a patient's independence, with dramatic socio-economic
336 implications [43]. Our analysis expands the results regarding functional outcomes to
337 a medical inpatient population, demonstrating similar results to those known from
338 geriatrics. Both quality of life and performance of daily activities measured by the EQ-
339 5D and the Barthel scale decreased with an increasing NRS score. Interestingly,
340 these associations were more pronounced for long-term outcomes and remained
341 significant in the fully adjusted statistical model. The Barthel scale, for instance, was
342 42% higher in patients scoring ≥ 5 points in the NRS than patients scoring 3 points.
343 Naturally the worsening of functional outcomes due to progression of sarcopenia
344 takes time to develop and the consequences of malnutrition only become evident
345 only after a certain period of time.

346

347 Fifth, as a new and clinically relevant main finding, we explored whether provision of
348 nutritional support influences the association between malnutrition and adverse
349 clinical outcomes. We focused on short-term outcomes because our intervention only
350 looked at the initial hospital stay and not the post-discharge period. Interestingly, the
351 association between NRS and mortality was only about half as strong in the
352 intervention group as compared to the control group. This indicates that the adverse
353 effects of malnutrition are at least partially modifiable. These findings again suggest
354 that patients as being identified as at risk of malnutrition according to NRS or a
355 similarly well-validated nutrition screening tool should receive more in-depth
356 assessment and individualised nutritional support, if indicated.

357 We used the NRS as a screening tool, as recommended by the European Society of
358 Parenteral and Enteral Nutrition (ESPEN)[18]. Other screening tools for malnutrition
359 such as the Mini Nutritional Assessment (MNA) and its shorter form (MNA-SF), as
360 well as the malnutrition universal screening tool (MUST) have been validated for
361 predicting mortality and adverse outcomes in previous studies, but it remains unclear
362 which of these tools best identifies patients who would benefit from nutritional
363 intervention [22, 23, 45].

364

365 This trial has several strengths and limitations worth mentioning. One of the strengths
366 of this study is that it consists on a secondary analysis of a prospective randomised
367 trial including a large unselected and heterogeneous population [12, 46, 47]. To the
368 best of our best knowledge this is the first adequately powered study to investigate
369 several short-term and long-term outcomes, and include functional outcomes.
370 Furthermore, while several observational studies investigated the predictive validity
371 of the NRS, we were the first to demonstrate that nutritional support has an influence
372 on the association of NRS and outcome and is thus an effect modifier. We were also
373 able to calculate multivariate regression models and adjust the analysis for important
374 confounders.

375

376 There are, however, some limitations to the underlying EFFORT trial; including the
377 non-blinding of patients and dieticians, some variation in compliance with the
378 nutritional protocol (with about 20% of patients not reaching their energy and protein
379 goals which, however, is a conservative bias towards the here relevant endpoints),
380 and the focus on one country which may limit external validity to other health care
381 systems. Also, we only included patients with an NRS score of at least 3 points and
382 thus have no data regarding patients with no nutritional risk as a control group. We

383 also did not include ICU patients and surgical patients and our findings thus only
384 applies to medical inpatients limiting external validity. Lastly the selection of co-
385 morbidities for inclusion in statistical models was based on the data collection within
386 the initial trial.

387

388 In conclusion, as it mirrors patients' individual nutritional risk, the NRS is a strong and
389 independent risk factor for mortality and adverse outcomes - which may in turn be
390 modified by the adequate provision of nutritional support.

391

392

393

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396

397 **Statement of Authorship**

398 LH, AB, LB and PS were responsible for the data analysis and interpretation of this
399 secondary analysis. LH, AB, LB and PS drafted the final manuscript with all authors
400 contributing to critical revision of the manuscript. PS was responsible for obtaining
401 funding. RF, VB, MG, MD, PT, NK, SS, CB, SM, CB were involved in data collection
402 and approved the final version of the manuscript.

403 FG, AK, TB, CH, VP, SB, SS, MB, CH, RT, JR, DA, NR, JD were involved in drafting
404 the trial protocol, supervision of study sites, drafting of the final manuscript and
405 approved the final version of the manuscript of the original EFFORT trial.

406 ZS and BM were involved in obtaining funding, drafting the trial protocol, supervision
407 of study sites, drafting of the final manuscript of the original EFFORT trial and
408 approved the final version of the current manuscript. The corresponding authors had
409 full access to all the data used and had a shared final responsibility for the accuracy
410 of the analysed data.

411

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Tables and Figure Legends

Table 1. Baseline Characteristics

Table 2. Association of NRS and clinical outcomes

Table 3: Association of NRS with short-term Outcomes, stratified by nutritional support (intervention vs control group).

Figure 1. Kaplan Meier estimate on 180-day mortality stratified by the NRS

Time to death shown for each NRS score upon admission ($p \leq 0.001$)

Figure 2. Subgroup analysis

Subgroup analysis for sociodemographic factors and main diagnosis. The overall effect is listed as the reference group (HR 1.31; CI 95% 1.17,1.48). "Other diagnosis" includes neuropsychological, renal, gastrointestinal and metabolic illnesses.

Figure 3. Subgroup analysis regarding mortality and non-elective readmission

Association of NRS and endpoints stratified by nutritional support (intervention vs control group). Adjusted Hazard ratios are shown for time to event outcome data, odds ratios for binary outcome data and coefficients for continuous outcomes.

Appendix

Figure 4. Subgroup analysis regarding adverse outcomes, major complications and decline in functional status

Effects of nutritional support on primary endpoints for patients compared to the control group. Odds ratios for binary outcome data and coefficients for continuous outcomes.

Figure 5. Subgroup analysis regarding Length of stay and Barthel index

Effects of nutritional support on primary endpoints for patients compared to the control group. Coefficients are shown for continuous outcomes.

Table 2. Association of NRS and clinical outcomes

	NRS 3 (N=624)	NRS 4 (N=775)	NRS ≥5 (N=629)	p-Value	Hazard ratio (HR), Odds ratio (OR), Coefficients	Regression analysis (not adjusted) (95%CI and p-value)	Regression analysis (adjusted) (95%CI and p-value)
Primary outcomes							
Short-term outcomes							
All-cause mortality within 30 days	41 (6.6%)	62 (8.0%)	70 (11.1%)	0.012	HR	1.33 (1.09 to 1.61) p=0.004	1.22 (1.00 to 1.48) p=0.048
Long-term outcomes							
All-cause mortality within 180 days	101 (16.2%)	169 (21.8%)	207 (32.9%)	<0.001	HR	1.51 (1.34 to 1.70) p<0.001	1.37 (1.22 to 1.55) p<0.001
Secondary outcomes							
Short-term outcomes							
Complications							
Adverse outcome within 30 days	141 (22.6%)	186 (24.0%)	177 (28.1%)	0.06	OR	1.16 (1.02 to 1.32) p=0.023	1.11 (0.97 to 1.27) p=0.130
Non-elective hospital readmission within 30 days	55 (8.8%)	64 (8.3%)	61 (9.7%)	0.64	HR	1.05 (0.87 to 1.27) p=0.589	1.03 (0.85 to 1.25) p=0.759
Admission to the intensive care unit within 30 days	13 (2.1%)	24 (3.1%)	12 (1.9%)	0.29	OR	0.96 (0.67 to 1.38) p=0.837	1.08 (0.74 to 1.57) p=0.696
Any major complication	45 (7.2%)	62 (8.0%)	43 (6.8%)	0.69	OR	0.97 (0.79 to 1.20) p=0.798	0.97 (0.78 to 1.21) p=0.804
Nosocomial infection	17 (2.7%)	34 (4.4%)	28 (4.5%)	0.19	OR	1.26 (0.94 to 1.68) p=0.116	1.22 (0.91 to 1.65) p=0.182
Major cardiovascular event	4 (0.6%)	4 (0.5%)	7 (1.1%)	0.41	OR	1.39 (0.72 to 2.69) p=0.332	1.35 (0.68 to 2.67) p=0.386
Acute kidney failure	20 (3.2%)	25 (3.2%)	18 (2.9%)	0.91	OR	0.94 (0.69 to 1.30) p=0.726	0.91 (0.66 to 1.27) p=0.592
Functional outcome							
Mean length of stay within 30 days (days)	8.8 (6.1)	9.8 (6.7)	9.9 (6.8)	0.005	Coefficient	0.54 (0.18 to 0.90) p=0.003	0.6 (0.23 to 0.97) p=0.002
Mean BARTHEL score (points) within 30 days	95.58 (9.12)	95.21 (9.42)	94.29 (10.6)	0.052	Coefficient	-0.65 (-1.18 to -0.11) p=0.018	-0.53 (-1.07 to 0.02) p=0.059
Decline in functional status of >10%	64 (10.3%)	90 (11.6%)	92 (14.6%)	0.052	OR	1.23 (1.04 to 1.46) p=0.018	1.16 (0.97 to 1.38) p=0.105
Long-term outcomes							
Complications							
Non-elective hospital readmission within 180 days	168 (26.9%)	204 (26.3%)	177 (28.1%)	0.74	HR	1.11 (1.00 to 1.24) p=0.051	1.07 (0.96 to 1.19) p=0.248
Accidental fall event within 180 days	74 (11.9%)	88 (11.4%)	58 (9.2%)	0.27	OR	0.87 (0.73 to 1.04) p=0.133	0.89 (0.74 to 1.07) p=0.200
Fracture within 180 days	8 (1.3%)	17 (2.2%)	7 (1.1%)	0.2	OR	0.94 (0.60 to 1.47) p=0.79	0.93 (0.58 to 1.48) p=0.749
Functional outcomes							
Mean EQ-5D index (points)†	0.77 (0.30)	0.75 (0.33)	0.69 (0.35)	<0.001	Coefficient	-0.04 (-0.06 to -0.02) p<0.001	-0.03 (-0.05 to -0.01) p=0.015
VAS index †	60 (26)	58 (27)	55 (29)	0.007	Coefficient	-2.57 (-4.23 to -0.91) p=0.002	-1.59 (-3.23 to 0.05) p=0.058
Mean EQ-5D VAS (points) within 180 days †	56.5 (32.5)	51.5 (34.6)	44.5 (37.3)	<0.001	Coefficient	-6.02 (-8.07 to -3.96) p<0.001	-4.22 (-6.16 to -2.28) p<0.001
Mean BARTHEL score (points) within 180 days †	73.1 (34.27)	68.34 (37.74)	58.08 (42.44)	<0.001	Coefficient	-7.51 (-9.63 to -5.39) p<0.001	-4.49 (-6.54 to -2.45) p<0.001
Decline in mean BARTHEL score (points) within 180 days	284 (47.3%)	369 (49.9%)	317 (53.2%)	0.12	OR	1.13 (1.01 to 1.26) p=0.040	1.12 (0.99 to 1.27) p=0.064
Continuous values as median and IQR, categorical/binary values as absolute number and percentage. NRS= Nutritional Risk Screening, EQ-5D= European Quality of Life 5 Dimensions index; VAS= visual-analogue scale Adjusted for sex, admission diagnosis, comorbidities, study centre and randomization. Comorbidities include: Coronary heart disease, chronic heart failure, hypertonia, stroke, chronic renal failure, diabetes mellitus, tumor, chronic obstructive pulmonary disease, peripheral artery disease and dementia HR= Hazard ratio; OR= Odds ratio							

Table 3: Short-term Outcomes in control versus intervention group

	<i>Hazard ratio (HR), Odds ratio (OR), Coefficients</i>	Regression analysis Control (non-adjusted) (odds ratio and 95%CI and p-value)	Regression analysis Intervention (non-adjusted) (odds ratio and 95%CI and p-value)	Regression analysis Control (adjusted) (odds ratio and 95%CI and p-value)	Regression analysis Intervention (adjusted) (odds ratio and 95%CI and p-value)
Primary outcomes					
All-cause mortality within 30 days	HR	1.43 (1.11 to 1.85) p=0.006	1.20 (0.89 to 1.61) p=0.232	1.33 (1.02 to 1.72) p=0.032	1.12 (0.83 to 1.52) p=0.454
Secondary outcomes					
Complications					
Adverse outcome within 30 days	OR	1.22 (1.02 to 1.46) p=0.026	1.10 (0.91 to 1.32) p=0.336	1.18 (0.98 to 1.42) p=0.087	1.05 (0.86 to 1.28) p=0.630
Any major complication	OR	0.98 (0.73 to 1.31) p=0.871	0.97 (0.72 to 1.31) p=0.842	0.95 (0.70 to 1.3) p=0.750	0.98 (0.72 to 1.34) p=0.914
Acute kidney failure	OR	1.17 (0.74 to 1.85) p=0.493	0.76 (0.48 to 1.20) p=0.243	1.07 (0.67 to 1.73) p=0.771	0.75 (0.47 to 1.19) p=0.223
Economic outcome					
Mean length of stay within 30 days (days)	Coefficient	0.37 (-0.11 to 0.84) p=0.132	0.72 (0.17 to 1.26) p=0.010	0.42 (-0.07 to 0.91) p=0.092	0.80 (0.24 to 1.36) p=0.005
Non-elective hospital readmission within 30 days	HR	1.14 (0.88 to 1.48) p=0.327	0.97 (0.74 to 1.26) p=0.822	1.18 (0.90 to 1.55) p=0.227	0.90 (0.69 to 1.18) p=0.436
Admission to the intensive care unit within 30 days	OR	1.00 (0.61 to 1.64) p=0.995	0.92 (0.55 to 1.57) p=0.769	1.05 (0.63 to 1.77) p=0.840	1.09 (0.62 to 1.92) p=0.753
Functional outcome					
Mean BARTHEL score (points) within 30 days	Coefficient	-0.33 (-1.1 to 0.44) p=0.395	-0.96 (-1.72 to -0.21) p=0.012	-0.15 (-0.94 to 0.64) p=0.702	-0.87 (-1.63 to -0.10) p=0.026
Decline in functional status of >10%	OR	1.17 (0.93 to 1.46) p=0.175	1.32 (1.01 to 1.72) p=0.040	1.08 (0.85 to 1.37) p=0.546	1.28 (0.97 to 1.69) p=0.078
Continuous values as median and IQR, categorical / binary values as absolute number and percentage. NRS= Nutritional Risk Screening, EQ-5D= Euroquo-5 Dimensions, VAS= Visual Analogue Scale *Adjusted for sex, admission diagnosis, comorbidities, study centre and randomization					

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Table 1. Baseline results of patients

Parameters	Stratified according to NRS				Stratified according to Mortality		
	NRS 3	NRS 4	NRS ≥5	p-Value	Survivors	Non-survivors	p-Value
N	624	775	629		1551	477	
Sociodemographics							
Mean age (years)	70.20 (15.2)	71.4 (14.9)	76.5 (10.6)	<0.001	71.5	76.2	<0.001
Age group							
<65 years	143 (22.9%)	162 (20.9%)	50 (7.9%)	<0.001	307 (19.8%)	48 (10.1%)	<0.001
65-75 years	215 (34.5%)	247 (31.9%)	209 (33.2%)	<0.001	517 (33.3%)	154 (32.3%)	<0.001
>75 years	266 (42.6%)	366 (47.2%)	370 (58.8%)	<0.001	727 (46.9%)	275 (57.7%)	<0.001
Male sex [no.] (%)	344 (55.1%)	402 (51.9%)	318 (50.6%)	0.250	773 (49.8%)	291 (61.0%)	<0.001
Nutritional assessment							
Mean Body Mass Index (kg/m ²)	26.0 (5.0)	24.8 (5.3)	23.6 (5.4)	<0.001	25 (5.5)	24.2 (4.6)	0.004
Mean bodyweight (kg)	74.2 (16.6)	71.1 (16.5)	67.1 (16.3)	<0.001	71.2 (17.0)	69.9 (15.5)	0.220
NRS 2002 score (%)							
3 points					523 (33.7%)	101 (21.2%)	<0.001
4 points					606 (39.1%)	169 (35.4%)	<0.001
5 points					357 (23.0%)	167 (35.0%)	<0.001
>5 points					65 (4.2%)	40 (8.4%)	<0.001
Weight loss - no. (%)							
≤5% in 3 month	434 (69.6%)	394 (50.8%)	242 (38.5%)	<0.001	858 (55.3%)	212 (44.4%)	<0.001
>5% in 3 month	94 (15.1%)	115 (14.8%)	76 (12.1%)	<0.001	200 (12.9%)	85 (17.8%)	<0.001
>5% in 2 month	70 (11.2%)	132 (17.0%)	55 (8.7%)	<0.001	182 (11.7%)	75 (15.7%)	<0.001
>5% in 1 month	26 (4.2%)	134 (17.3%)	256 (40.7%)	<0.001	311 (20.1%)	105 (22.0%)	<0.001
Loss of appetite - no. (%)							
No	99 (15.9%)	74 (9.5%)	56 (8.9%)	<0.001	200 (12.9%)	29 (6.1%)	<0.001
Yes	525 (84.1%)	701 (90.5%)	573 (91.1%)	<0.001	1351 (87.1%)	448 (93.9%)	<0.001
Normal required food intake preceding week - no. (%)							
>75%	89 (14.3%)	69 (8.9%)	47 (7.5%)	<0.001	181 (11.7%)	24 (5.0%)	<0.001
50-75%	336 (53.8%)	202 (26.1%)	101 (16.1%)	<0.001	501 (32.3%)	138 (28.9%)	<0.001
25-50%	184 (29.5%)	378 (48.8%)	277 (44.0%)	<0.001	614 (39.6%)	225 (47.2%)	<0.001
<25%	15 (2.4%)	126 (16.3%)	204 (32.4%)	<0.001	255 (16.4%)	90 (18.9%)	<0.001
Severity of illness - no. (%)							
Very mild	33 (5.3%)	22 (2.8%)	0 (0.0%)	<0.001	53 (3.4%)	2 (0.4%)	<0.001
Mild	482 (77.2%)	548 (70.7%)	286 (45.5%)	<0.001	1021 (65.8%)	295 (61.8%)	<0.001
Moderate	105 (16.8%)	200 (25.8%)	330 (52.5%)	<0.001	458 (29.5%)	177 (37.1%)	<0.001
Severe	4 (0.6%)	5 (0.6%)	13 (2.1%)	<0.001	19 (1.2%)	3 (0.6%)	<0.001
Admission diagnosis							
Cardiovascular disease	78 (12.5%)	76 (9.8%)	51 (8.1%)	0.034	148 (9.5%)	57 (11.9%)	0.130
Infection	166 (26.6%)	234 (30.2%)	213 (33.9%)	0.02	517 (33.3%)	96 (20.1%)	<0.001

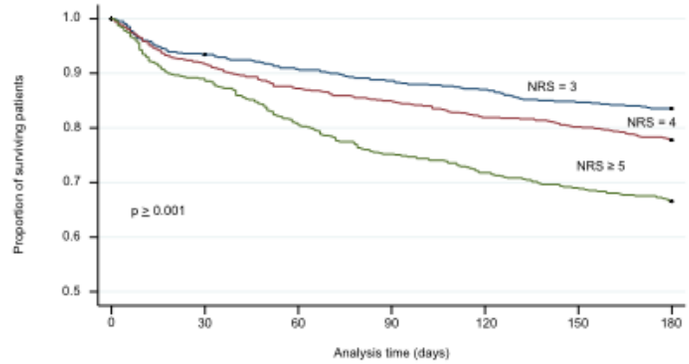
Metabolic disease	20 (3.2%)	28 (3.6%)	14 (2.2%)	0.31	54 (3.5%)	8 (1.7%)	0.045
Gastrointestinal disease	57 (9.1%)	72 (9.3%)	35 (5.6%)	0.02	136 (8.8%)	28 (5.9%)	0.042
Renal disease	13 (2.1%)	29 (3.7%)	26 (4.1%)	0.098	52 (3.4%)	16 (3.4%)	1.000
Cancer	91 (14.6%)	129 (16.6%)	154 (24.5%)	<0.001	188 (12.1%)	186 (39.0%)	<0.001
Lung disease	39 (6.2%)	50 (6.5%)	36 (5.7%)	0.85	98 (6.3%)	27 (5.7%)	0.600
Neurological disease	44 (7.1%)	34 (4.4%)	17 (2.7%)	0.001	89 (5.7%)	6 (1.3%)	<0.001
Reduced general condition	71 (11.4%)	76 (9.8%)	47 (7.5%)	0.061	167 (10.8%)	27 (5.7%)	<0.001
Other	21 (3.4%)	20 (2.6%)	14 (2.2%)	0.44	42 (2.7%)	13 (2.7%)	0.980
Comorbidity							
Coronary heart disease	175 (28.0%)	208 (26.8%)	183 (29.1%)	0.64	423 (27.3%)	143 (30.0%)	0.25
Congestive heart failure	120 (19.2%)	123 (15.9%)	110 (17.5%)	0.26	239 (15.4%)	114 (23.9%)	<0.001
Hypertension	305 (48.9%)	435 (56.1%)	369 (58.7%)	0.001	839 (54.1%)	270 (56.6%)	0.34
Stroke	51 (8.2%)	58 (7.5%)	53 (8.4%)	0.79	121 (7.8%)	41 (8.6%)	0.58
PAD	64 (10.3%)	72 (9.3%)	50 (7.9%)	0.36	137 (8.8%)	49 (10.3%)	0.34
Chronic kidney disease	184 (29.5%)	219 (28.3%)	238 (37.8%)	<0.001	459 (29.6%)	182 (38.2%)	<0.001
Diabetes	124 (19.9%)	171 (22.1%)	133 (21.1%)	0.61	311 (20.1%)	117 (24.5%)	0.036
COPD	89 (14.3%)	115 (14.8%)	99 (15.7%)	0.76	231 (14.9%)	72 (15.1%)	0.91
Dementia	25 (4.0%)	33 (4.3%)	17 (2.7%)	0.27	55 (3.5%)	20 (4.2%)	0.51
Malignant disease	178 (28.5%)	213 (27.5%)	276 (43.9%)	<0.001	388 (25.0%)	279 (58.5%)	<0.001

Continuous values as median and IQR, categorical / binary values as absolute number and percentage.

NRS-2002= Nutritional Risk Screening 2002, PAD= Peripheral Artery Disease, COPD= Chronic Obstructive Pulmonary Disease

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Figure 1. Kaplan Meier estimate on 180 day mortality stratified by NRS



Number at risk

NRS = 3	624	583	551	538	528	514	507
NRS = 4	775	712	656	639	616	603	585
NRS ≥ 5	629	560	497	462	441	424	409

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Figure 2. Subgroup analysis

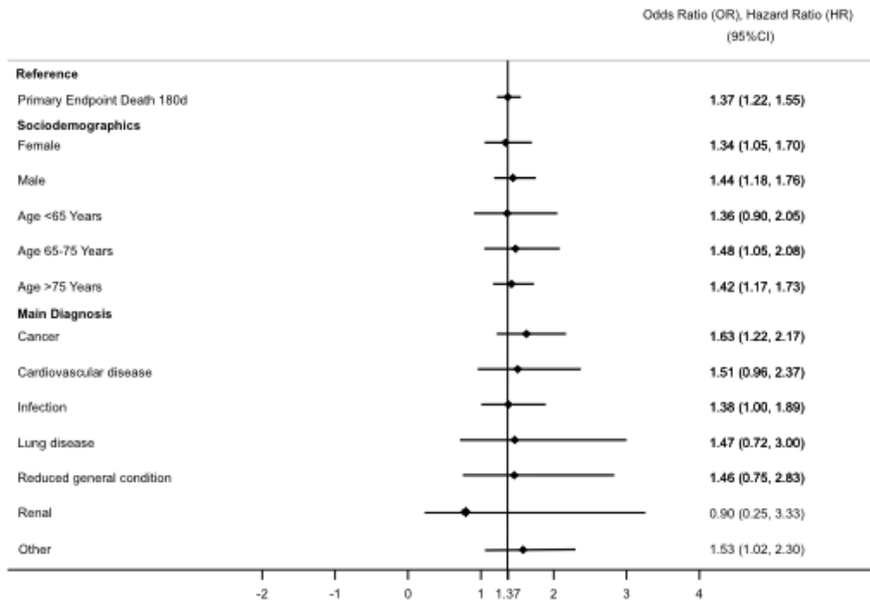
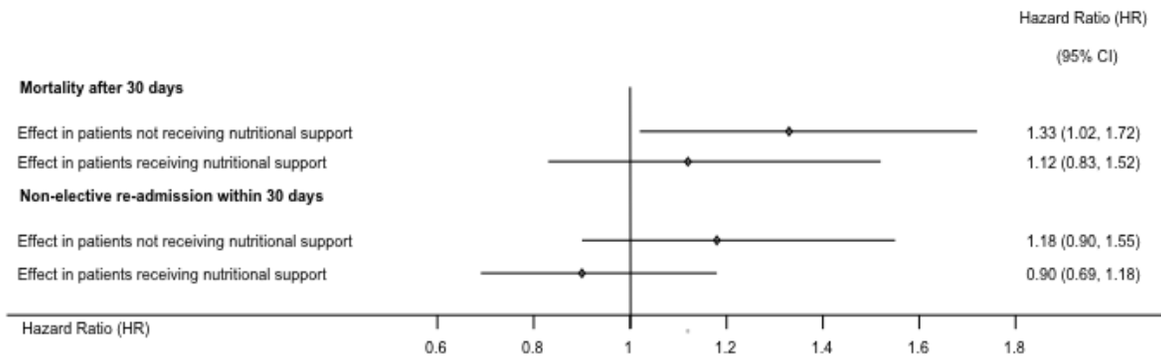


Figure 3. Analysis regarding nutritional intervention in primary endpoints



Conflict of Interest Statement and Funding sources

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