



Intravenous Vitamin C for Patients Hospitalized With COVID-19: Two Harmonized Randomized Clinical Trials

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1 Intravenous vitamin C for patients hospitalized with COVID-19: a prospective harmonization of
2 two randomized clinical trials

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4 behalf of the Canadian Critical Care Trials Group, and for the REMAP-CAP Investigators

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149

150

151

152 **Key Points**

153

154 Question: Does intravenous vitamin C administered to patients hospitalized with COVID-19
155 improve organ-support free days (a composite outcome of in-hospital mortality and duration of
156 intensive care unit–based respiratory or cardiovascular support) up to day 21?

157

158 Findings: In two prospectively harmonized randomized clinical trials, vitamin C, compared to
159 placebo or no vitamin C, yielded posterior probabilities of efficacy of 8.6% among 1568
160 critically ill patients and 2.9% among 1022 non-critically ill patients, regarding the odds of
161 improvement in organ-support free days.

162

163 Meaning: Among hospitalized patients with COVID-19, there was a low probability that vitamin
164 C improved organ-support free days.

165 **Abstract**

166

167 Importance: The efficacy of vitamin C for hospitalized patients with COVID-19 is uncertain.

168 Objective: To determine whether vitamin C improves outcomes for COVID-19 inpatients.

169 Design, Setting, Participants: Two prospectively harmonized randomized clinical trials enrolled
170 critically ill patients receiving organ support in an intensive care unit (ICU, 90 sites), and non-
171 critically ill patients (40 sites), from 23July2020 to 15July2022, in 4 continents.

172 Interventions: Patients were randomized to receive intravenous vitamin C or control (placebo/no
173 vitamin C) for up to 96hr.

174 Main outcomes and measures: The primary outcome was a composite of organ-support free days,
175 defined as days alive and free of ICU-based respiratory and cardiovascular organ support, up to
176 day 21, and survival to hospital discharge. Values ranged from -1 for in-hospital death to 22 for
177 survivors with no organ support. The primary analysis used a Bayesian cumulative logistic
178 model. Odds ratio (OR) >1 represented efficacy (improved survival, more organ-support free
179 days, or both), OR <1 represented harm, and OR <1.2 represented futility.

180 Results: Enrollment was terminated after statistical triggers for harm and futility were met. The
181 trials enrolled 1568 critically ill patients (1041 vitamin C, 537 control; median age 60yr; 35.9%
182 female) and 1022 non-critically ill patients (464 vitamin C; 572 control; median age 62yr, 39.6%
183 female). Among critically ill patients, median organ-support free days (vitamin C vs. control)
184 were 7 (interquartile range [IQR] -1, 17) vs. 10 (IQR -1, 17); OR 0.88, 95% credible interval [CrI]
185 0.73-1.06; posterior probabilities were 8.6% (efficacy), 91.4% (harm), and 99.9% (futility).

186 Among non-critically ill patients, median organ-support free days (vitamin C vs. control) were
187 22 (IQR 18, 22) vs. 22 (IQR 21, 22); OR 0.80, 95%CrI 0.60-1.01; posterior probabilities were

188 2.9% (efficacy), 97.1% (harm), and >99.9% (futility). Survival to hospital discharge (vitamin C
189 vs. control) in the critically ill was 61.9% (642/1037) vs. 64.6% (343/531) [OR 0.92 (95%CrI
190 0.73-1.17)] and in the non-critically ill was 85.1% (388/456) vs. 86.6% (490/566) [OR 0.86
191 (95%CrI 0.61-1.17)], with 24.0% and 17.8% posterior probability of efficacy, respectively.

192 Conclusions and Relevance: In hospitalized patients with COVID-19, vitamin C did not improve
193 organ-support free days or hospital survival.

194

195 Trial Registration: ClinicalTrials.gov identifiers: NCT04401150 (LOVIT-COVID);

196 NCT02735707 (REMAP-CAP).

197 As of September 2023, World Health Organization (WHO) has reported at least 770 million
198 cases and 6.9 million deaths due to coronavirus disease 2019 (COVID-19).¹ For hospitalized
199 patients, immunomodulatory and anti-viral therapies are effective but imperfect,² and global
200 availability remains disparate.³

201

202 Vitamin C is widely available and its use in septic shock increased pre-pandemic⁴ until clinical
203 trials failed to demonstrate benefit.⁵⁻⁷ At the beginning of the COVID-19 pandemic, a WHO
204 report highlighted it as a potential immunomodulatory agent.⁸ Vitamin C attenuates oxidative
205 stress and microvascular thrombosis,⁹ two features of COVID-19, and hospitalized patients with
206 COVID-19 were found to have low serum vitamin C levels.¹⁰ A meta-analysis in patients with
207 COVID-19 reported that vitamin C may reduce hospital mortality.¹¹

208

209 We harmonized two initially separate randomized clinical trials to investigate the effect of
210 intravenous vitamin C on need for organ support and hospital survival in hospitalized patients
211 with COVID-19, hypothesizing that vitamin C would increase days alive and free of organ
212 support.

213

214 **Methods**

215 Trial design

216 Before recruitment commenced, the investigators harmonized and decided to pool data from two
217 clinical trials designed to evaluate the same vitamin C regimen. The Lessening Organ
218 dysfunction with VITamin C-COVID (LOVIT-COVID) trial was initially designed as a
219 frequentist blinded trial enrolling in Canada. The Randomized, Embedded, Multifactorial

220 Adaptive Platform Trial for Community-Acquired Pneumonia (REMAP-CAP) trial is an
221 international, adaptive unblinded platform trial in patients with severe pneumonia;¹² this report
222 includes patients enrolled in the COVID-19 stratum. Both trials prospectively adopted the same
223 intervention, outcomes, statistical analysis plan, and reporting, but control groups were different:
224 placebo in LOVIT-COVID, and no vitamin C in REMAP-CAP. Development of the harmonized
225 trial and essential details of LOVIT-COVID and REMAP-CAP are in supplement 1 (eMethods
226 and eTable 1); full protocols for both trials are in supplement 2. To account for observed racial
227 and ethnic differences in outcomes during the pandemic, REMAP-CAP collected self-reported
228 race and ethnicity from either participants or their surrogates, according to each region's
229 standards.

230

231 The research ethics committee and regulatory authority in each jurisdiction approved the relevant
232 trial protocol. Informed consent was obtained, either before randomization or afterwards, from
233 all patients or their surrogates, in accordance with applicable legislation. Both trials had separate
234 steering committees (with common co-chairs) and Data and Safety Monitoring Boards (DSMBs).
235 Neither trial incorporated accruing data from the other in interim analyses, but their DSMBs
236 exchanged information regarding respective trial progress.

237

238 Patients

239 Eligible patients were adults admitted to hospital with suspected or proven COVID-19. Patients
240 admitted to an intensive care unit (ICU) and receiving respiratory or cardiovascular organ
241 support at the time of randomization were classified as critically ill and all others as non-
242 critically ill. This prospective classification was undertaken because of previous reports

243 suggesting differential treatment effects in these two populations.¹³⁻¹⁵ Respiratory support was
244 defined by receipt of invasive ventilation, non-invasive ventilation, or high-flow nasal oxygen,
245 and cardiovascular support by a vasopressor or inotrope infusion. In LOVIT-COVID, critically
246 ill patients were enrolled while receiving respiratory support; cardiovascular support was an
247 exclusion criterion. Detailed selection criteria appear in eMethods.

248

249 Randomization, interventions, and follow-up

250 Randomization in both trials was concealed via separate computer-based randomization systems.
251 Patients in LOVIT-COVID were assigned in a 1:1 ratio to vitamin C or placebo. In REMAP-
252 CAP, randomization was stratified by state (critically ill vs. non-critically ill), and patients could
253 participate in other domains (eTable 2). The initial randomization ratio of vitamin C to no
254 vitamin C was 1:1, with patients subsequently assigned preferentially to the arm that appeared
255 more favorable after each adaptive analysis (protocol, supplement 2).

256

257 In both trials, patients in the intervention group received intravenous vitamin C, 50 mg/kg body
258 weight, infused over 30-60 minutes, every 6 hours for 96 hours, up to a maximum of 16 doses.
259 All sites used locally available vitamin C formulations (eMethods). In LOVIT-COVID, glucose
260 monitoring for patients receiving insulin or oral hypoglycemic agents was protocolized to
261 account for interference of vitamin C with bedside glucometers (eMethods). In REMAP-CAP,
262 this protocol was advised for patients randomized to vitamin C. All other aspects of care were at
263 clinicians' discretion. Patients were followed in hospital, with survivors or their relatives (all in
264 LOVIT-COVID, and a subset in REMAP-CAP) telephoned at 6 months for additional outcomes.

265

266 Trial outcomes

267 The primary outcome was a composite of an ordinal measure of organ-support free days, defined
268 as days free of respiratory and cardiovascular organ support delivered in the ICU up to day 21,
269 and survival to hospital discharge. This hospital-based outcome is associated with 180-day
270 survival.¹⁶ Deaths within the hospital were assigned the worst outcome (-1). Among hospital
271 survivors, respiratory and cardiovascular organ support-free days were calculated up to day 21; a
272 higher number represents faster recovery. Survival to hospital discharge was censored at 90 days.
273 Non-critically ill patients who survived without needing any organ support were assigned the
274 best outcome (22 organ support-free days).

275

276 Secondary outcomes were pre-specified in the statistical analysis plan (supplement 2) and
277 included death or persistent organ dysfunction¹⁷ (receipt of invasive ventilation, a vasopressor
278 infusion, or new kidney replacement therapy) at trial day 28, which was the primary outcome in
279 the LOVIT trial of vitamin C in sepsis.⁷

280

281 Site investigators reported serious adverse events considered at least possibly related to a trial
282 procedure to the coordinating center and then to the DSMB and national regulatory authorities,
283 as required. In LOVIT-COVID, data on hemolysis and hypoglycemia were collected as safety
284 outcomes. Additional in-hospital outcomes collected only in LOVIT-COVID and post-discharge
285 outcomes¹⁶ were not included in the statistical analysis plan and will be reported separately.

286

287 Statistical analysis

288 Following harmonization of both trials, the original fixed LOVIT-COVID sample size was
289 replaced by the REMAP-CAP Bayesian design with no maximum sample size. Adaptive
290 analyses were performed and response-adaptive randomization continued until reaching a pre-
291 defined statistical trigger, initially specified as efficacy, inferiority, and equivalence.

292

293 The statistical analysis plan for the harmonized trial specified that the trial outcomes would be
294 reported from a merged dataset created after both trials had stopped (additional details in
295 eMethods). The analysis used Bayesian cumulative logistic models, which calculated posterior
296 probability distributions based on accumulated trial evidence and a neutral prior distribution.
297 Distinct treatment effects of vitamin C compared to control were estimated in critically ill and
298 non-critically ill patients using a hierarchical prior that dynamically borrowed information
299 between groups. The hierarchical prior distribution was centered on an overall intervention effect
300 estimated with a prior assuming no treatment effect (standard normal prior on the log-odds ratio).
301 The primary statistical model, used to estimate the effect of vitamin C on organ support-free days,
302 and a similar model for hospital survival and for 28-day death or persistent organ dysfunction,
303 adjusted for trial (LOVID-COVID vs. REMAP-CAP); other interventions, and eligibility and
304 randomization in vitamin C domain (within REMAP-CAP); location (site, nested within country);
305 age (categorized into six groups); sex; and time-period (two-week calendar epochs) to account
306 for changes in clinical care and outcomes during the pandemic. Statistical models were fit using
307 a Markov Chain Monte Carlo algorithm that drew iteratively (20,000 draws) from the joint
308 posterior distribution. There were no terms for vitamin C interactions with other interventions.
309 The model included patients enrolled in all other domains of REMAP-CAP, including those that

310 remained blinded, to provide robust estimation of covariate effects. The Statistical Analysis
311 Committee conducted the analysis for patients with COVID-19 randomized up to July 15, 2022.
312
313 Patients were analyzed according to group assignment. Missing outcomes were not imputed.
314 Posterior odds ratios with 95% credible intervals (CrI) were calculated, with odds ratio >1
315 corresponding to superiority of vitamin C to control. The probabilities of efficacy (odds ratio >1),
316 harm (odds ratio <1), futility (odds ratio <1.2), and equivalence (odds ratio between $1/1.2$ and
317 1.2) were calculated. For the primary outcome, an ordinal scale with 24 categories (worst
318 category, death, and best category, alive with 21 days free of organ support), the odds ratio
319 denotes the relative odds of being in the category $>i$ vs. $\leq i$, for i equals -1 to 21 . The robustness
320 of the proportional odds assumption was assessed for the primary ordinal regression model. For
321 90-day survival, an adjusted hazard ratio with 95% CrI was calculated.
322
323 The original pre-defined statistical triggers for trial conclusions were based on posterior
324 probabilities of efficacy ($>99\%$, odds ratio for vitamin C >1), inferiority ($>99\%$, odds ratio <1),
325 and equivalence ($>90\%$, odds ratio between $1/1.2$ and 1.2). After LOVIT found that vitamin C
326 increased the risk of 28-day death or persistent organ dysfunction in sepsis,⁷ statistical triggers for
327 futility ($>95\%$, odds ratio <1.2) and harm ($>90\%$, odds ratio <1) were added.
328
329 Sensitivity analyses for the primary outcome and 28-day death or persistent organ dysfunction,
330 and analyses of all secondary outcomes, used data from patients enrolled in REMAP-CAP
331 domains that had stopped and were unblinded at the time of analysis to inform covariate
332 adjustment. Additional sensitivity analyses with different analysis populations, and pre-specified

333 subgroup analyses, are in the statistical analysis plan. One such analysis included 63 patients
334 with COVID-19 enrolled in LOVIT.⁷ Data management and summaries were created using R
335 version 4.1.2, and the primary analysis was computed in R version 4.1.3 using the rstan package
336 version 2.21.0 (R Foundation for Statistical Computing, Vienna, Austria).

337

338 **Results**

339 Patients

340 The first patient was randomized in LOVIT-COVID on August 23, 2020 and in the vitamin C
341 domain of REMAP-CAP on July 23, 2020. Both trials stopped recruitment on July 15, 2022 as
342 advised by their DSMBs, as statistical triggers for futility and harm had been met for both
343 critically ill and non-critically ill strata in REMAP-CAP. Interim analysis reports of both trials
344 are in eResults, and response-adaptive randomization proportions over time in REMAP-CAP are
345 shown in eFigure 1.

346

347 Of 2613 randomized patients, 7 were assessed as non-eligible, 15 withdrew consent for follow-
348 up, and one critically ill patient in the control group contributed baseline data but had a missing
349 primary outcome (Figure 1 and eFigures 2-3). The population for the primary statistical model
350 included 2590 randomized and evaluable patients, with 1493 patients assigned to vitamin C and
351 1097 assigned to control. There were 1568 critically ill patients from 90 sites and 1022 non-
352 critically ill patients from 40 sites, with 2206 enrolled in the vitamin C domain of REMAP-CAP
353 and 384 in LOVIT-COVID. Two critically ill patients included in the analysis withdrew consent
354 for follow-up but allowed for collected data to be used; their last known status was carried

355 forward for the primary outcome. Accrual rates over time are shown in eFigures 4-5. Covariate
356 effects were estimated from 9771 patients from any REMAP-CAP domain and LOVIT-COVID.
357
358 Baseline characteristics are reported in Table 1 and eTables 3-8. Patients were recruited from
359 Asia (34.7%), North America (28.5%), Europe (27.7%), and Australia (9.2%). Among critically
360 ill patients, respiratory support at enrollment included invasive ventilation (28.0%), non-invasive
361 ventilation (36.2%), and high-flow nasal oxygen (35.1%). Among non-critically ill patients, most
362 were receiving no respiratory support or low-flow oxygen (90.7%). Most patients received
363 corticosteroids (96.4%). In LOVIT-COVID, 96.1% of patients received $\geq 90\%$ of scheduled
364 doses (eTable 9); in REMAP-CAP, 95.2% of patients had no treatment delivery-related deviation
365 (eTable 10).

366

367 Primary outcome

368 Among critically ill patients, median organ-support free days were 7 (interquartile range [IQR] –
369 1, 17) in the vitamin C group vs. 10 (IQR –1, 17) in the control group (Table 2; Figure 2). The
370 odds ratio for vitamin C was 0.88 (95%CrI 0.73-1.06), yielding posterior probabilities of 8.6%
371 for efficacy, 91.4% for harm, and 99.9% for futility. Among non-critically ill patients, median
372 organ-support free days were 22 (IQR 18, 22) in the vitamin C group vs. 22 (IQR 21, 22) in the
373 control group (Table 3; Figure 3). The odds ratio for vitamin C was 0.80 (95%CrI 0.60-1.01),
374 yielding posterior probabilities of 2.9% for efficacy, 97.1% for harm, and >99.9% for futility.

375

376 Among critically ill patients, survival to hospital discharge was 61.9% (642/1037) in the vitamin
377 C group vs. 64.6% (343/531) in the control group. The odds ratio for vitamin C was 0.92

378 (95%CrI 0.73-1.17), with posterior probabilities of 24.0% for efficacy, 76.0% for harm, and 98.4%
379 for futility. Among non-critically ill patients, survival to hospital discharge was 85.1% (388/456)
380 in the vitamin C group vs. 86.6% (490/566) patients in the control group. The odds ratio for
381 vitamin C was 0.86 (95%CrI 0.61-1.17), with posterior probabilities of 17.8% for efficacy, 82.2%
382 for harm, and 98.1% for futility.

383

384 Secondary outcomes

385 Among critically ill patients, 90-day survival was 59.8% (617/1032) in the vitamin C group vs.
386 62.1% (328/528) in the control group (Table 2; Figure 2). The hazard ratio for vitamin C was
387 0.94 (95%CrI 0.80-1.11), with 22.4% posterior probability for efficacy. Among non-critically ill
388 patients, 90-day survival was 81.5% (370/454) in the vitamin C group vs. 82.8% (466/563)
389 patients in the control group (Table 3; Figure 3). The odds ratio for vitamin C was 0.93 (95%CrI
390 0.74-1.19), with 27.2% posterior probability of efficacy. Survival to 28 days without persistent
391 organ dysfunction was similar in critically ill patients (Table 2; odds ratio for vitamin C, 0.90 (95%
392 CrI 0.72-1.12; 16.4% probability of efficacy) and in non-critically ill patients (Table 3; odds ratio
393 for vitamin C, 0.92 (95% CrI 0.68-1.23; 26.6% probability of efficacy)

394

395 Posterior probabilities of superiority of vitamin C vs. control were less than 33% for all other
396 secondary outcomes (Tables 2-3; eFigures 6-7). Serious adverse events were reported in 1.8%
397 (27/1493) patients assigned to vitamin C and 0.8% (9/1098) assigned to control (eTable 11).
398 There were four serious adverse events possibly or probably related to vitamin C, including one
399 patient with methemoglobinemia, two with hypoglycemia, and one with hemolytic anemia
400 subsequently discovered to have glucose-6-phosphate dehydrogenase deficiency.

401

402 Sensitivity, subgroup, and exploratory analyses

403 Sensitivity analyses of organ support-free days, hospital survival, and 28-day mortality or
404 persistent organ dysfunction using different analysis populations were consistent with the
405 primary analyses (eTables 12-14). Credible intervals were wider in LOVIT-COVID compared to
406 REMAP-CAP, with no convincing evidence of divergent effect estimates (eTable 15). There
407 were no differential effects among subgroups (eTable 16). Exploratory analyses showed that the
408 in-hospital mortality rates by group in REMAP-CAP shifted over time (eFigure 8), with the
409 effect of vitamin C on organ support-free days varying over successive periods defined by
410 randomization ratio (eTable 17). *Post hoc* analyses of treatment effect by continent and by
411 dominant SARS-CoV-2 strain by month in each country of enrollment did not explain this
412 variation (eTables 18-19).

413

414 **Discussion**

415 In this large, harmonized, multinational randomized trial, vitamin C administered to hospitalized
416 patients with COVID-19 did not improve organ-support-free days or hospital survival. On the
417 contrary, there were high posterior probabilities (>90% for organ support-free days and >75%
418 for hospital survival) that vitamin C worsened both outcomes in critically ill and non-critically ill
419 patients. These effects were consistent across predefined subgroups and in sensitivity analyses.

420

421 The regimen of vitamin C was based on a previous trial in sepsis showing sustained elevation of
422 serum vitamin C levels over the treatment course, in addition to lower mortality, a secondary
423 outcome.¹⁸ The current results, from a critically ill population with mainly COVID-19

424 respiratory failure and a non-critically ill population, are consistent with the LOVIT trial among
425 septic patients treated with vasopressors.⁷ Existing analyses do not elucidate mechanisms of
426 harm, and while future biomarker analyses from LOVIT-COVID may be informative,¹⁹ the same
427 biomarkers measured in LOVIT were comparable between vitamin C and placebo groups.⁷ A
428 previous meta-analysis of nine trials, with the largest randomizing 100 patients, found a reduced
429 odds of mortality in COVID-19 patients receiving vitamin C.¹¹ These divergent results may be
430 explained by more extreme effects observed in small trials.²⁰

431

432 Several methodological issues are noteworthy. First, the initial decision to limit statistical
433 stopping triggers to efficacy, inferiority, and equivalence facilitated investigation of a small
434 treatment benefit. Although the current results do not exclude the possibility of any beneficial
435 effect of vitamin C in COVID-19, it is more likely that vitamin C is ineffective or harmful.
436 Second, this report provides separate effects of vitamin C in critically ill and non-critically ill
437 patients, consistent with the design. An alternative approach would have included all randomized
438 patients and generated a more precise overall treatment effect, with testing for a subgroup effect.
439 Nonetheless, the current model allowed for statistical borrowing between critically ill and non-
440 critically ill strata, thus mitigating the loss of statistical power. Third, treatment effects are
441 presented in relative terms, rather than as absolute effects better suited for shared decision-
442 making. The difference of 1.5 organ-support free days is considered minimally important by the
443 Food and Drug Administration,²¹ but patients' views are unknown. Finally, response-adaptive
444 randomization in REMAP-CAP, designed to favor assignment to the group with superior
445 outcomes at interim analyses, led to 69% of critically ill patients assigned to vitamin C, despite
446 lack of efficacy in both strata. This situation arose because early results in critically ill patients

447 favored vitamin C, without reaching a statistical trigger, with the final adaptive analysis
448 conducted 10 months after the penultimate one due to implementation of new processes for
449 international data flow. During this period, over 50% of enrollment occurred, without changes to
450 domain selection criteria or trial procedures. This analysis reported a reversed direction of
451 treatment effect, unexplained *post hoc*, underscoring the early instability of treatment effect
452 estimates in trials.²²⁻²⁴ Because the inferiority trigger was never reached, the trial may have
453 continued, even with more frequent analyses, until harm and futility triggers were introduced due
454 to external evidence.⁷ Options for avoiding this situation include frequent adaptive analyses or
455 forcing the randomization ratio to remain closer to 1:1.^{25,26}

456

457 Strengths of this report include selection of a vitamin C regimen based on promising initial
458 evaluations,^{18,27} excellent treatment adherence and follow-up, and enhanced generalizability
459 based on a broad geographical enrollment.²⁸

460

461 Limitations

462 This report combines data from two trials, initially designed differently, in an attempt to improve
463 efficiency and reduce waste in pandemic research.²⁹ Fewer patients were enrolled in the placebo-
464 controlled LOVIT-COVID trial, with differential post-randomization care possible for patients
465 enrolled in the open-label REMAP-CAP trial. Analyses showing comparable treatment effects in
466 these two trials were underpowered. Data on individual participants' vaccination status, vitamin
467 C product received, and baseline vitamin C levels were unavailable to inform subgroup analyses,
468 although a subgroup analysis by baseline vitamin C level in LOVIT was uninformative.⁷

469

470 In conclusion, in hospitalized patients with COVID-19, treatment with vitamin C did not
471 improve organ support-free days or hospital survival.

472

473

474

475 Author contributions:

476 Drs Adhikari and Lamontagne had full access to all the data in the study and take responsibility
477 for the integrity of the data and the accuracy of the data analysis. Drs Adhikari and Lamontagne
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479

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522

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611

612 Table 1 Baseline characteristics

	Critically ill		Non-critically ill	
	Vitamin C (n = 1037)	Control (n = 532)	Vitamin C (n = 456)	Control (n = 566)
Age in years, median (IQR)	60.0 (49.0-69.0)	61.0 (50.0-72.0)	63.0 (51.0-73.0)	62.0 (51.0-72.0)
Age category, n (%)				
18-49	268 (25.8)	122 (22.9)	97 (21.3)	132 (23.3)
50-69	512 (49.4)	253 (47.6)	204 (44.7)	258 (45.6)
70+	257 (24.8)	157 (29.5)	155 (34.0)	176 (31.1)
Female sex, n (%)	382 (36.8)	182 (34.2)	189 (41.4)	216 (38.2)
Male sex, n (%)	655 (63.2)	350 (65.8)	267 (58.6)	350 (61.8)
Body mass index, median (IQR) ^a	29.6 (25.7-35.3) (n=837)	29.6 (26.0-35.1) (n=437)	28.4 (25.0-33.8) (n=358)	28.4 (25.1-32.5) (n=435)
Continent, n (%)				
Asia	373 (36.0)	134 (25.2)	156 (34.2)	235 (41.5)
Australia	136 (13.1)	65 (12.2)	15 (3.3)	22 (3.9)
Europe	365 (35.2)	180 (33.8)	74 (16.2)	98 (17.3)
North America	163 (15.7)	153 (28.8)	211 (46.3)	211 (37.3)
Race / Ethnicity, ^b n / N (%)				
Asian	32/417 (7.7)	10/219 (4.6)	3/123 (2.4)	4/133 (3.0)
Black	16/417 (3.8)	12/219 (5.5)	14/123 (11.4)	13/133 (9.8)
Mixed or multiple	6/417 (1.4)	1/219 (0.5)	0/123 (0.0)	0/133 (0.0)
White	298/417 (71.5)	159/219 (72.6)	97/123 (78.9)	108/133 (81.2)
Other	65/417 (15.6)	37/219 (16.9)	9/123 (7.3)	8/133 (6.0)
APACHE II score, ^c median (IQR)	12.0 (8.0-18.0) (n=1031)	14.0 (8.0-21.0) (n=531)	8.0 (5.0-12.0) (n=278)	8.0 (5.0-11.0) (n=358)
Clinical Frailty Score, ^d median (IQR)	3.0 (2.0-3.0) (n=979)	3.0 (2.0-3.0) (n=492)	3.0 (2.0-3.0) (n=358)	3.0 (2.0-3.0) (n=463)
Preexisting condition, ^e n / N (%)				
Diabetes	323 (31.1)	159 (29.9)	133 (29.2)	138 (24.4)
Respiratory disease	167/1006 (16.6)	89/505 (17.6)	75/386 (19.4)	86/495 (17.4)
Kidney disease	68/919 (7.4)	46/446 (10.3)	24/371 (6.5)	37/488 (7.6)
Severe cardiovascular disease	42 (4.1)	32 (6.0)	25/455 (5.5)	35/565 (6.2)
Any immunosuppressive condition	35/998 (3.5)	33/496 (6.7)	21/367 (5.7)	20/468 (4.3)
Time to enrollment, median (IQR)				
From hospital admission, days ^f	1.1 (0.8-2.7)	1.1 (0.8-2.5)	1.0 (0.7-2.1)	1.0 (0.7-2.1)
From ICU admission, hours ^g	15.0 (8.5-19.9) (n=1034)	15.2 (8.8-20.0) (n=531)	15.6 (9.6-20.1) (n=219)	15.0 (7.2-21.0) (n=283)
Acute respiratory support, ^h n / N (%)				
Invasive mechanical ventilation	287/1036 (27.7)	151/531 (28.4)	0 (0.0)	0 (0.0)
Noninvasive ventilation only	393/1036 (37.9)	175/531 (33.0)	6 (1.3)	8 (1.4)
High-flow nasal oxygen	350/1036 (33.8)	200/531 (37.7)	35 (7.7)	46 (8.1)
None or low-flow oxygen	6/1036 (0.6)	5/531 (0.9)	415 (91.0)	512 (90.5)
Vasopressor support, n / N (%)	152/1036 (14.7)	76/531 (14.3)		

Concomitant therapies, n / N (%) [†]				
Remdesivir	403/974 (41.4)	174/463 (37.6)	211/355 (59.4)	267/471 (56.7)
Corticosteroids	990/1035 (95.7)	518/531 (97.6)	420 (92.1)	531/564 (94.1)
Tocilizumab or sarilumab	296/974 (30.4)	151/463 (32.6)	30/355 (8.5)	52/471 (11.0)

617

618 APACHE, Acute Physiology and Chronic Health Evaluation; IQR, interquartile range.

619 Control patients include all patients randomized to control who were also eligible to be randomized to vitamin C, i.e., direct concurrent controls.

620 Trial-specific baseline characteristics may be found in eTables 5-8.

621

622 Percentages may not sum to 100 because of rounding.

623

624 ^a The body mass index (BMI) is the weight in kilograms divided by the square of the height in meters.

625 ^b Collection of ethnicity data was approved in UK, Australia, and USA only, and data were not collected in LOVIT-COVID. “Other” includes any other racial or ethnic group reported.

626 ^c Acute Physiology and Chronic Health Evaluation II scores range from 0 to 71, with higher scores indicating greater severity of illness and higher risk of death.

627 ^d Scores on the Clinical Frailty Scale range from 1 to 9, with higher scores indicating greater frailty.

628 ^e Kidney disease was determined from the most recent serum creatinine level prior to this hospital admission, except in patients who were receiving dialysis.

629 Abnormal kidney function was defined as a creatinine level of 130 µmol/L or greater (1.5 mg/dL) for males or 100 µmol/L or greater (1.1 mg/dL) for females not previously receiving dialysis. Cardiovascular disease was defined as New York Heart Association class IV symptoms. In LOVIT-COVID, immunosuppressive conditions included receipt of recent chemotherapy or chronic immunosuppressive medications (excluding steroids), neutropenia, solid organ or stem cell transplantation, or human immunodeficiency virus positive status. In REMAP-CAP, these conditions included acquired immunodeficiency syndrome, metastatic cancer, specific hematological malignancies or other hematological conditions, or other inherited, primary, or secondary immune deficiencies.

630 ^f In LOVIT-COVID, hospital admission was recorded when the patient left the Emergency Department or when care in the Emergency Department was assumed by an inpatient service, depending on the hospital. In REMAP-CAP, time to enrolment from hospital admission explicitly includes all time spent in the Emergency Department.

631 ^g Patients in an intensive care unit but not receiving respiratory or cardiovascular organ support were prospectively classified as non-critically ill.

632 ^h Non-invasive ventilation and high-flow nasal oxygen delivered outside an intensive care unit did not fulfil the trial definition of critical illness.

633 ⁱ Concomitant therapies were given at baseline or within 48 hours of randomization (REMAP-CAP), or at baseline or on the day of or the day after randomization (LOVIT-COVID). Data on remdesivir and tocilizumab or sarilumab were specifically collected in REMAP-CAP, but could be recorded under ‘antiviral’ or ‘immunomodulator’ in LOVIT-COVID.

642

643

644 Table 2. Primary and secondary outcomes in critically ill participants

	Intravenous vitamin C	Control	Adjusted proportional Odds Ratio (95% CrI) ^a	Probability of efficacy / harm, %
Primary outcome				
Organ support-free days to day 21 ^b	Median (q1, q3) [N=1037] 7 (-1 to 17)	Median (q1,q3) [N=532] 10 (-1 to 17)	0.88 (0.73, 1.06)	8.6 / 91.4 ^c
Component of primary outcome				
Survival to hospital discharge	No. of patients/total no. (%) 642/1037 (61.9)	No. of patients/total no. (%) 343/531 (64.6)	0.92 (0.73, 1.17)	24.0 / 76.0 ^d
Secondary outcomes				
Survival without persistent organ dysfunction at day 28 ^e	No. of patients/total no. (%) 592/1037 (57.1)	No. of patients/total no. (%) 323/532 (60.7)	0.90 (0.72, 1.12)	16.4 / 83.6
Vasopressor/inotrope-free days through day 28	Median (q1, q3) [N=1037] 26 (-1, 28)	Median (q1, q3) [n=532] 27 (-1, 28)	0.84 (0.75, 0.94)	0.9 / 99.1
Respiratory support-free days through day 28	Median (q1, q3) [N=1037] 13 (-1, 24)	Median (q1, q3) [N=531] 16 (-1, 24)	0.89 (0.73, 1.01)	3.2 / 96.8
Endotracheal intubation through day 28	No. of patients/total no. (%) 266/750 (35.5)	No. of patients/total no. (%) 124/381 (32.5)	0.74 (0.56, 0.99)	2.1 / 97.9
Extracorporeal support through day 28 ^f	No. of patients/total no. (%) 12/1034 (1.2)	No. of patients/total no. (%) 7/532 (1.3)		
Survival to day 28	No. of patients/total no. (%) 671/1032 (65.0)	No. of patients/total no. (%) 356/530 (67.2)	0.94 (0.75, 1.19)	31.2 / 68.8
Discharge alive from the ICU ^g			0.96 (0.84, 1.10)	28.4 / 71.6
Discharge alive from the hospital ^g			0.93 (0.82, 1.05)	12.1 / 87.9
90-day survival ^h	No. of patients/total no. (%) 617/1032 (59.8)	No. of patients/total no. (%) 328/528 (62.1)	0.94 (0.80, 1.11)	22.4 / 77.6
WHO ordinal scale at day 14 ⁱ			0.89 (0.75, 1.07)	11.0 / 89.0

645

646 CrI, credible interval; ICU, intensive care unit; IQR, interquartile range; WHO, World Health Organization

647 ^a The odds ratio is for vitamin C relative to control.
648 ^b The model assigns hospital decedents a value of -1 organ support-free days.
649 ^c The probability of futility was 99.9%.
650 ^d The probability of futility was 98.4%.
651 ^e The outcome is the complement of 28-day mortality or persistent organ dysfunction to preserve the interpretation
652 of odds ratio >1 denoting superiority of vitamin C.
653 ^f No model was constructed for this outcome, as per the statistical analysis plan.
654 ^g Crude results are not provided because the model assigns hospital decedents a length of stay of 90 days.
655 ^h The 90-day survival proportions exclude from the denominator patients censored alive prior to 90 days (8 critically
656 ill patients were censored).
657 ⁱ The WHO ordinal scale measures the patient's overall status at day 14; range: 0-8, where 0 denotes no illness, 1-7
658 denote increasing level of care, and 8 denotes death.³⁰ In this analysis, categories 0, 1, and 2 have been condensed
659 into one category for all patients discharged from hospital. In LOVIT-COVID, states 3 and 4 were collapsed into
660 one category.
661
662

663 Table 3. Primary and secondary outcomes in non-critically ill participants

	Intravenous vitamin C	Control	Adjusted proportional Odds Ratio (95% CrI) ^a	Probability of efficacy / harm, %
Primary outcome				
Organ support-free days to day 21 ^b	Median (q1, q3) [N=456] 22 (18 to 22)	Median (q1,q3) [N=566] 22 (21 to 22)	0.80 (0.60, 1.01)	2.9 / 97.1 ^c
Component of primary outcome				
Survival to hospital discharge	No. of patients/total no. (%) 388/456 (85.1)	No. of patients/total no. (%) 490/566 (86.6)	0.86 (0.61, 1.17)	17.8 / 82.2 ^d
Secondary outcomes				
Survival without persistent organ dysfunction at day 28 ^e	No. of patients/total no. (%) 381/456 (83.6)	No. of patients/total no. (%) 477/566 (84.3)	0.92 (0.68, 1.23)	26.6 / 73.4
Vasopressor/inotrope-free days through day 28	Median (q1, q3) [N=456] 28 (28, 28)	Median (q1, q3) [n=566] 28 (28, 28)	0.77 (0.65, 0.90)	0.5 / 99.5
Respiratory support-free days through day 28	Median (q1, q3) [N=456] 28 (26, 28)	Median (q1, q3) [N=566] 28 (27, 28)	0.83 (0.64, 0.99)	1.9 / 98.1
Endotracheal intubation through day 28	No. of patients/total no. (%) 63/456 (13.8)	No. of patients/total no. (%) 50/566 (8.8)	0.59 (0.38, 0.83)	0.1 / 99.9
Extracorporeal support through day 28 ^f	No. of patients/total no. (%) 2/456 (0.4)	No. of patients/total no. (%) 4/566 (0.7)		
Survival to day 28	No. of patients/total no. (%) 385/454 (84.8)	No. of patients/total no. (%) 480/563 (85.3)	0.94 (0.68, 1.26)	32.9 / 67.1
Discharge alive from the hospital ^g			0.92 (0.81, 1.05)	10.6 / 89.4
90-day survival ^h	No. of patients/total no. (%) 370/454 (81.5)	No. of patients/total no. (%) 466/563 (82.8)	0.93 (0.74, 1.19)	27.2 / 72.8
WHO ordinal scale at day 14 ⁱ			0.89 (0.71, 1.12)	15.6 / 84.4

664

665 CrI, credible interval; ICU, intensive care unit; IQR, interquartile range; WHO, World Health Organization

666 ^a The odds ratio is for vitamin C relative to control.
667 ^b The model assigns hospital decedents a value of -1 organ support-free days.
668 ^c The probability of futility was >99.9%.
669 ^d The probability of futility was 98.1%.
670 ^e The outcome is the complement of 28-day mortality or persistent organ dysfunction to preserve the interpretation
671 of odds ratio >1 denoting superiority of vitamin C.
672 ^f No model was constructed for this outcome, as per the statistical analysis plan.
673 ^g Crude results are not provided because the model assigns hospital decedents a length of stay of 90 days.
674 ^h The 90-day survival proportions exclude from the denominator patients censored alive prior to 90 days (4 non-
675 critically ill patients were censored).
676 ⁱ The WHO ordinal scale measures the patient's overall status at day 14; range: 0-8, where 0 denotes no illness, 1-7
677 denote increasing level of care, and 8 denotes death.³⁰ In this analysis, categories 0, 1, and 2 have been condensed
678 into one category for all patients discharged from hospital. In LOVIT-COVID, states 3 and 4 were collapsed into
679 one category.
680

681 **Figure 1** Flow of patients through the harmonized trial. Additional details are provided in
682 Figures S2 and S3. ITT, intention to treat; SC, steering committee. SDM: Surrogate decision
683 maker.

684
685 ^a Other reasons why patients were excluded in LOVIT-COVID: 7 Had known G6PD deficiency;
686 3 Had known sickle cell anemia, 2 Had known allergy to vitamin C, 17 Had known kidney
687 stones within the past 1 year, 1 Received IV vitamin C (not incorporated into parenteral nutrition).

688 ^b Other reasons why eligible patients were not enrolled in LOVIT-COVID: 12 Had SDM that
689 was unable to be reached, 18 Were missed (off-business hours), 1 Was enrolled in a trial for
690 which co-enrollment was not allowed, and 129 for: 74 had no reason, 33 were diabetic patients
691 (glucose monitoring requiring too much work for the nursing staff), 5 were asymptomatic
692 COVID patients hospitalized for another reason, 4 were discharged before the responsible
693 physician get back to the research team on patient's eligibility, 2 were disoriented or had
694 dementia and no SDM, 2 were palliative or deemed palliative, 2 had a language barrier, 1 had
695 passive decline, 1 was being discharged, 1 was transferred to another hospital after intubation, 1
696 had acute kidney injury, 1 had planned renal transplant, 1 was not enrolled due to research team
697 workload, 1 was due for several interventions with no possibility of approach within 24
698 hours.

699 ^c Patients could meet more than one ineligibility criterion.

700 ^d Other reasons in Vitamin C Domain active and not enrolled in another domain: 10 Received IV
701 vitamin C during this hospital admission, 5 Patients randomized to another trial of vitamin C.

702 ^e Other reasons in Vitamin C Domain active: 19 Patients randomized to another trial of vitamin
703 C, 12 Reveal of allocation not completed, 1 Other.

704 ^f Randomization was stratified by site in LOVIT-COVID and by population (critically ill vs.
705 non-critically ill) in REMAP-CAP.

706 ^g The principal investigators designed both LOVIT-COVID and the vitamin C domain of
707 REMAP-CAP, and with support of the respective steering committees, *a priori* decided to use a
708 common vitamin C treatment regimen, collect a set of common outcomes, and conduct a merged
709 analysis after both trials had completed recruitment.

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714 **Figure 2** Critically ill patients. Panel A: The cumulative proportion (y-axis) for vitamin C (blue
715 line) or control (red line) by day (x-axis) of organ support-free days, with death listed first.
716 Curves that rise more slowly indicate a more favorable distribution in the number of days alive
717 and free of organ support. Panel B: Organ support-free days as horizontally stacked proportions
718 by intervention group. Red represents worse outcomes and blue represents better outcomes. The
719 median adjusted odds ratio from the primary analysis was 0.88 (95% credible interval, 0.73 to
720 1.06), yielding 8.6% probability of vitamin C being superior to control. Panel C: 90-day survival.
721 There were 415/1032 deaths (40.2%) in the vitamin C group and 200/528 deaths (37.9%) in the
722 control group. Denominators exclude censored patients. The blue line represents vitamin C and
723 the red line represents control. Data was available on all patients through death or 90 days except
724 for 8 patients that were censored alive prior to 90 days.
725

726 **Figure 3** Non-critically ill patients. Panel A: The cumulative proportion (y-axis) for vitamin C
727 (blue line) or control (red line) by day (x-axis) of organ support-free days, with death listed first.
728 Curves that rise more slowly indicate a more favorable distribution in the number of days alive
729 and free of organ support. Panel B: Organ support-free days as horizontally stacked proportions
730 by intervention group. Red represents worse outcomes and blue represents better outcomes. The
731 median adjusted odds ratio from the primary analysis was 0.80 (95% credible interval, 0.60 to
732 1.01), yielding 2.9% probability of vitamin C being superior to control. Panel C: 90-day survival.
733 There were 84/454 deaths (18.5%) in the vitamin C group and 97/563 deaths (17.2%) in the
734 control group. Denominators exclude censored patients. The blue line represents vitamin C and
735 the red line represents control. Data was available on all patients through death or 90 days except
736 for 4 patients that were censored alive prior to 90 days.
737
738

LOVIT-COVID

2 027 Adults hospitalized with confirmed COVID-19 were assessed for eligibility

1 208 Were excluded^a
 443 Were receiving or have received vasopressors during the current hospitalisation
 510 Had more than 24 hours elapsed since receipt of non-invasive ventilatory support (high-flow nasal cannula or continuous positive airway pressure or non-invasive ventilation) or invasive mechanical ventilation
 143 Were expected to be discharged from the hospital in the next 24 hours
 29 Had more than 14 days elapsed since the commencement of hospital admission with respiratory illness
 28 Were pregnant or were breastfeeding
 25 Had expected death or withdrawal of life sustaining therapies within 48 hours

427 Were eligible but not enrolled^b
 187 Patients or SDM declined consent
 80 Had a treating physician who declined consent
 129 Were not enrolled for other reasons

392 Underwent randomization^f

194 Were assigned to the vitamin C group 198 Were assigned to the control group

84 Were critically ill	110 Were non-critically ill	98 Were critically ill	100 Were non-critically ill
	3 Were not eligible per adjudication by two SC members. 1 Withdrew consent for all data	1 Was not eligible per adjudication by two SC members	3 Were not eligible per adjudication by two SC members
84 Had complete follow-up for primary outcome	106 Had complete follow-up for primary outcome	97 Had complete follow-up for primary outcome	97 Had complete follow-up for primary outcome

Vitamin C Domain of REMAP-CAP

22 452 Eligibility assessments for patients admitted with suspected or proven COVID-19 on or before 15th July 2022

8 463 Ineligible for platform
 2 479 Site not active for Vitamin C Domain and not enrolled in another domain
 1 489 Vitamin C Domain active and not enrolled in another domain^{c,d}
 889 More than 24 hours since admission to ICU
 50 Contraindication to agents in domain
 179 Not considered in the patient's best interests
 560 Prospective consent declined or not obtained

10 021 Randomizations enrolled in one or more REMAP-CAP domains

7 800 Ineligible or not assessed for Vitamin C Domain
 7 201 Site not active for Vitamin C Domain
 599 Vitamin C Domain active^{c,e}
 280 More than 24 hours since admission to ICU
 25 Received IV vitamin C during this hospital admission
 29 Contraindication to agents in domain
 104 Not considered in the patient's best interests
 151 Prospective consent declined or not obtained

2 221 Underwent randomization^f

1310 Were assigned to receive vitamin C 911 Were assigned to receive no vitamin C

957 Were critically ill	353 Were non-critically ill	439 Were critically ill	472 Were non-critically ill
4 Withdrew consent	3 Withdrew consent	4 Withdrew consent. 1 Had outcome not available	3 Withdrew consent
953 Had complete follow-up for primary outcome	350 Had complete follow-up for primary outcome	434 Had complete follow-up for primary outcome	469 Had complete follow-up for primary outcome

7 800 Assigned to receive an intervention in another domain

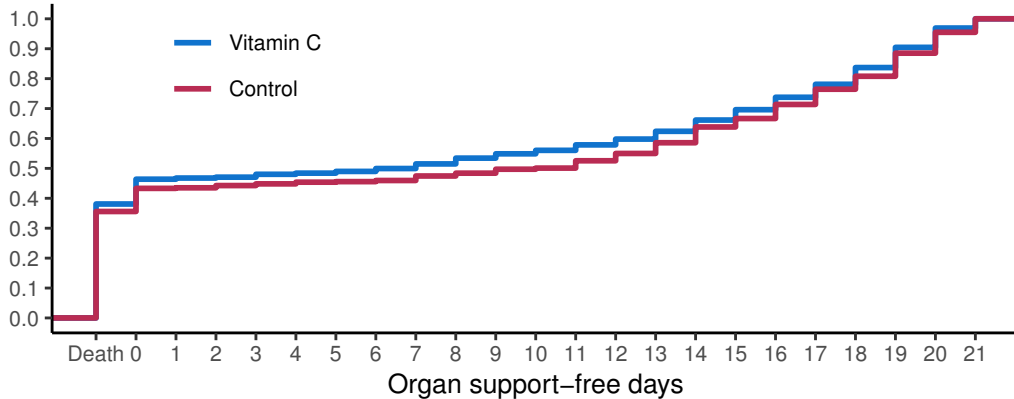
162 Withdrew consent
 2 Outcome not available
 424 Not randomized to a modeled domain

7 212 Used for covariate adjustment

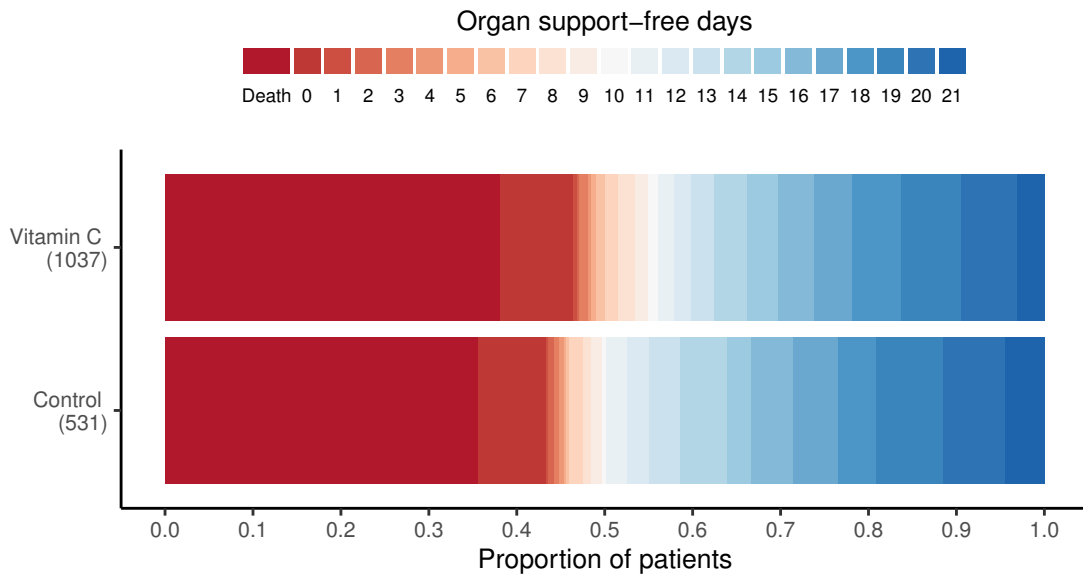
Harmonization^g

Vitamin C, critically ill	Vitamin C, non-critically ill	Placebo or no vitamin C, critically ill	Placebo or no vitamin C, non-critically ill
1037 (84+953) Had complete follow-up for primary outcome	456 (106+350) Had complete follow-up for primary outcome	531 (97+434) Had complete follow-up for primary outcome	566 (97+469) Had complete follow-up for primary outcome

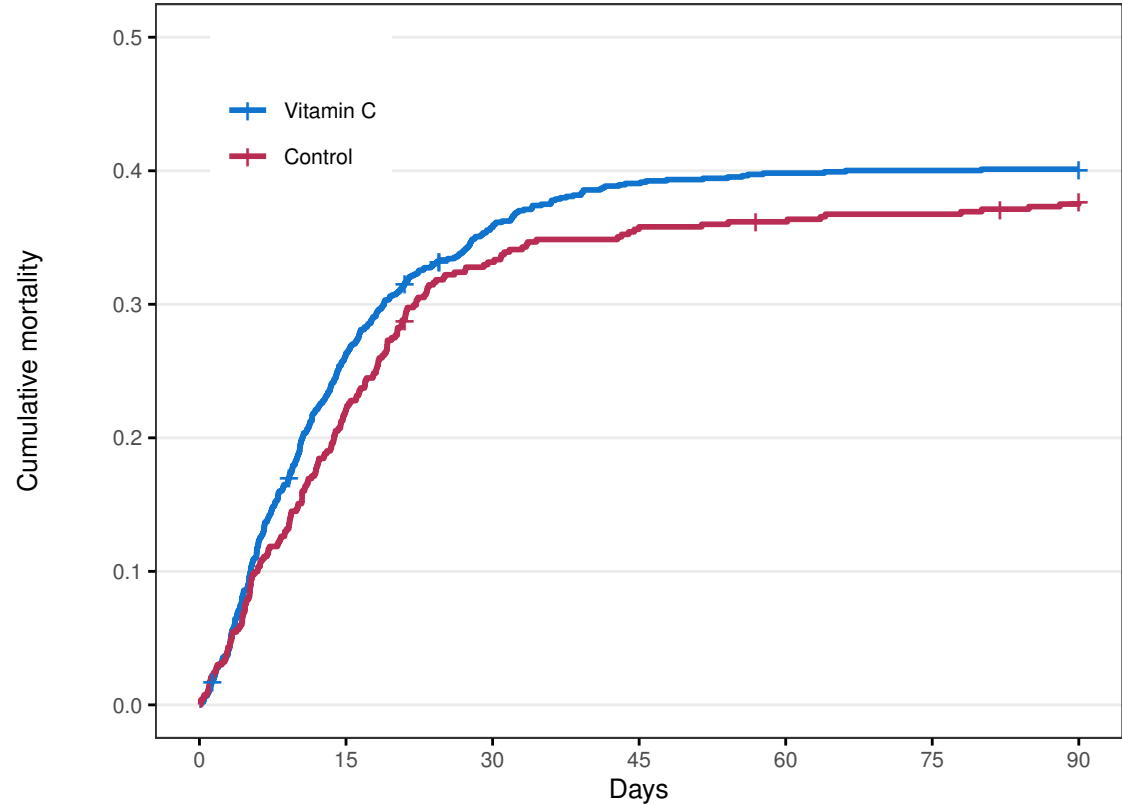
A) Cumulative proportion of organ support-free days



B) Stacked barplot of organ support-free days



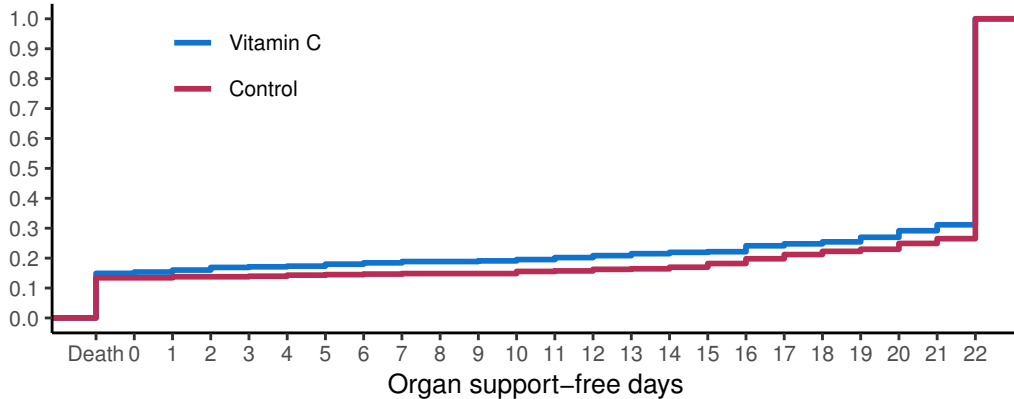
C) Survival through 90 days



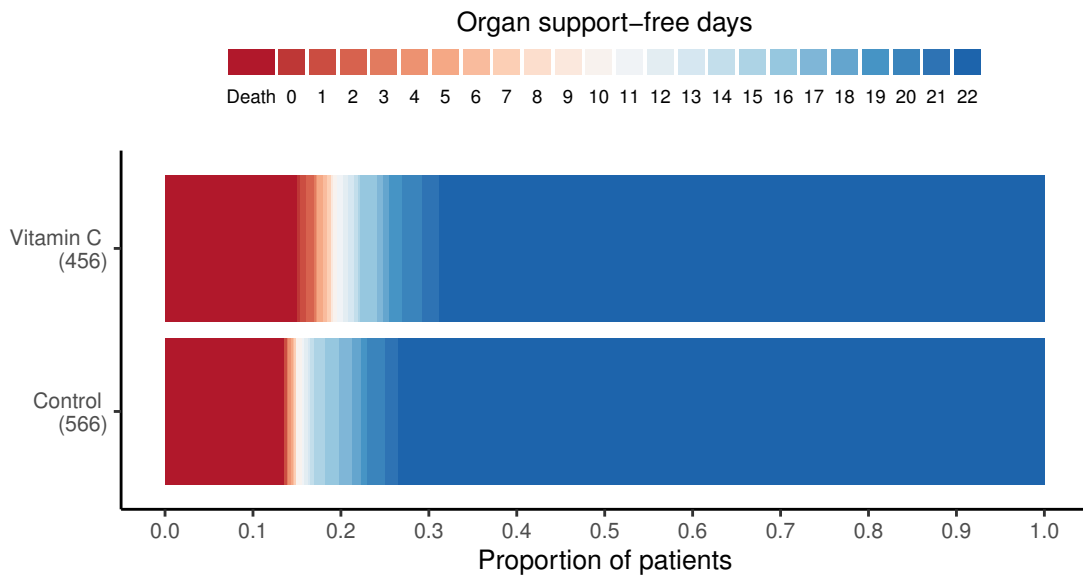
Number at risk

	0	15	30	45	60	75	90
Vitamin C	1037	763	662	628	620	618	617
Control	531	414	354	340	337	334	329

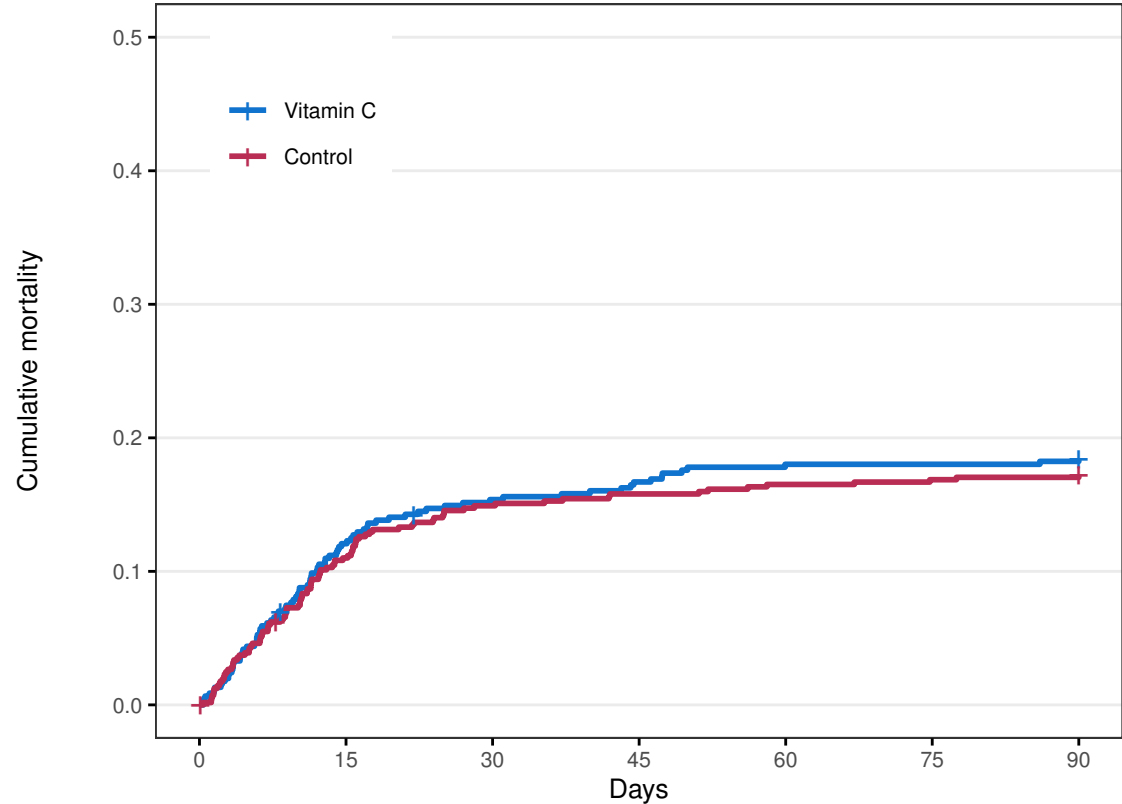
A) Cumulative proportion of organ support-free days



B) Stacked barplot of organ support-free days



C) Survival through 90 days



Number at risk

	0	15	30	45	60	75	90
Vitamin C	456	400	384	378	372	372	371
Control	565	501	479	474	470	468	467