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### ESPEN guideline on clinical nutrition in acute and chronic pancreatitis

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ESPEN guideline on clinical nutrition in acute and chronic pancreatitis

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#### ESPEN guideline on clinical nutrition in acute and chronic pancreatitis 1

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37	nutritional support, medical nutrition

39 Abbreviations:

ACS, Acute Compartmental Syndrome; ANP, Acute Necrotizing Pancreatitis; AP, Acute 40 41 Pancreatitis; BMI, Body Mass Index; CP, Chronic Pancreatitis; DPEJ, Direct Percutaneous 42 Endoscopic Jejunostomy; DXA, Dual-energy X-ray Absorptiometry; EN, Enteral Nutrition; IAH, Intra-abdominal Hypertension; IAP, Intra-abdominal Pressure; MCT, 43 Medium Chain Triglycerides; MUST, Malnutrition Universal Screening Tool; NAFLD, Non 44 Alcoholic Fatty Liver Disease; ONS, Oral Nutritional Supplements; PEG-J, Percutaneous 45 46 Endoscopic Gastrostomy with Jejunal Extension; PEI, Pancreatic Exocrine Insufficiency; PERT, Pancreatic Enzyme Replacement Therapy; PN, Parenteral Nutrition; PPI, Proton 47 Pump Inhibitor; RCT, Randomized Controlled Trial; SIBO, Small Intestinal Bacterial 48 49 Overgrowth; VARD, Video-assisted Retroperitoneal Debridement;

OUTAN

### 50 Abstract

51 Both acute and chronic pancreatitis are frequent diseases of the pancreas, which, despite 52 being of benign nature, are related to a significant risk of malnutrition and may require 53 nutritional support. Acute necrotizing pancreatitis is encountered in 20% of patients 54 with acute pancreatitis, is associated with increased morbidity and mortality, and may 55 require artificial nutrition by enteral or parenteral route, as well as additional 56 endoscopic, radiological or surgical interventions. Chronic pancreatitis represents a 57 chronic inflammation of the pancreatic gland with development of fibrosis. Abdominal 58 pain leading to decreased oral intake, as well as exocrine and endocrine failure are 59 frequent complications of the disease. All of the above represent risk factors related to malnutrition. Therefore, patients with chronic pancreatitis should be considered at risk, 60 61 screened and supplemented accordingly. Moreover, osteoporosis and increased facture risk should be acknowledged in patients with chronic pancreatitis, and preventive 62 measures should be considered. 63

64

### 65 **1. Introduction**

66 Acute pancreatitis (AP) is the most common acute gastrointestinal disease requiring 67 hospital admission (1), with the outcome being favorable in most cases (80%) (2). 68 However, acute necrotizing pancreatitis (ANP) may develop in up to 20% of patients and 69 is associated with significant rates of early organ failure (38%), need for intervention (38%), and death (15%) (2). Catabolism is very high in this setting; therefore, 70 71 nutritional support is one of the cornerstones of management (3). A significant amount 72 of research has shown the superiority of enteral over parenteral nutrition in ANP, 73 creating a paradigm shift a decade ago and modifying the management strategy (3). 74 Nevertheless, additional questions regarding the timing, route and type of enteral 75 nutrition (EN), as well as the place of oral refeeding, are still the objects of clinical 76 investigations.

77 Chronic pancreatitis (CP) is a disease in which recurrent inflammatory episodes lead to 78 replacement of the pancreatic parenchyma by fibrous connective tissue (4). The major 79 consequence of CP is the loss of functional exocrine and endocrine pancreatic tissue, thus resulting in both exocrine and endocrine insufficiency (4). Pain is also frequently 80 81 encountered in patients with CP, and seems to be related to a multitude of factors such 82 as pancreatic neural remodeling and neuropathy, increased intraductal and parenchymal pressure, pancreatic ischemia and acute inflammation during an acute 83 84 relapse (5). Both pain and loss of pancreatic function can lead to malnutrition in patients 85 with CP (4). Moreover, other long-term consequences such as osteoporosis are 86 frequently overlooked, despite their potential impact on quality of life in patients with 87 CP. Therefore, screening for malnutrition and nutritional support play a crucial part in 88 the multimodal management required in this setting.

Although recent guidelines for AP (2) and CP (4) have been published, a dedicatedconsensus on nutritional support in pancreatic diseases is lacking.

### 91 2. Methods

92 The present guideline was developed according to the standard operating procedure for
93 ESPEN guidelines (6). The guideline was developed by an expert group of 13 authors
94 from eleven European countries.

### 95 Methodology of guideline development

96 Based on the standard operating procedures for ESPEN guidelines and consensus 97 papers, the first step of the guideline development was the formulation of so-called PICO 98 questions which address specific **p**atient groups or **p**roblems, interventions, **c**ompare 99 different therapies and are outcome-related (6). In total, 31 PICO questions were 100 created and split into two main chapters, "Acute pancreatitis" and "Chronic 101 Pancreatitis". To answer these PICO questions, a literature search was performed to 102 identify suitable meta-analyses, systematic reviews and primary studies, published from 103 1977 up to December 2018. The PICO questions were allocated to subgroups/experts 104 for the different subjects who created 42 recommendations and seven statements. For 105 grading the literature, the grading system of the Scottish Intercollegiate Guidelines 106 Network (SIGN) was used (7). Allocation of studies to the different levels of evidence is 107 shown in Table 1. Supportive of the recommendations, the working group added 108 commentaries to the recommendations where the bases of the recommendations are 109 explained.

The recommendations were graded according to the levels of evidence assigned (Table
2). The wording of the recommendations reflect the grades of recommendations, level A
is indicated by "shall", level B by "should" and level 0 by "can/may". The good practice

point (GPP) is based on experts' opinions due to the lack of studies, here, the wordingcan be chosen deliberately.

115 Online voting on the recommendations was performed on the guideline-services.com 116 platform. All ESPEN members were invited to agree or disagree with the 117 recommendations and to comment on them. A first draft of the guideline was also made 118 available to the participants; on that occasion 36 recommendations and all seven 119 statements reached an agreement of >90%, six recommendations reached an agreement 120 of 75-90% and no recommendation an agreement of <75%. Those recommendations with an agreement of >90%, which means a strong consensus (Table 3) were passed 121 122 directly; all others were revised according to the comments and voted on again during a consensus conference, which took place on 29th April 2019. All recommendations 123 received an agreement of >90%. During the consensus conference, one of the original 124 125 recommendations was considered redundant and one statement was transformed into a 126 recommendation. Therefore, the guideline comprises 42 recommendations and six 127 statements. To support the recommendations and the assigned grades of 128 recommendation, the ESPEN guideline office created evidence tables of relevant meta-129 analyses, systematic reviews and randomized controlled trials (RCTs). These evidence 130 tables are available online as supplemental material to this guideline.

### 131 Search strategy

A comprehensive literature research including systematic reviews, controlled clinical trials and cohort studies, with the keywords and filters presented in Table 4 was performed. We initially searched Pubmed, Cochrane Library and EMBASE for recent, rigorous systematic reviews and meta-analyses that answered our clinical questions. In the absence of these, we looked for comparative studies, whether randomized or not.

137 The search phrases included the following terms: (acute pancreatitis OR acute 138 necrotizing pancreatitis OR chronic pancreatitis OR pancreatitis OR 139 hypertriglyceridemic pancreatitis OR hyperlipidemic pancreatitis) AND (nutritional 140 status OR nutritional assessment OR nutritional screening OR malnutrition OR oral 141 feeding OR enteral nutrition OR tube feeding OR parenteral nutrition OR intravenous 142 nutrition OR timing OR formula OR formulation OR nasogastric tube OR nasojejunal tube OR digestive intolerance OR necrosectomy OR minimally invasive OR increased intra-143 abdominal pressure OR abdominal compartment syndrome OR open abdomen OR 144 145 immunonutrition OR glutamine OR antioxidants OR probiotics OR enzyme 146 supplementation OR enzyme replacement therapy OR micronutrients OR 147 macronutrients OR nutrient deficiency OR diet OR fat OR nitrogen OR dietary protein OR carbohydrates oral supplementation OR medium chained triglycerides OR osteoporosis 148 149 OR osteopenia).

150 Finally, 88 articles were selected for the AP chapter, and 111 articles for the CP chapter.

151

152

### 153 **3. Results**

- 154 I. Acute pancreatitis
- 155 1. Which patients with AP are considered at nutritional risk?
- 156 Statement 1
- 157 Patients with AP should be considered at moderate to high nutritional risk,
- 158 because of the catabolic nature of the disease and because of the impact of the
- 159 nutritional status for disease development.
- 160 **Strong consensus (97% agreement)**
- 161

### 162 **Recommendation 1**

All patients with predicted mild to moderate AP should be screened using
validated screening methods such as the Nutritional Risk Screening - 2002 (NRS2002); however, the patients with predicted severe AP should always be
considered at nutritional risk.

### 167 Grade of Recommendation B – Strong consensus (100% agreement)

168

### 169 **Commentary**

Fortunately, the majority of patients with AP have predicted mild or moderately severe forms of the disease that are self-limited with fully recovery in less than a week, in whom oral feeding can be started within few days after the onset of AP (9). Gut-barrier dysfunction may occur in up to 60% of patients with AP; mostly in severe AP and it is thought to lead to bacterial translocation and infection of necrosis (10). Along with the increased catabolic state related to the disease, patients with predicted severe AP are

176 considered at nutritional risk (11). Nevertheless, malnourished patients should also be 177 considered at nutritional risk, even if they have predicted mild AP, because of their pre-178 existing condition. Similarly, patients with increased alcohol consumption are frequently 179 malnourished (12). Scoring systems such as the NRS 2002 (13), can be helpful in 180 identifying these patients (14-17). These scores have been validated in hospitalized, as 181 well as critically ill patients. Nevertheless, no studies have validated these scoring 182 systems in a specific population of patients with AP (18). 183 A low body mass index (BMI) may also identify patients who are at nutritional risk. 184 Nevertheless, obesity is a known risk factor for severe AP and is, therefore, a disease

- 185 severity-related nutritional risk (19).
- 186
- 187 2. Is early oral feeding feasible in patients with predicted mild AP?
- 188 **Recommendation 2**
- 189 Oral feeding shall be offered as soon as clinically tolerated and independent of
- 190 serum lipase concentrations in patients with predicted mild AP.
- 191 Grade of Recommendation A Strong consensus (100% agreement)
- 192
- 193 **<u>Recommendation 3</u>**
- 194 Low-fat, soft oral diet shall be used when reinitiating oral feeding in patients with

195 **mild AP.** 

- 196 Grade of Recommendation A Strong consensus (100% agreement)
- 197
- 198 Commentary

199 Most patients with AP suffer from disease of a mild to moderate severity, non-200 necrotizing type with an uncomplicated clinical course. Four RCTs have shown that 201 patients with mild to moderate AP can tolerate early oral feeding and this strategy is 202 related with a shorter length of stay compared with conventional oral feeding 203 (introduced after enzyme decrease, pain resolution and bowel movement) (9, 20-23). 204 Furthermore, one of these trials revealed that oral food intake is safe and well-tolerated 205 independently of the course and normalization of serum lipase (20). Immediate oral 206 feeding with a soft diet seems to be more beneficial regarding caloric intake and equally 207 tolerated compared with clear liquid diets (23-25). A meta-analysis confirmed that early 208 oral feeding was feasible in patients with predicted mild AP and reduced length of stay 209 (26). A recent meta-analysis including 17 studies identified that 16.3% of patients with 210 AP will subsequently have intolerance to oral feeding (27). Predictive factors included 211 the presence of pleural effusions and/or collections and severity (higher 212 Ranson/Glasgow and Balthazar scores).

Hyperlipidemia is the third most common cause of AP and accounts for 4-10% of cases 213 (28). It was reported that hyperlipidemia is associated with a worse prognosis of AP 214 215 than other etiological factors (28-30). The initial management of hyperlipidemic AP is 216 the same as for all other causes of the disease, but subsequent management in addition 217 to generalized supportive measures may include etiology-specific targeted therapies. 218 These include initially putting patients on a nil by mouth regimen for 24-48 hours, 219 subsequent dietary modifications, medical management with the different classes of 220 anti-hyperlipidemic agents, in-hospital pharmacological treatment with insulin and/or 221 heparin and plasmapheresis. Whilst these measures are effective in lowering triglyceride concentrations, they do not appear to affect the outcome of AP (28, 29). 222 However, tight regulation of triglyceride concentration after presentation with AP was 223

## 224 found to reduce the risk of recurrence. These include a low fat diet, encouragement of weight loss and treatment with a fibrate, with the addition of a statin if 225 226 hypercholesterolemia is present in addition to hypertriglyceridemia (28). 227 228 3. If required, what type of medical nutrition (enteral or parenteral) is preferable in 229 patients with AP? 230 **Recommendation 4** 231 In patients with AP and inability to feed orally, EN shall be preferred to parenteral 232 nutrition (PN). 233 Grade of Recommendation A - Strong consensus (97% agreement) 234 235 **Commentary** 236 EN is supposed to preserve the integrity of the gut mucosa, stimulate intestinal motility, 237 prevent bacterial overgrowth, and increase the splanchnic blood flow (10). Currently 238 there are twelve RCTs and eleven systematic reviews/meta-analyses including a 239 Cochrane-standard meta-analysis which clearly prove that in patients with severe AP, 240 EN is safe and well-tolerated, with significant decreases in complication rates, multi-241 organ failure, and mortality, compared with PN (31-41). The meta-analysis by Al-Omran

*et al.* was performed to Cochrane-standards on the basis of eight RCTs with 348 patients
and clearly shows that early EN when compared with initial total PN, significantly
decreases mortality by 50% (OR 0.50 [95% CI 0.28 to 0.91]), rate of infection (OR 0.39
[95% CI 0.23 to 0.65]), multi-organ failure (0.55 [95% CI 0.37 to 0.81]) as well as the
necessity for operation (OR 0.44 [95% CI 0.29 to 0.67]) (35). Furthermore if only
patients with severe AP were included in this meta-analysis, mortality further decreased

248	by more than 80% [0.18 [95 % CI 0.006 to 0.58]) (35). These results were confirmed by
249	more recent meta-analyses, including a latest publication including only critically ill
250	patients with AP (39). Compared with PN, EN was associated with a significant
251	reduction in overall mortality (RR 0.36, 95% CI 0.20 to 0.65, p=0.001) and the rate of
252	multiple organ failure (RR 0.39, 95% CI 0.21 to 0.73, p=0.003).
253	
254	4. What is the optimal timing for initiating EN in patients with AP?
255	Recommendation 5
256	EN should be started early, within 24-72 hours of admission, in case of intolerance
257	to oral feeding
258	Grade of Recommendation B – Strong consensus (92% agreement)
259	
260	Commentary
261	Soveral meta-analyses have investigated the clinical effects and telerance of early FN in

261 Several meta-analyses have investigated the clinical effects and tolerance of early EN in 262 patients with AP either within 24 hours (42-44) or 48 hours (45-47) of admission. All 263 these meta-analyses clearly reveal that early EN is feasible, safe and well-tolerated and 264 associated with substantial clinical benefits regarding mortality, organ failure and 265 infectious complications for both time-points compared with delayed EN. Nevertheless, 266 a potential bias could be that five of these meta-analysis included studies which had 267 patients receiving PN in their control groups (42-46). One meta-analysis, compared 268 early (within 24 hours) with late enteral nutrition (after 72 hours), but no comparison 269 was made between 24 and 48 hours (44).

In contrast to these data from the aforementioned meta-analyses that provided strong
evidence for early EN within 24-48 hours, a multicenter RCT (208 patients with

272 predicted severe AP) found no difference in the rate of major infection or death between 273 early EN, started within 24 hours after admission, and an oral diet initiated 72 hours 274 after admission (48). A second RCT (214 patients with AP) confirmed these results, 275 showing no significant reduction in persistent organ failure and mortality in patients 276 receiving early EN compared with patients receiving no nutritional support (49). A 277 plausible explanation could be that these trials included mostly patients with mild or 278 moderate AP (in the Bakker trial there were only 63% of cases with necrotizing AP 279 (48)); therefore, the beneficial effect of early EN could be less pronounced.

Finally, a prospective cohort study including 105 patients with AP concluded that the third day after hospital admission was the best cut-off time for early EN (with an area under the curve of 0.744), by reducing the risk of secondary infection and improving the nutritional status of patients, with a better tolerance (50).

284

285 5. What type of EN is indicated?

- 286 **Recommendation 6**
- 287 In patients with AP a standard polymeric diet shall be used.
- 288 Grade of Recommendation A Strong consensus (97% agreement)
- 289

### 290 **Commentary**

Most studies that evaluated the clinical benefits of early EN in comparison with total PN used semi-elemental formulae while the recent studies were performed with polymeric formulae. In all studies both types of formulae were proven to be feasible, safe and welltolerated. One small RCT in 30 patients found that both formulae were safe and welltolerated (based on a visual analogue scale and number of stools per day) with some

296 clinical benefits for semielemental diets, including length of stay  $(23 \pm 2 vs. 27 \pm 1 days)$ 297 p = 0.006) and weight maintenance (51). On the other hand an indirect adjusted meta-298 analysis of Petrov et al. on 428 patients using PN as a reference treatment showed no 299 differences regarding tolerance, rate of infection and mortality between both formulae 300 (52). Finally, a second, more recent meta-analysis, including 15 trials (1376 301 participants), showed no evidence to support a specific enteral formula (53). 302 Nevertheless, a subgroup of patients with severe AP may have malabsorption and 303 therefore, semi-elemental diets could be of interest.

304

305 6. What route should be used for EN in patients with AP?

### 306 **Recommendation 7**

307 If EN is required in patients with AP, it should be administered via a nasogastric

308 tube. Administration via a nasojejunal tube should be preferred in case of

309 digestive intolerance.

310 Grade of Recommendation B – Strong consensus (95% agreement)

311

### 312 **Commentary**

Three RCTs compared nasojejunal with nasogastric support route in patients with severe AP (54-56) showed no differences regarding tolerance, complications rates and mortality. Four meta-analyses (57-60) conclude that nasogastric tube feeding is feasible, safe and well-tolerated and, compared with nasojejunal tube feeding, does not increase complication rate, mortality, refeeding pain recurrence or prolong hospital stay in patients with severe AP. Compared with nasojejunal tubes, nasogastric tubes are much easier to place, more convenient and cheaper. Nevertheless, about 15% of patients will

320	experience digestive intolerance, mostly because of delayed gastric emptying and gastric
321	outlet syndrome (57, 58) and in this situation, nasojejunal tube feeding is required.
322	Furthermore, potential bias arises from the small number of patients included in the
323	aforementioned trials and the use of different criteria to define severe AP.
324	
325	7. In patients with AP, when should PN be initiated?
326	Recommendation 8
327	PN should be administered in patients with AP who do not tolerate EN or who are
328	unable to tolerate targeted nutritional requirements, or if contraindications for
329	EN exist.
330	Grade of Recommendation GPP – Strong consensus (97% agreement)
331	
332	Commentary
333	The primary nutritional route in all patients with severe AP should be enteral, as this
334	
	route has been shown to have benefits over other regimens. However, PN is indicated in
335	route has been shown to have benefits over other regimens. However, PN is indicated in patients with severe AP who do not tolerate EN or who are unable to tolerate targeted
335 336	
	patients with severe AP who do not tolerate EN or who are unable to tolerate targeted
336	patients with severe AP who do not tolerate EN or who are unable to tolerate targeted requirements, or if there exists contraindication for EN overall. Complications of severe
336 337	patients with severe AP who do not tolerate EN or who are unable to tolerate targeted requirements, or if there exists contraindication for EN overall. Complications of severe AP, which may occur and represent a contraindication for EN, include bowel obstruction,
336 337 338	patients with severe AP who do not tolerate EN or who are unable to tolerate targeted requirements, or if there exists contraindication for EN overall. Complications of severe AP, which may occur and represent a contraindication for EN, include bowel obstruction, abdominal compartment syndrome, prolonged paralytic ileus and mesenteric ischemia
336 337 338 339	patients with severe AP who do not tolerate EN or who are unable to tolerate targeted requirements, or if there exists contraindication for EN overall. Complications of severe AP, which may occur and represent a contraindication for EN, include bowel obstruction, abdominal compartment syndrome, prolonged paralytic ileus and mesenteric ischemia (61). Similar to critically ill patients with other diseases, approximately 20% of patients

	Journal Pre-proof
343	8. How should medical nutrition be provided in case of necrosectomy (endoscopically or by
344	minimally invasive surgery) in patients with severe AP?
345	Recommendation 9
346	Oral food intake in patients undergoing minimally invasive necrosectomy is safe
347	and feasible and should be initiated in the first 24 hours after the procedure, if the
348	clinical state (hemodynamic stability, septic parameters, gastric emptying) of the
349	patient allows it.
350	Grade of Recommendation GPP – Strong consensus (95% agreement)
351	
352	Recommendation 10
353	In patients undergoing minimally invasive necrosectomy who are unable to be fed
354	orally, EN is indicated via nasojejunal as preferred route.
355	Grade of Recommendation B – Strong consensus (91% agreement)
356	
357	Recommendation 11
358	PN is indicated in patients undergoing minimally invasive necrosectomy who do
359	not tolerate EN or who are unable to tolerate targeted nutritional requirements,
360	or if there exist contraindications for EN.
361	Grade of Recommendation GPP – Strong consensus (94% agreement)
362	
363	Commentary

364 Approximately 10-20% of patients with AP will develop necrosis of the pancreas and/or 365 peripancreatic tissue (ANP) (1, 2). These patients with ANP have moderate or severe 366 forms of AP, and a higher risk for development of multiple organ failure, secondary 367 infection of the necrosis, and death (62). After proven benefits of the "step-up" 368 (minimally invasive approach) over the open approach for the treatment of ANP (63), 369 minimally invasive techniques have been used extensively (64). Furthermore, the Dutch 370 Pancreatitis Study Group recently showed a lower rate of pancreatic fistula and better 371 cost benefits of endoscopic over surgical step-up approach for infected necrotizing 372 pancreatitis (65). Unfortunately, to date there are no published data on nutritional 373 support in patients with AP treated by the minimally invasive approach. In the aforementioned trial (65), all patients received oral nutrition, if tolerated. If this was not 374 375 tolerated, a nasojejunal feeding tube was introduced and EN was started. If 376 gastrointestinal feeding was contraindicated, the patient received PN. No specific data 377 were reported regarding nutrition-related outcomes.

378 In the RCT by Bakker *et al.* (48), there was no superiority of early (first 24 hours) 379 nasojejunal tube feeding when compared with an oral diet after 72 hours in reducing the 380 rate of infection or death in patients with predicted severe AP. In this trial interventional 381 procedures due to necrotizing pancreatitis included percutaneous catheter drainage, 382 endoscopic transgastric drainage or necrosectomy and surgical necrosectomy (without 383 information on the type of surgery performed – minimally invasive or open approach). 384 The authors did not find any difference in the number of patients who underwent 385 interventions between groups (24 percutaneous drainages in early EN group vs. 46 in 386 the on demand feeding group, p = 0.13; eight endoscopic transgastric drainage or necrosectomy in the early EN group vs. six in the on-demand feeding group, p = 0.53; 387 and three surgical necrosectomy in the early EN group vs. seven in the on-demand 388

389 feeding group, p = 0.49). In this trial PN was not used, as it was not mentioned in the 390 feeding protocol of the study. In a retrospective series of 37 patients undergoing 391 laparoscopic transgastric necrosectomy, an oral food intake 24-48 hours after the 392 procedure was feasible and safe (66). In one prospective study on video-assisted 393 retroperitoneal debridement (VARD) the feeding regimen was reported but without 394 specified time of initiation and reasons for shifting oral nutrition to EN or PN (67). Forty 395 patients in that study were fed by nasojejunal tube as the preferred route when 396 tolerated; otherwise, PN was given (67). Therefore, based on small series, nasojejunal 397 feeding seems safe in patients having undergone minimally invasive necrosectomy. 398 Nevertheless, definitive data are missing.

399

9. How should medical nutrition (EN and PN) be provided in critically patients with severe
AP (intra-abdominal hypertension (IAH), abdominal compartment syndrome (ACS) with
need for open abdomen)?

403

#### 404 Recommendation 12

In patients with severe AP and intraabdominal pressure (IAP) < 15 mmHg early</li>
EN shall be initiated via nasojejunal, as the preferred route, or nasogastric tube.
IAP and the clinical condition of patients during EN shall be monitored
continuously.

409 Grade of Recommendation A – Strong consensus (91% agreement)

410

411 **Recommendation 13** 

412 In patients with severe AP and IAP > 15 mmHg EN should be initiated via 413 nasojejunal route starting at 20 mL/hour, increasing the rate according to the 414 tolerance. Temporary reduction or discontinuation of EN should be considered when IAP values further increase under EN. 415 416 Grade of Recommendation B - Strong consensus (94% agreement) 417 418 **Recommendation 14** 419 In patients with severe AP and IAP > 20 mmHg or in the presence of ACS, EN 420 should be (temporarily) stopped and PN should be initiated. Grade of Recommendation GPP - Strong consensus (94% agreement) 421 422 423 **Recommendation 15** 424 In patients with severe AP and open abdomen EN should be administered, at least 425 in a small amount. If required for achievement of nutritional requirements, 426 supplementary or total PN should be added. 427 Grade of Recommendation B - Strong consensus (97% agreement) 428 429 **Commentary** 430 The mortality of patients with severe AP who develop IAH/ACS during the course of the 431 disease rises from 25% up to 66% (68, 69). Energy expenditure in patients with AP is 432 increased by 1.49 (1.08 to 1.78)  $\times$  the predicted resting energy expenditure; 58% of

patients with severe AP have an increase in energy expenditure, approximate net
nitrogen loses are 20-40 grams per day, and proteolysis can be increased by 80% (70,
71). There are no data available regarding energy requirements in patients with both AP
and IAH/ACS, however, energy expenditure in such patients may be increased due to
several reasons (decreased splanchnic blood flow, acidosis and bacterial translocation)
(17, 72).

439 It has been clearly demonstrated that EN in patients with severe AP reduces mortality 440 and infectious complications, decreases organ failure and surgical intervention rate, has 441 a trend towards reduction of hospital stay, and is safer and more effective than PN (17). 442 Nevertheless, it has been reported that EN may increase intraluminal pressure with 443 subsequent elevation of IAP and development of severe complications (73, 74). 444 Therefore, it is recommended that EN should be administered with caution when IAP 445 reaches 15 mmHg and over (74). In an observational study, 274 patients with AP had 446 IAH and 103 developed ACS. The intolerance of EN was more frequent in patients with 447 grade III and IV IAH (n=105) and 62/105 (59%) required PN (75). In only one RCT 448 including 60 patients, comparing early with delayed EN in patients with IAH and severe AP, it was found that early EN had benefits in patients with IAP < 15 mmHg preventing 449 450 development of IAH. In patients with IAP above 15 mmHg abdominal distension was 451 more frequent in the early EN group. The group of patients with early EN experienced 452 feeding intolerance more often than patients in delayed EN group. However, early EN 453 did not increase IAP and was able to ameliorate clinical course of the disease (76). 454 Because the majority of patients with IAH have gastrointestinal symptoms and signs 455 (absence of bowel movements, abdominal distension, high gastric residual volume, etc.), 456 EN should be initiated via nasojejunal tube (77). From a practical point of view, in patients with severe AP and IAH the initiation of EN should be at 20 mL/hour, increasing 457

the rate according to the tolerance. The reduction of EN from higher rates to 20 mL/h
should be considered when IAP increases between 15 and 20 mmHg. In patients with
IAP above 20 mmHg or in the presence of ACS, EN should be (temporarily) stopped (74).
When it is impossible to meet nutritional goals with EN only, supplementary or total PN
should be considered.

463 A decompressive laparotomy (laparostomy) may be necessary in up to 74% of patients 464 who develop ACS during course of AP (72). Patients with an open abdomen are in a 465 hyper-catabolic state with high nitrogen losses and negative nitrogen balance. It has 466 been estimated that such patients have nitrogen loss of almost 2 g/L of abdominal fluid 467 output and, therefore, nutritional therapy in patients with an open abdomen is essential 468 (78). Several cohort studies reported that initiation and feeding by EN was feasible and 469 safe despite a relatively high rate of digestive intolerance, ranging from 48-67% (78-83). 470 Two studies concluded that that early EN in patients with an open abdomen resulted in 471 higher fascial closure rates, lower fistula rates, reduced nosocomial infections and lower 472 hospital costs (82, 83). In the multicenter analysis by Burlew et al., out of 597 with an 473 open abdomen patients, EN was successfully initiated in 39% (81). For the 307 patients without a bowel injury, logistic regression indicated that EN was associated with higher 474 475 fascial closure rates (OR 5.3; p < 0.01) decreased complication rates (OR, 0.46; p = 0.02), 476 and decreased mortality (OR 0.30; p = 0.01) (81).

477

478 10. Is there any role for immunonutrition (glutamine, antioxidants) in severe AP?

### 479 **Recommendation 16**

When EN is not feasible or contraindicated and PN is indicated, parenteral
glutamine should be supplemented at 0.20 g/kg per day of L-glutamine. Otherwise,
there is no role for immunonutrition in severe AP.

- 483 Grade of Recommendation B Strong consensus (94% agreement)
- 484

### 485 **Commentary**

486 An initial meta-analysis including eleven RCTs assessed the effect of antioxidants (five RCTs on glutamine and six on various other antioxidants) on the outcome of patients 487 488 with AP (84). Among patients with AP, antioxidant therapy resulted in a borderline significant reduction in hospital stay (mean difference 1.74; 95% CI 3.56 to 0.08), a 489 490 significant decrease in complications (RR 0.66; 95% CI 0.46 to 0.95) and a non-491 significant decrease in mortality rate (RR 0.66; 95% CI 0.30 to 1.46). Nevertheless, these 492 results were mostly attributed to the effect of glutamine. Recently, a Cochrane Review 493 assessed the effects of different pharmacological interventions including antioxidants in 494 patients with AP (85). Very low-quality evidence suggested that none of the 495 pharmacological treatments decreased short-term mortality in patients with AP.

496 Regarding glutamine, four meta-analyses have been published. A meta-analysis of ten 497 RCTs including 433 patients with severe AP revealed a significant decrease in the 498 incidence of infectious complications and mortality in the patient group with glutamine-499 enriched nutrition (86). Another meta-analysis of twelve RCTs (including 505 patients) 499 demonstrated a significantly reduced infection rate and mortality after glutamine 501 supplementation in patients with AP (87). In the subgroup analyses, only patients who 502 received total PN demonstrated a significant benefit in terms of study outcomes. Two

503 recently published meta-analyses showed beneficial effects of glutamine 504 supplementation in patients with AP in the terms of elevation of serum albumin 505 concentrations, decrease in serum concentrations of C-reactive protein, and reductions 506 in infectious complications, mortality and hospital stay (84, 88). Nevertheless, the risk of 507 bias of the included studies is important due to many reasons: (i) small sample size in 508 most of the studies, (ii) possible heterogeneity in disease severity and (iii) confounding 509 factors such as other interventions that may change outcome (drainage, debridement or 510 surgery).

511

512 11. Is there any role for probiotic use in severe AP?

### 513 Recommendation 17

- 514 **Probiotics cannot be recommended in patients with severe AP.**
- 515 **Grade of Recommendation 0 Consensus (89% agreement)**
- 516

### 517 **Commentary**

A meta-analysis of six RCTs including 536 patients revealed no significant benefit of probiotics on pancreatic infection rate, overall infection rate, operation rate, length of hospital stay and mortality (89). Significant heterogeneity was observed in the type, dose and treatment duration of probiotics in these trials. In one of these RCTs the patient group assigned to a particular combination of probiotic strains showed similar pancreatic infection rate but increased mortality when compared with the placebo group (90).

525

526 12. Is there any role for the use of oral enzyme supplementation in AP?

### 527 Recommendation 18

- 528 **Pancreatic enzymes should not be supplemented generally except in patients with**
- 529 obvious pancreatic exocrine insufficiency (PEI).
- 530 Grade of Recommendation B Strong consensus (97% agreement)
- 531

### 532 **Commentary**

There are only two RCTs with a total of 78 patients randomized to pancreatic enzyme 533 534 supplementation or placebo (91, 92). In the study by Kahl *et al.* 20 of the 56 patients 535 showed low fecal elastase values indicating PEI. Although the pancreatic enzyme 536 supplement group showed a tendency for better outcome this did not reach statistical significance (91). In the second small study by Patankar et al. there was also no 537 538 significant difference in laboratory or clinical outcomes (92). Therefore, no conclusion 539 can be drawn, but enzyme supplementation should be considered in patients with 540 proven or obvious exocrine insufficiency and malabsorption with steatorrhea.

541

### 542 II. Chronic pancreatitis

543

544 13. What are the risks of developing malnutrition in patients with CP?

545 **Statement 2** 

546 **Risk of malnutrition in CP is high and malnutrition is common in patients with CP**.

547 **Strong consensus (100% agreement)** 

548

### 549 **Commentary**

550 CP is a disease with progressive and irreversible inflammatory changes in the pancreas 551 that result in permanent structural damage with fibrosis, which can lead to abdominal 552 pain and to impairment of exocrine (pancreatic insufficiency) and often endocrine 553 function (4, 93-95).

554 Malnutrition is often a late, but important manifestation in the course of CP and depends 555 on the intensity and duration of the underlying disease. There are differences in the 556 onset of pancreatic insufficiency and malnutrition between patients with alcoholic and 557 idiopathic CP. The latency between onset of first symptoms and signs of CP, including 558 pain and malabsorption/malnutrition is between five to ten years in alcoholic, but 559 delayed in non-alcoholic pancreatitis (94).

Despite the inconsistency of the data there is an evident risk of malnutrition in patients with CP (95-97). According to a recent study medium or higher risk for malnutrition based on Malnutrition Universal Screening Tool (MUST) score of one or higher was found in 31.5% patients (98). Similarly, 26% underweight patients with a nutritional risk were identified in a study of outpatients with CP (99).

At the same time a recent prospective cohort study on 62 patients with CP and 66 controls showed that over half of the patients with CP were overweight or obese (100). Nevertheless, significant differences in handgrip strength were shown in patients with CP when compared with controls.

569 In patients with CP with moderate to severe weight loss, decreased lean body mass and 570 sarcopenia may lead to decreased functional capacity, which may have an impact on 571 quality of life (101, 102). In addition, PEI leads to the increased risk of developing 572 significant bone loss and severe osteoporosis (103, 104). A recent prospective study 573 (102) including 182 patients with CP showed that sarcopenia was present in 17% (74%) 574 of patients with CP had a BMI > 18.5 kg/m<sup>2</sup>). During follow-up, sarcopenia was 575 associated with an increased risk of hospitalization (OR 2.2; 95% CI 0.9 to 5.0; p = 0.07), 576 increased number of in-hospital days (p < 0.001), and reduced survival (HR 6.7; 95% CI 577 1.8 to 25.0; p = 0.005).

578

579 14. What are the causes of malnutrition in patients with CP?

### 580 Statement 3

581 Pancreatic insufficiency, abdominal pain, alcohol abuse, lower food intake,
582 diabetes mellitus and smoking are the main causes of malnutrition in CP.

583 **Strong consensus (97% agreement)** 

584

### 585 **Commentary**

Multiple risk factors for developing nutrient deficiencies and malnutrition co-exist in
patients with CP. First of all, pancreatic insufficiency (exocrine but also often endocrine)

588 can lead to maldigestion and malabsorption. Clinical signs of PEI include steatorrhea,

abdominal pain, weight loss and malnutrition (4). Recent data showed endocrine insufficiency and/or clinical steatorrhea in 41% and 36% of 809 patients (93). Moreover, increased resting energy expenditure can be seen in up to 50% of patients with CP, thus leading to a negative energy balance and malnutrition (105). Furthermore, abdominal pain, which is frequent in patients with CP, can lead to suboptimal dietary intake and also contribute to malnutrition (4).

595 Tobacco is an independent risk factor for CP, and can also be a disease modifier, acting

in synergy with alcohol intake, and therefore, adds to the nutritional risk factors (93).

597

598 15. Which diagnostic tests are preferred to assess nutritional status in patients with CP?

### 599 Recommendation 19

600 Nutritional status should be assessed according to symptoms, organic functions,

anthropometry, and biochemical values. Solely BMI should not be used, because it

602 does not register sarcopenia in the obese patient with CP.

603 Grade of Recommendation GPP – Strong consensus (97% agreement)

604

### 605 **Commentary**

Studies assessing malnutrition have identified many biochemical factors that are associated with malnutrition (106, 107) and prevalence studies show a diverse presentation of malnutrition. Olesen *et al.* identified that 26% of patients with CP were underweight in a cross-sectional study of 166 patients with CP (99), whereas Duggan *et al.* highlighted that over half of the patients in their prospective controlled cohort study (n = 128) fell into the overweight/obese category using BMI (100). However, patients had lower muscle stores and reduced functional status assessed using hand-grip

613	strength than healthy controls. Consequently, BMI alone is not considered an adequate
614	method of assessing nutritional status. Percentage weight loss is considered a more
615	reliable indicator of the onset of malnutrition and is associated with an increased risk in
616	the surgical setting (108).
617	Consequently, nutritional assessment should allow for detection of simple malnutrition,
618	sarcopenia and micronutrient deficiencies in addition to identifying symptoms that may
619	predispose patients to worsening malnutrition (Error! Reference source not found.5).
620	
621	16. What is the frequency of screening for micro- and macro-nutrient deficiencies in
622	patients with CP?

### 623 Recommendation 20

- 624 Patients should undergo screening for micro- and macronutrient deficiencies at
- 625 least every twelve months; screening may need to occur more frequently in those
- 626 with severe disease or uncontrolled malabsorption.
- 627 Grade of Recommendation GPP Strong consensus (100% agreement)

628

### 629 **Commentary**

Patients with CP are at high risk of malnutrition, both in terms of body weight and altered body composition (100). This has an impact on quality of life (99) and survival after surgery (109, 110). Nutritional intervention can improve nutritional markers and is associated with reduced pain (111) and, therefore, routine screening to trigger nutritional intervention should be undertaken. Deficiencies in micronutrients (vitamin B12, folic acid, vitamin A, D and E, zinc, selenium, iron) are well documented in patients with exocrine insufficiency, these are diverse in presentation with some studies

637 reporting biochemical deficiencies (100, 103, 112) and case reports document clinical 638 manifestations including night blindness (113, 114). However, there are no data 639 recommending the frequency of assessment or the likely timing of progression to 640 micronutrient deficiency. As clinical manifestation of deficiency represents a late 641 presentation, routine screening should be implemented to detect early signs of 642 deficiency.

- 643
- 644 17. What recommendations regarding diet and intake of fat, carbohydrates and proteins
- 645 should be given in patients with CP?
- 646 <u>Statement 4</u>
- 647 Patients with CP do not need to follow a restrictive diet.
- 648 Strong consensus (94% agreement)
- 649
- 650 **Recommendation 21**
- 651 **CP** patients with a normal nutritional status should adhere to a well-balanced diet.
- 652 Grade of Recommendation GPP Strong consensus (94% agreement)
- 653
- 654 **<u>Recommendation 22</u>**
- 655 Malnourished patients with CP should be advised to consume high protein, high-
- 656 **energy food in five to six small meals per day.**
- 657 Grade of Recommendation GPP Strong consensus (94% agreement)
- 658

### 659 **Recommendation 23**

- 660 In patients with CP, diets very high in fiber should be avoided.
- 661 **Grade of Recommendation B Strong consensus (91% agreement)**

662

- 663 Statement 5
- 664 In patients with CP, there is no need for dietary fat restriction unless symptoms of

665 steatorrhea cannot be controlled.

666 Strong consensus (100% agreement)

667

### 668 Commentary

There are very little data to suggest the optimal dietary management for patients with CP. Historically, patients were encouraged to have a low-fat diet, and studies in the Netherlands suggest 48-58% of patients still restrict dietary fat (104, 115). International guidelines are consistent in their recommendation that patients should have a balanced diet and avoid fat restriction (4, 116-119).

674 The role of dietary fat has been examined in small studies, suggesting an improvement 675 in dyspeptic symptoms in patients with very mild pancreatic disease who did not 676 consume alcohol regularly when a very low fat diet was consumed (< 20 g fat per day) 677 (120) and patients who consumed a higher fat diet were thought to be diagnosed at a 678 younger age, and had an increased probability of continuous abdominal pain (121) 679 suggesting a potential role in the initial development of CP. However once CP was 680 diagnosed, there was no difference in severity or complications of disease. An RCT 681 comparing dietary counselling and nutritional supplements in a cohort of 60

682 malnourished patients with CP found that nutritional intervention in which 33% of 683 energy was derived from fat was well tolerated (111). Improvements in nutritional 684 status and pain control were observed in patients receiving nutritional intervention and 685 the authors did not report any adverse events (111).

Patients consuming very high fiber diets reported increased flatulence, and increased fecal weight and fat losses were observed in a small trial (n = 12) in patients with CP. This study suggested that very high fiber diets may inhibit pancreatic enzyme replacement therapy, thus resulting in malabsorption. Thus, very high fiber diets are not recommended in this patient group (122).

691

692 18. Are oral supplements, with or without medium-chain triglycerides (MCTs), indicated in693 patients with CP?

### 694 Recommendation 24

Oral nutritional supplements (ONS) should be prescribed to undernourished
patients only if oral nutrition is insufficient for reaching the calorie and protein
goals.

698 Grade of Recommendation GPP – Strong consensus (100% agreement)

699

### 700 Recommendation 25

701 If adequate enzyme supplementation and exclusion of bacterial overgrowth has

not led to relief of malabsorption and its accompanying symptoms, ONS with MCT

703 can be administered.

704 **Grade of Recommendation 0 – Strong consensus (97% agreement)** 

### 705

### 706 Commentary

707 Very few studies have investigated the benefit of ONS in patients with CP. Eighty percent
708 of patients can be treated with diet and enzyme supplementation, the rest need oral
709 supplementation (96).

ONS can be of benefit in undernourished patients with CP, especially if the caloric and
protein goals cannot be reached with normal meals and counselling. ONS are a simple
way to improve oral intake, but long-term compliance may be a problem.

There are no RCTs investigating the relative efficacy of different formulae (e.g. standard or peptide-based with MCT). However, in the presence of PEI, enteral formulae consisting of pre-digested products and a mixture of long chain fatty acids and MCT would seem, theoretically, to have potential advantage. MCTs are less dependent on lipase activity for their absorption (123).

A reduction in oral fat intake or the replacement of dietary fat with MCT risks a 718 reduction in energy intake and, therefore, a negative energy balance. MCTs have an 719 720 unpleasant taste and are associated with adverse effects like cramps, nausea, and diarrhea. Up to now, studies have not shown any clear benefit of MCTs over standard 721 722 long-chain triglycerides when used in combination with enzyme supplementation (123, 723 124). One RCT investigated the efficacy of ONS in patients with CP and severe 724 malnutrition (111). Dietary counselling achieved equal results compared with the use of 725 a commercial supplement enriched with MCTs. Both groups also received enzyme 726 supplementation and so it is not possible to explain the additional gain from dietary 727 MCTs over enzyme supplementation.

728 If MCTs are being considered, their dose should be increased slowly depending on the729 patient's tolerance.

730	
731	19. When is micronutrient supplementation indicated in patients with CP (not including
732	osteoporosis prevention)?
733	Recommendation 26
734	Fat-soluble (A, D, E, K) and water-soluble (vitamin B12, folic acid, thiamine)
735	vitamins as well as minerals such as magnesium, iron, selenium and zinc should
736	be monitored (if available) and administered if low concentrations are detected
737	or if clinical signs of deficiency occur. Supplementation should be proposed to
738	patients with known malabsorption.
739	Grade of Recommendation GPP – Strong consensus (95% agreement)
740	
741	Commentary
742	The reported prevalence of deficiency of fat-soluble vitamins is 3–14.5% for vitamin A
743	deficiency (100, 103, 125), 58–77.9% for vitamin D deficiency (100, 103, 125, 126), 9-
744	24% for vitamin E deficiency (100, 103, 106, 125, 126) and 13–63% for vitamin K
745	deficiency (100, 103, 125, 126). In a prospective controlled cohort study of 128 subjects
746	and 66 age/gender-matched controls, 14.5% and 24.2% were deficient in vitamins A
747	and E, respectively, with a significant difference compared with controls. Nineteen
748	percent of patients had excess serum vitamin A concentrations (100). This must be
749	taken in account and a blind supplementation of all fat-soluble vitamins for all patients
750	with CPs is not advised.
751	Deficiencies of water-soluble vitamins in patients with CP are less frequent. A recent
752	study with 301 patients with CP and 266 controls showed that patients with CP had

significantly lower concentrations of vitamins A, D and E, but no difference regarding

754	vitamin B12 (103). Similarly, another cohort study of 114 patients with CP (33% with
755	exocrine failure) did not show any significant deficiencies of vitamin B12 (0%) and folic
756	acid (2.2%) (127).

- 757 Thiamine deficiency secondary to concomitant alcoholism must be considered (106).
- 758 Minerals and trace elements deficiencies have been reported in patients with CP in some
- 759 case-control studies. The results are conflicting. Lower concentrations of zinc, selenium
- 760 (106) and magnesium (127) have been observed. Furthermore, low magnesium
- 761 concentrations seemed to correlate with exocrine failure (127).
- 762
- 763 20. When is EN indicated in patients with CP and how should it be administered?
- 764 Recommendation 27
- 765 EN should be administered in patients with malnutrition who are not responding
- 766 to oral nutritional support.
- 767 Grade of Recommendation GPP Strong consensus (100% agreement)
- 768
- 769 Recommendation 28
- 770 EN should be administered via the nasojejunal route in patients with pain, delayed
- 771 gastric emptying, persistent nausea or vomiting and gastric outlet syndrome.
- 772 Grade of Recommendation GPP Strong consensus (100% agreement)

773

774	Recommendation 29
775	Long-term jejunostomy access (percutaneous endoscopic gastrostomy with
776	jejunal extension (PEG-J) or direct percutaneous endoscopic jejunostomy (DPEJ)
777	or surgical jejunostomy) can be used in those requiring EN for more than 30 days.
778	Grade of Recommendation GPP – Strong consensus (97% agreement)
779	
780	Recommendation 30
781	Semi-elemental formulae with medium chain triglycerides can be used if standard
782	formulae are not tolerated.
783	Grade of Recommendation GPP – Strong consensus (94% agreement)
784	
785	Recommendation 31
786	Pancreatic enzymes should be supplemented in patients requiring EN, if signs of
787	exocrine failure manifest.
788	Grade of Recommendation GPP – Strong consensus (100% agreement)
789	
790	Commentary
791	Oral nutritional support with dietary counselling is usually sufficient to improve
792	nutritional status in patients with CP (111). EN is indicated in approximately 5% of
793	patients with CP (97). Regarding indications and outcomes of EN in these patients,
794	evidence is based on few cohort studies and RCTs are generally lacking (4).
795	Four retrospective series have shown the benefits of EN in patients with CP regarding
796	weight gain and pain control (128-131). Two of them included 58 (129) and 50 patients

(131) respectively, in whom a naso-jejunal tube was placed. Long-term access with PEGJ or DPEJ was used in 57 (128) and 58 patients (130). All studies showed that this type
of nutritional support was safe and effective in patients with CP, even in case of gastric
outlet syndrome (130, 131).

There is limited high quality evidence for the composition of enteral formulae in patients with CP. However, there is a rationale that semi-elemental enteral formulae with MCTs are more adapted for jejunal nutrition, compared with polymeric formulae (132). In two of the aforementioned studies (129, 131), semi-elemental formulae were used with good digestive tolerance. Nevertheless, the cost of these feeds is higher and data on cost-effectiveness are also lacking.

807 In patients with exocrine failure, who do not improve with semi-elemental formulae,

808 pancreatic enzymes can be administered with the formula (133). This involves opening

809 the capsules and suspending the enzyme microspheres in thickened acidic fluid (such as

810 the mildly thickened or "nectar-thick" fruit juice used for dysphagia) for delivery via the

811 feeding tube.

812

813 21. When is PN indicated in patients with CP and how should it be administered?

814 **<u>Recommendation 32</u>** 

815 **PN may be indicated in patients with gastric outlet obstruction and in those with** 

816 complex fistulating disease, or in case of intolerance of EN.

817 Grade of Recommendation GPP – Strong consensus (100% agreement)

818

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oun	lai			U.

## 819 **Recommendation 33**

- 820 For PN the preferable route is central venous access.
- 821 Grade of Recommendation GPP Strong consensus (100% agreement)
- 822

## 823 Commentary

824 PN is infrequently uses in patients with CP (4, 97). EN preserves immune function and 825 mucosal architecture and decreases the possibility for hyperglycemia while PN also 826 increases the risk of catheter-related infections and septic complications (96, 119). PN 827 is, therefore, only indicated when it is impossible to use EN (e.g. presence of gastric 828 outlet obstruction, the need for gastric decompression, when it is impossible to 829 introduce a tube into the jejunum, or a complicated fistula is present) or if requirements 830 are only partly reached by EN. PN is mainly administered over a short-term period and 831 long-term studies are lacking. In this case, a standard nutritional solution should be 832 administered via central venous access such as a peripherally inserted central catheter. 833 Contraindications to PN do not differ from general contraindications to medical 834 nutrition.

835

836 22. What are the indicators for starting pancreatic enzyme replacement therapy (PERT) in837 patients with CP?

## 838 **Recommendation 34**

When PEI is diagnosed through clinical signs and symptoms and/or laboratory
tests of malabsorption, PERT shall be initiated. An accurate nutritional
assessment is mandatory to detect signs of malabsorption.

# 842 Grade of Recommendation A – Strong consensus (100% agreement)

843

### 844 **Commentary**

PEI is defined as an insufficient secretion of pancreatic enzymes (acinar function) 845 and/or sodium bicarbonate (ductal function) (4). Diagnosis of PEI can be challenging in 846 847 practice because pancreatic function and secretion are not solely reliant on the quantity 848 or quality of pancreatic tissue (134) but also depend on complex pancreatic stimulatory 849 mechanisms (135). Moreover, different PEI biomarkers and their threshold values have 850 been used in the current literature (136). For these reasons a wide range (from 22% to 94%) of prevalence rates for PEI among patients with CP has been reported (98, 106, 851 852 137-146).

The most frequent clinical sign of PEI is steatorrhea (147), defined as presence of fat in the stool, and associated generally with flatulence, bloating, dyspepsia, urgency to pass stools, and cramping abdominal pain. In a recent systematic review, including 14 studies on pancreatic enzyme supplementation in patients with CP, the criteria for the diagnosis of PEI were the measurement of the coefficient of fat absorption with a threshold < 80% or the fecal fat absorption less than 7 - 15 g of fat per day (136).

Overt steatorrhea is not expected unless there is severe or decompensated PEI (i.e. when secretion of pancreatic lipase is less than < 10% of normal). However, the absence of overt steatorrhea is not always an indicator of adequate absorption and nutritional status. PEI is consistently associated with biochemical and clinical signs of malnutrition (148). Management of PEI involves replacing the inadequate pancreatic enzymes, which should be used to maintain weight and improve the symptoms of maldigestion (149).

Awareness of PEI among many physicians is poor outside of referral centers and especially among physicians in primary care (115). Consequentially, patients who

867 present with symptoms of PEI may be overlooked or advised to adopt inappropriate 868 dietary restrictions in an attempt to control the symptoms. A study identified that the 869 primary unmet patient need was the difficulty in managing gastrointestinal symptoms, 870 diet, and digestion; indeed, many of these patients and caregivers cited delays in dietary 871 assessment and initiation of PERT causing additional distress that could have been 872 prevented (150). Untreated PEI has also a deleterious impact on the quality of life of 873 patients (151). As the quantitative measurement of fecal fat is often omitted, it is 874 recommended that enzyme replacement is started when clinical signs of malabsorption, 875 or anthropometric and/or biochemical signs of malnutrition are present (96, 127, 152-876 154). Symptoms include weight loss, alteration of body compartments at bioimpedance 877 analysis, and low nutritional markers (albumin, cholinesterase, prealbumin, retinol-878 binding protein, and magnesium) (127). Although it is assumed that steatorrhea is the 879 most important clinical manifestation of PEI, several studies have shown reduced 880 absorption of fat-soluble vitamins even in patients with mild to moderate PEI (155-158). Non-alcoholic fatty liver disease (NAFLD) is also a poorly recognized complication of 881 882 PEI. The mechanisms underlying NAFLD in PEI is different from NAFLD associated with 883 metabolic syndrome, because it is mainly due to malabsorption of essential amino acids 884 such as choline which leads to a decrease in plasma concentrations of apoprotein B 885 (159), a major component of very-low-density lipoprotein.

886

887 23. What are the enzyme preparations of choice for PERT?

888 **Recommendation 35** 

889 pH-sensitive, enteric-coated microspheres pancreatic enzyme replacement
890 preparations shall be used for treating PEI.

#### 891 Grade of Recommendation A – Strong consensus (100% agreement)

892

# 893 Commentary

There are multiple pancreatic enzyme replacement preparations that are now licensed around the world. All are of porcine origin and contain, with varying concentrations and mixtures, pancreatic lipase, amylase, protease, and other pancreas-derived proteins and nucleic acids. Several factors affect the efficacy of pancreatic enzyme supplementation: (a) mixture with meal; (b) gastric emptying with meal; (c) mixing with chyme and bile acids and rapid release of enzymes in duodenum (160).

Nowadays, most of the pancreatic enzyme preparations are formulated as pH-sensitive, enteric-coated, capsules containing microspheres or tablets that protect the enzymes from gastric acidity and allow them to disintegrate rapidly at pH > 5.5 in the duodenum (160, 161). Non enteric-coated, conventional powder or tablet formulations have been abandoned because they are less effective in treating PEI as pancreatic enzymes are partially inactivated by pepsin and gastric acidity (162).

906 The efficacy of these more recent formulations has been demonstrated in several recent 907 studies (163-166) and in a recent meta-analysis (136). A Cochrane review on the 908 efficacy of pancreatic enzyme preparations in patients with pancreatic insufficiency 909 demonstrated a higher efficacy for enteric-coated microspheres compared with enteric-910 coated tablets (167). Mini-microspheres 1.0 - 1.2 mm in diameter seem to be associated 911 with higher therapeutic efficacy compared with 1.8 - 2.0 mm microspheres that still 912 have an optimal therapeutic action (168). Another trial compared two enteric-coated 913 pancreatic enzyme preparations. One moisture-resistant, formulated to contain between 914 90% to 110% labeled lipase content over the shelf life of the product and the other 915 potentially unstable in the presence of moisture and degradable over time. The

916	characteristics of the moisture-resistant formulation should have allowed more accurate
917	dosing, both providing more predictable therapeutic effects and reducing the risk of
918	overdose, which is assumed as a potential risk factor for fibrosing colonopathy. The
919	results suggested a comparable efficacy and safety in patients with cystic fibrosis for the
920	treatment of PEI (169).
921	
922	24. How should enzyme supplementation be administered?
923	Recommendation 36
924	Oral pancreatic enzymes should be distributed along with meals and snacks.
925	Grade of Recommendation B – Strong consensus (100% agreement)
926	
927	Commentary
928	The efficacy of pancreatic enzyme supplements presupposes the mixing of enzymes and
929	chyme (161). While one study evaluating the impact of the scheduling of PERT

930 administration on fat malabsorption suggested the optimal timing of administration was 931 during or after meals, no significant difference was observed when patients took PERT 932 immediately before meals (170). In practice, although many patients prefer to take 933 PERT at the beginning of meals, they should be encouraged to spread the capsules out 934 over a meal when using multiple capsules or with larger meals (162, 170). If the patient 935 is taking the older preparations of pancreas powder, they should take about a third of 936 the dose immediately before, one third during, and one third immediately after the meal. 937 This concerns only meals and snacks that contain fat (e.g. not for fruit).

938

939 25. What is the optimal dosage of enzyme supplementation?

#### 940 **Recommendation 37**

The posology aims at individual needs and depends on the severity of the disease
and the composition of the meal. In practice, a minimum lipase dose of 20,000 50,000 PhU (based on the preparation) shall be taken together with main meals,
and half that dose with snacks.

945 Grade of Recommendation A – Strong consensus (100% agreement)

946

# 947 **Commentary**

948 The dosage recommended depends on the patient's clinical response, but the dosage and
949 dosing will need to be monitored carefully, as well as altered, depending on patient's
950 food intake/pattern of eating, method of cooking, portion sizes, and disease evolution.

For the digestion of a normal meal a minimum activity of 30,000 IU of naturally secreted pancreatic lipase is required. The recommended initial dose is about 10% of the physiologically secreted dose of lipase after a normal meal (171). Since 1 IU of naturally secreted lipase equals 3 PhU in commercial preparations, the minimum amount of lipase needed for digestion of a normal meal is 90,000 PhU (endogenous plus orally administered lipase).

The results of several RCTs have proven the efficacy of pancreatic enzyme replacement therapy with enteric-coated mini-microspheres at a dose ranging from 40,000 - 80,000 PhU of lipase per main meal, and half dose per snack (165, 166, 170, 172-174). Studies evaluating enteric-coated microspheres have shown a similar efficacy for doses ranging from 10,000 - 40,000 PhU of lipase per meal, indicating the lack of a dose-response relationship with these preparations (175, 176).

Dose escalation may be warranted according to response. In adults there is no upper limit to dosing, as there is no risk of overdose because pancreatic enzymes exceeding the needs are eliminated through stools. Caution for dosage should be placed in children in whom colonic strictures have been described after high dose of the enteric coated, delayed release preparations (177).

968

969 26. How should the efficacy of enzyme supplementation be evaluated?

#### 970 Recommendation 38

971 The efficacy of PERT should be evaluated by the relief of gastrointestinal 972 symptoms and the improvement of nutritional parameters (anthropometric and 973 biochemical). In patients who do not respond, the evaluation should be extended 974 to pancreatic function tests (fecal fat excretion or <sup>13</sup>C-MTG-breath test).

975 Grade of Recommendation B – Strong consensus (97% agreement)

976

#### 977 **Commentary**

978 The aforementioned recent meta-analysis including 14 RCTs (136) showed that PERT increased the coefficient of fat absorption, as well as improved gastrointestinal 979 980 symptoms, compared with baseline or placebo. Two open label extensions up to one 981 year from RCTs included in the meta-analysis demonstrated significant improvement in 982 nutritional parameters and weight (164, 178). A review of reported data (106) as well as 983 the recent guidelines on the therapy for CP (4) support the use of nutritional parameters 984 as an optimal way to assess the efficacy of PERT. Dietary intake and nutritional status should be monitored regularly to maximize patient compliance and specialist dietetic 985 986 assessment sought in patients with underlying malnutrition (179).

987 In patients who do not respond, pancreatic function tests (136) while on PERT can 988 monitor effectiveness. <sup>13</sup>C-MTG-breath test is a useful method that can replace the 989 somewhat cumbersome fecal fat excretion tests and can be used for patients on PERT 990 (180).

991

992 27. What should be done in cases of unsatisfactory clinical response?

## 993 Recommendation 39

In case of unsatisfactory clinical response, PERT dosage should be increased or a
protein pump inhibitor (PPI) should be added. If these methods fail, other causes
of malabsorption such as small intestinal bacterial overgrowth (SIBO) should be
excluded.

998 Grade of Recommendation B – Strong consensus (97% agreement)

999

#### 1000 **Commentary**

The recommended dose of 20,000 - 50,000 PhU with main meals has been shown to improve symptoms in more than half the patients (136). Dose escalation may be warranted according to response. In adults there is no upper limit to dosing, as there is no risk of overdose because pancreatic enzymes exceeding the needs are eliminated in the stool. Caution for high PERT dosage should be exercised in children, in whom colonic strictures have been described after high dose of the enteric coated, delayed release preparations (177).

1008 The inhibition of gastric acid secretion by PPIs can lead to a significant improvement 1009 and even normalization of fat digestion in patients with an incomplete response to 1010 PERT, as shown in a prospective cohort study of 21 patients with CP (43% had an initial

incomplete response to PERT, and 29% normalized their function after addition of a
PPI) (181). Nevertheless, a review including 34 clinical trials failed to show
improvement in the efficacy of PERT with PPI or histamine-2 receptor antagonists
(182). It is noteworthy that the populations included and the therapeutic schemes were
very heterogeneous, therefore, suggesting significant bias.

SIBO can also explain persistent symptoms. A recent prospective case-control studyrevealed that SIBO was present in 15% of patients with CP whereas no healthy control

1018 was tested positive by means of a fasting glucose hydrogen breath test (183).

1019

1020 28. Does the surgical technique for treating CP affect PERT and nutritional status?

# 1021 Recommendation 40

1022 Long-term PERT and nutritional status are similarly affected by all surgical
1023 procedures. Tissue-preserving procedures shall be preferred.

1024 Grade of Recommendation A – Strong consensus (100% agreement)

1025

#### 1026 **Commentary**

Surgical intervention is effective in carefully selected patients. Common indications for
surgical intervention in CP include poorly controlled pain, duodenal, biliary and
pancreatic duct obstruction, and suspicion of cancer (184).

1030 Surgery for CP can be broadly classified into three categories: drainage procedures,

1031 partial pancreatic resection including or not the duodenum, and total pancreatectomy.

1032 Recently, Kamper *et al.* (185), reviewed all the available techniques in detail. In drainage

- 1033 procedures a dilated pancreatic duct is cut open and anastomosed to the proximal
- 1034 jejunum. The most common drainage procedures are the modified Puestow procedure,

also known as lateral pancreatico-jejunostomy, and the Frey procedure, which in
addition to a pancreaticojejunostomy includes coring of the pancreatic head. In patients
with persistent inflammation of the pancreatic head without upstream ductal dilatation,
a resective surgery such as a classic pancreaticoduodenectomy or a duodenumpreserving head resection (Beger procedure) can be performed.

Theoretically, the type of procedure may deeply affect short- and long-term nutritional
outcomes, since the extension of the parenchyma resection, as well as the preservation
of the duodenum and bile natural transit, and pancreatic secretion may represent key
factors for endocrine and exocrine functions (186, 187).

Meta-analyses showed better postoperative pain relief and improved quality of life with the Beger procedure compared with conventional pancreaticoduodenectomy (188, 189). However, the studies included had a high grade of heterogeneity and a recent large prospective large RCT showed no significant difference between procedures in the longterm nutritional status, quality of life, and preservation of the exocrine pancreatic function (190).

A 2015 meta-analysis of 23 studies compared outcomes of the Frey procedure with pancreaticoduodenectomy and the Berger procedure (191). Short-term quality of life and pancreatic function outcomes were more favorable in patients who had the Frey procedure than in those who had pancreaticoduodenectomy. Long-term follow-up data from an RCT comparing the Frey and Berger procedures for CP showed no significant difference in endocrine or exocrine insufficiency more than a decade after surgery 1056 (192).

1057

1058 29. What is the risk of developing osteoporosis or osteopenia in patients with CP?

47

### 1059 **Statement 6**

- 1060 Patients with CP are at risk for osteoporosis (almost one out of four) and at high
- 1061 risk (about two out of three), for osteopathy (either osteoporosis or osteopenia).
- 1062 **Strong consensus (97% agreement)**
- 1063

## 1064 **Commentary**

Osteoporosis is characterized by structural deterioration of bone tissue and low bone 1065 1066 mass, leading to bone fragility and increased risk of fracture (193). Osteoporosis and osteopenia are defined by the World Health Organization according to T-scores (a T-1067 1068 score between -1.0 and -2.5 standard deviations is defined as osteopenia; a T-score 1069 below 2.5 standard deviations is defined as osteoporosis), T-scores compare bone 1070 density values with those of young adults (peak bone mass) (194). Osteoporosis and 1071 osteopenia can also be defined according to Z-score (Z-score < -1 defined as osteopenia, 1072 Z-score < -2 defined as osteoporosis). The Z-scores represents gender- and age-matched 1073 controls for the evaluation of secondary osteoporosis, they are usually used in 1074 premenopausal women, men under the age of 50, and in children (195).

1075 A systematic review and meta-analysis including ten studies applied the definition in 1076 accordance with the T-scores in eight and the Z-scores in two studies. It revealed that, 1077 based on the random-effects model of the total 513 patients with CP included, a pooled 1078 prevalence rate of osteoporosis of 24.3% (95% CI 16.6 to 32.0%) and osteopathy (either 1079 osteoporosis or osteopenia) of 65% (95% CI 54.7 to 74.0%) (196). Two of the included 1080 studies revealed osteoporosis rate for controls respectively 8.6 and 10.2%. All the 1081 included studies had relatively small sample sizes (< 100) and considerable 1082 heterogeneity; therefore, subgroup analyses were not acquiescent. Certain patterns

were, however, evident from the studies included, like an association between pancreatic enzyme insufficiency and lower bone mineral density. On the contrary, the available data failed to show direct associations between serum vitamin D concentrations and low bone mineral density. These data suggest that vitamin D deficiency is not the sole driver of bone demineralization, other factors that may be of importance for premature bone demineralization in CP are heavy smoking, low physical activity, and chronic inflammation (197).

1090 The important clinical endpoint of osteoporosis is bone fracture. Two large 1091 retrospective studies shed light on this regarding patients with CP. The first is a cohort 1092 database study, examining patients with CP at a single tertiary care center. A total of 1093 3,192 patients with CP and 1,436,699 controls were included in the study. The fracture 1094 prevalence (patients with fracture per total patients) was 1.1% in controls 1095 (16,208/1,436,699) and 4.8% in patients with CP (154/3192); in comparison Crohn's 1096 disease revealed a risk of 3.0% (182/6057); liver cirrhosis 4.8% (805/16,658) and celiac disease 5.0% (74/1480) (198). 1097

The second, a Danish retrospective cohort study including 2594 patients with CP
revealed an adjusted hazard ratio for any fracture of 1.7 (95% CI 1.6 to 1.8) (199).
Patients with CP receiving PERT for fat malabsorption had a lower risk of fractures than
other CP patients (HR 0.8; 95% CI 0.7 to 0.9).

1102

1103 *30. What methods should be used to identify patients who are at risk?* 

## 1104 **Recommendation 41**

Dual-energy X-ray absorptiometry (DXA) shall be used to identify patients with CP
with osteopathy.

#### 1107 Grade of Recommendation A – Strong consensus (100% agreement)

1108

## 1109 **Commentary**

1110 The American College of Radiology aims to rate the appropriateness of several radiological modalities for specific patient populations. Although they do not mention CP 1111 1112 explicitly, they do state that in premenopausal females and males 20 - 50 years of age 1113 with malabsorption, DXA of the lumbar spine and hip(s) or distal forearm is usually an 1114 appropriate diagnostic modality to identify low bone mineral density (200). It is not yet 1115 well defined when and to whom these tests should be offered in patients with CP. 1116 However, there are recommendations from the American Gastroenterological 1117 Association on the detection of osteoporosis in other gastrointestinal diseases: 1118 recommending that patients with at least one additional osteoporosis risk factor should 1119 undergo initial screening with DXA (201). This recommendation was specifically for 1120 inflammatory bowel disease, celiac disease, and post-gastrectomy patients. The recently 1121 published HaPanEU guidelines on CP argued that bone density testing by DXA should be 1122 extended to patients with CP with an additional risk; post-menopausal women, those 1123 with previous low-trauma fractures, men over 50 years and those with malabsorption 1124 (4). They further stated that considering the associated morbidity and cost of bone 1125 fractures when prevention is within range (202), a baseline bone density assessment for 1126 all patients with CP may be worth considering.

1127

1128 31. What is the recommended management for the prevention and treatment of these1129 conditions?

#### 1130 **Recommendation 42**

1131 Basic preventive measures should be advised to all patients with CP including 1132 adequate calcium/vitamin D intake and, if indicated, pancreatic enzyme 1133 supplementation, regular weight-bearing exercise and smoking and alcohol 1134 avoidance. Additional pharmacologic treatment should be reserved for patients 1135 with osteopathy and, in particular, osteoporosis.

1136 **Grade of Recommendation GPP – Strong consensus (97% agreement)** 

1137

#### 1138 **Commentary**

1139 The reasons for osteopathy in CP are multifactorial; (i) low serum vitamin D concentrations due to impaired absorption of fat-soluble vitamin D, poor dietary intake 1140 1141 (including calcium) and/or sunshine exposure, (ii) smoking and alcohol intake, (iii) low 1142 physical activity, and (iv) chronic inflammation, all contribute. Therefore, basic 1143 preventive measures should be advised to all patients with CP including adequate 1144 calcium/vitamin D intake and PERT if indicated, regular weight-bearing exercise and 1145 avoidance of smoking and alcohol (4). Research on pharmaceutical supplementation of 1146 vitamin D and calcium in patients with osteopenia and adding bisphosphonates in 1147 osteoporosis has mainly been performed in post-menopausal women and elderly 1148 patients. Based on these findings, and bearing in mind that the cost and side effects are 1149 limited, one could consider in patients with osteopathy to supplement vitamin D (800 1150 IU) and calcium (500 - 1,000 mg) daily (149). In patients with osteopenia it is 1151 recommended to repeat the DXA every two years, whereby in patients with osteoporosis 1152 there are no specific recommendations beside appropriate medication, screening for 1153 other causes and/or referral to a bone specialist (4).

Journal Pression

# 1155 **Conflict of interest**

1156 The expert members of the working group were accredited by the ESPEN Guidelines Group, the ESPEN Education and Clinical Practice Committee, and the ESPEN executive. 1157 1158 All expert members have declared their individual conflicts of interest according to the 1159 rules of the International Committee of Medical Journal Editors (ICMJE). If potential 1160 conflicts were indicated, they were reviewed by the ESPEN guideline officers and, in cases of doubts, by the ESPEN executive. None of the expert panel had to be excluded 1161 1162 from the working group or from co-authorship because of serious conflicts. The conflict of interest forms are stored at the ESPEN guideline office and can be reviewed by ESPEN 1163 1164 members with legitimate interest upon request to the ESPEN executive.

1165

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1169

## 1170 Author Contributions

All authors contributed: literature research, PICO questions and writing the
corresponding recommendation and comments; MA: overall manuscript writing and
editing; DNL and SCB: critical revision of the final manuscript; all authors approved the
final submitted version of the manuscript.

1175

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1179

# 1180 Figure legends

- 1181 <u>Figure 1:</u> Algorithm suggesting nutritional management in acute pancreatitis. HTG:
- 1182 hypertriglyceridemia; EN: enteral nutrition; PN: parenteral nutrition. Adapted from
- 1183 Adiamah et al. (28).

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# Table 1. Levels of evidence

1					
1++	High quality meta-analyses, systematic reviews of RCTs, or RCTs with a very low risk of bias				
1+	Well-conducted meta-analyses, systematic reviews, or RCTs with a low risk of bias				
1-	Meta-analyses, systematic reviews, or RCTs with a high risk of bias				
2++	High quality systematic reviews of case control or cohort studies. High quality case control or				
	cohort studies with a very low risk of confounding or bias and a high probability that the				
	relationship is causal				
2+	Well-conducted case control or cohort studies with a low risk of confounding or bias and a				
	moderate probability that the relationship is causal				
2-	Case control or cohort studies with a high risk of confounding or bias and a significant risk that				
	the relationship is not causal				
3	Non-analytic studies, e.g. case reports, case series				
4	Expert opinion				

According to the Scottish Intercollegiate Guidelines Network (SIGN) grading system. Source: SIGN 50: A guideline developer's

handbook. Quick reference guide October 2014 [SIGN 50]. RCT=randomized controlled trial

# Table 2. Grades of recommendation (6)

А	At least one meta-analysis, systematic review, or RCT rated as 1++, and directly applicable to the target population; or A body of evidence consisting principally of studies rated as 1+, directly applicable to the target population, and demonstrating overall consistency of results				
В	A body of evidence including studies rated as 2++, directly applicable to the target population; or A body of evidence including studies rated as 2+, directly applicable to the target population and demonstrating overall consistency of results; or Extrapolated evidence from studies rated as 1++ or 1+				
0	Evidence level 3 or 4; or Extrapolated evidence from studies rated as 2++ or 2+				
GPP	Good practice points/expert consensus: Recommended best practice based on the clinical experience of the guideline development group				

RCT=randomized controlled trial

Strong consensus	Agreement of >90% of the participants
Consensus	Agreement of >75-90% of the participants
Majority agreement	Agreement of 50-75 % of the participants
No consensus	Agreement of <50% of the participants

According to the AWMF (Arbeitsgemeinschaft der Wissenschaftlichen Medizinischen Fachgesellschaften, Association of the Scientific

Medical Societies in Germany) methodology (8)

# Table 4. Criteria for systematic search for literature - databases, filters and

# keywords

Publication date	From 1977 to December 2018
Language	English
Databases	Pubmed, EMBASE, Cochrane library
Filters	human
Publication type	Cohort study, controlled trial, systematic review
Keywords	Acute pancreatitis, chronic pancreatitis, nutrition

# Table 5: Nutritional assessment in the patient with chronic pancreatitis

		2	
Anthropometric	Biochemical	Symptom	Body
assessment	assessment	assessment	composition
<ul> <li>Change in body weight</li> <li>Functional assessment: Hand-grip strength dynamometry / 6- minute walk tests / sit to stand tests.</li> <li>Skin fold thickness, waist circumference and mid arm muscle</li> </ul>	<ul> <li>assessment</li> <li>Fat soluble vitamins (A, D, E, K)</li> <li>Bone health (Parathyroid hormone)</li> <li>Trace elements (magnesium, selenium, zinc)</li> <li>Anemia screen (iron studies, B12, folate,</li> </ul>	<ul> <li>assessment</li> <li>Change in dietary intake</li> <li>Appetite</li> <li>Presence of symptoms that impact on oral intake (nausea / pain / indigestion / early satiety)</li> <li>Presence of exocrine /</li> </ul>	<ul> <li>CT / US imaging of muscle stores (muscle mass)</li> <li>DXA scanning (bone mineral density)</li> </ul>
circumference.	ferritin and CRP)	endocrine	
Presence of ascites / edema	• Glycemic control: HbA1c and random glucose	dysfunction	

CRP = C-reactive protein, HbA1c = hemoglobin A1c, CT = computed tomography, US = ultrasound, DXA = dual-energy X-ray

absorptiometry

