Original Article

Systematic Review: Components of a Comprehensive Geriatric Assessment in Inflammatory Bowel Disease—A Potentially Promising but Often Neglected Risk Stratification

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Abstract

Background: The population of older patients with inflammatory bowel disease [IBD] is increasing. Patient age does not fully account for poor outcomes and its clinical utility for risk stratification is limited. Comprehensive geriatric assessment [CGA], comprising a somatic, functional, mental, and social assessment or frailty, could be a predictor tool.

Aims: To systematically review literature on the kind of components of a CGA being used in adult IBD patients and the association of these components with adverse health outcomes.

Methods: An electronic literature search was performed on January 16, 2018, using PubMed, Embase, Web of Science, the Cochrane Library, CENTRAL, Emcare, and PsycINFO. Longitudinal studies relating somatic, functional, mental, and social assessment or frailty to adverse health outcomes during follow-up in IBD patients were included. The Newcastle-Ottawa scale was used to assess individual study quality.

Results: Of 4080 identified citations, 27 studies were included, reporting 169 associations. Median sample size was 108 patients (interquartile range [IQR] 60–704). No studies performed subgroup analyses on older patients, and the highest mean age reported was 52.7 years. Somatic and functional assessments were used in three studies, mental in 24, and social in five. No study assessed cognitive status, functional performance, or frailty. In 62 associations [36.7%], components of a CGA were significantly associated with adverse health outcome measurements.

Conclusions: Components of a CGA were associated with adverse health outcomes in IBD patients, but older patients were under-represented. More studies among older patients with IBD are warranted to further establish the clinical impact of a CGA.

Key Words: Crohn’s disease; ulcerative colitis; comprehensive geriatric assessment
1. Introduction

The number of older patients with inflammatory bowel disease [IBD] is increasing.¹ This increase can be explained by both a rising prevalence due to the ageing of the population and a rising incidence of IBD in older patients,²³ and also by greater availability of treatment options.⁴ A recent population-based epidemiological study from The Netherlands reported a doubling of the IBD incidence in older patients, from 11.71 per 100,000 persons in 1991 to 23.66 per 100,000 persons in 2010.³ Older patients with IBD are at higher risk of IBD-related hospitalisation and surgery than younger patients.⁴ They are also at higher risk of developing serious adverse events during IBD treatment, such as infections or lymphoproliferative disorders.⁵ Older patients show a larger heterogeneity in their somatic, functional, mental, and social abilities or frailty compared with younger patients.⁶ A comprehensive geriatric assessment (CGA) aims to systematically explore these components of a patient's health.⁷ In other medical fields such as oncology and nephrology, research performed in older patients shows a relationship between impairments found during a CGA and adverse health outcomes, which could be helpful in clinical decision making.⁸⁹ In IBD, preliminary baseline results from our cohort study in 135 IBD patients, aged ≥65 years, indicated a high prevalence of frailty, measured with the Geriatric 8 questionnaire, and impaired physical capacity, measured using hand-grip strength.¹⁰ However, how impairments in these components of a CGA may be related to adverse health outcomes in IBD patients has not been systematically evaluated.

Therefore, the aim of this systematic review is to study the literature on the different components of a CGA used in adult IBD patients and the association of these components with adverse health outcomes, impaired quality of life [QoL], and functional or cognitive decline after follow-up.

2. Materials and Methods

2.1. Search strategy

Our literature search aimed to identify original longitudinal studies in IBD patients, in which the association between components of a CGA at baseline and IBD-related adverse health outcomes, non-IBD-related adverse health outcomes, health-related [HR]QoL questionnaires, and functional or cognitive decline after follow-up, was examined. In our search strategy, IBD was defined as Crohn's disease [CD] or ulcerative colitis [UC]. If a study included patients with IBD-unclassified [IBD-U] or indeterminate colitis [IC], these results were taken into account as well.¹¹

2.2. Components of a comprehensive geriatric assessment

The purpose of a CGA is to systematically explore four different domains as a reflection of patients' health, namely the somatic, functional, mental, and social domains.⁷ The somatic domain includes malnutrition by using malnutrition screening tools, taking a medical history, medication use, and anthropometrics. This domain is usually included as part of routine care. The functional domain includes functional performance, and can be measured with questionnaires such as [instrumental] activities of daily living ([I]ADL) as well as physical capacity, measured with tests such as handgrip strength, gait speed, or balance, or measured with questionnaires. The mental domain includes both cognitive status (measured with tests such as the Six Item Cognitive Impairment Test [6CIT] or the Mini-Mental State Examination [MMSE]) and depression or anxiety (measured with questionnaires such as the Geriatric Depression Scale [GDS]). The social domain assesses social support and is measured by questionnaires assessing living situation or marital status. The above-mentioned domains are integrated into an assessment of the overall level of frailty. Frailty is a state of increased vulnerability to poor resolution of homeostasis following a stress. Its presence, which can be assessed using frailty indices such as the Groningen Frailty Indicator,¹² increases the risk of adverse outcomes.¹¹²

2.3. Outcome parameters

Outcome parameters were categorised in IBD-related adverse health outcomes, non-IBD-related adverse health outcomes, [HR]QoL questionnaires, and functional or cognitive decline after follow-up. The following outcomes were considered IBD-related: an exacerbation or flare-up of disease measured with IBD disease activity scores such as [simplified] Crohn's Disease Activity Index ([S]CDAI), Simple Clinical Colitis Activity Index [SCCAI], Harvey-Bradshaw Index [HBI], Partial Mayo score [PMS], or Modified Truelove and Witts Activity Index [MTWAI], or with biological parameters such as C-reactive protein, faecal calprotectin, haemoglobin, haematocrit, mean corpuscular volume, leukocytes, platelet count, or erythrocyte sedimentation rate, or established with endoscopic/radiological examination. The need to step up medication, use of corticosteroids, the need for IBD-related surgery, and the occurrence of IBD-related complications such as strictures, fistulas, and extra-intestinal manifestations, were also considered to be relevant IBD-related outcomes. The [Short] Inflammatory Bowel Disease Questionnaire ([S]IBDQ) was considered to be an IBD-related outcome parameter because of the amount of questions considering IBD symptoms. The following outcomes were considered to be non-IBD-related adverse health outcomes: emergency department visits, outpatient department visits, all-cause hospitalisation, any surgery or any abdominal surgery, length of any hospital stay, and mortality. Outcome parameters reporting on [HR]QoL, functional decline (using questionnaires such as [I]ADL), or cognitive decline (using measurements or questionnaires such as the 6CIT) were also considered relevant outcome measures.

2.4. Literature search

On January 16, 2018, seven online databases [PubMed, Embase, Web of Science, the Cochrane Library, CENTRAL, Emcare, and PsyNFO] were searched using synonyms of IBD, combined with synonyms of different components of a CGA. As we surmised that the number of studies addressing components of a CGA in an older IBD population would be low, we included all studies that investigated components of a CGA known to influence adverse health outcomes in older patients among adult patients. After the initial search, a second search was performed solely regarding anxiety terms. For full details of the search strategy for PubMed, see Supplementary Material A, available as Supplementary data at ECCO-JCC online. The searches were restricted to articles in Dutch and English. Also, conference and meeting abstracts were excluded. There were no restrictions on publication date.

2.5. Study selection

The eligibility of all studies identified by the search was independently evaluated by at least two authors [VA, FLK, or EK]. For any article that seemed potentially relevant based on the title and abstract, the full text was retrieved and screened. Studies were included when containing original data reporting on an association between
any component of a CGA at baseline and an outcome of interest after follow-up in IBD patients in a longitudinal study design. In case of disagreement on the eligibility of studies, consensus was reached after discussion with at least one additional author [FvD, SM, or PM]. Discussion with additional authors because of disagreement on eligibility took place in 24 out of 4080 studies, which represents a 99.4% agreement on the selection of studies during evaluation of eligibility. Cross-referencing was performed using the reference list of the included publications, to ensure all relevant studies were identified.

2.6. Data extraction and quality assessment
The following items were extracted from each study: Publication data [author, year, and journal], study design, setting, duration of follow-up, patient characteristics [sample size, mean age, disease type, inclusion criteria], type of geriatric assessment [somatic, functional, mental, and social assessment or frailty], correction for confounding factors, the outcome and conclusion of the study. To assess the methodological quality and risk of bias of the studies included, we adapted the Newcastle-Ottawa scale to the purpose of this review [Supplementary Material B, available as Supplementary data at ECCO-JCC online].14 Two authors [VA and FLK] performed data extraction and quality assessment; in case of disagreement, a third author was consulted [PM]. The Preferred Reporting Items for Systematic Reviews and Meta-analyses [PRISMA] checklist, which is a checklist for an evidence-based minimum set of items for reporting in systematic reviews, is available in Supplementary Material C, available as Supplementary data at ECCO-JCC online.15

2.7. Data presentation
Study characteristics are presented in tables per included study. Accumulated descriptive statistics of the included studies are presented by calculating the percentage of studies reporting on associations between components of a CGA with outcomes. The overall sample size of included studies is expressed as median and interquartile range [IQR]. In this review, an ‘association’ implies an investigated, but not necessarily a statistically significant, relationship between a component of a CGA and an outcome of interest. The main findings of the included studies regarding the associations of components of a CGA with outcomes of interest are presented in tables. When authors performed an adjustment for potential confounders, these confounders are tabulated per included study; and when hazard ratio [HR], odds ratio [OR], or relative risk [RR] was adjusted for confounders, this is reported as an adjusted ratio [aHR, aOR, or aRR]. When possible, the fully adjusted model was reported.

2.8. Supplementary analysis
Because of the low median sample size in the included studies, a supplementary analysis was performed. The six studies with the largest sample size were analysed, and the association of components of a CGA with outcomes of interest was described.

3. Results
3.1. Search results and study selection
The first database search identified 3296 unique citations [Figure 1]. After initial screening of title and abstract, 246 studies were potentially eligible and the full text was screened. After full-text review, 226 were excluded and the remaining 20 studies were included. A second additional database search identified 784 citations and yielded five studies [for flowchart see Supplementary Figure 1, available as Supplementary data at ECCO-JCC online]. Cross-referencing yielded two additional relevant studies, which resulted in a total of 27 studies included in this review.

3.2. Study characteristics
Table 1 shows an overview of the included studies. The median sample size of all included studies was 108 patients [IQR 60–704]. Out of the 27 included studies, 22 [81.5%] were performed in the USA or Europe.16–37 The majority of the included studies had an observational prospective study design [77.8%],18,20,24–26,31,33–42 the median follow-up time was 12 months [IQR 10–22.9], and follow-up data were extracted from medical records or insurance data in 38.6% of associations,16,17,19,23,27,32,42 assessed during hospital visits in 33.1%,18,21,23,24,27,29,31,33,34,36–38 and self-reported in 28.6%,20,22,24,26,28,33,39–41 Twelve [44.4%] studies included both CD and UC,14,17,19,20,22,28,32–37,40,41 and of two of these included IBD-U or IC as well.19,40 Ten studies included only CD patients,16,23–27,31,36,39,42 and five studies only UC,21,29,30,37,39 None of the studies was specifically designed for older patients or performed subgroup analyses on older patients.

3.3. Reported components of a comprehensive geriatric assessment
The included studies reported on a total of 169 associations in which the relationship between a component of a CGA and an outcome measurement was investigated. An ‘association’ therefore implies an investigated, but not necessarily a statistically significant, relationship. Somatic and functional assessment were measured in 39 associations [23.1%],27,32,42 mental in 117 associations [69.2%],16–22,24–26,28,41 and social in 13 associations [7.7%].16,20,22,23,30 None of the studies used a measurement of functional performance, cognitive status, or frailty [Figure 2].

3.4. Reported outcomes
IBD-related adverse health outcomes were the main outcome of interest, reported as an outcome measurement in 125 associations [74.0% of all associations].17,18,20,23,25,31,33–42 [HR]QoL was used as an outcome measurement in eight associations [4.7%].18,20,22,24,26,28,33,39–41 One of the included studies [1.2% of the associations] used a composite outcome measurement comprising [HR]QoL, disease progression, and any readmissions or hospitalisations.24 None of the studies used functional or cognitive decline as an outcome measurement [Figure 2].

3.5. Association of geriatric impairments and outcomes
Table 2 shows an overview of the investigated associations of components of a CGA with adverse health outcomes. A significant association between a component of a CGA and an outcome of interest, in which more geriatric impairment leads to worse outcome, was presented as ‘+’+. A significant association between a component of a CGA and an outcome of interest, in which more geriatric impairment leads to better outcome, was presented as ‘−’. A non-significant association was presented as ‘ns’. In Supplementary Table 1 [available as Supplementary data at ECCO-JCC online], the available association measures are presented. In 62 associations [36.7%] there was a statistically significant association between an impairment in somatic, functional, mental, or social assessment and a higher risk of adverse
When the authors performed an adjustment for potential confounders, these confounders are tabulated in Supplementary Table 2, available as Supplementary data at ECCO-JCC online. The detected effect of geriatric impairments on the outcomes of interest is summarised in Figure 4.

### 3.6. Somatic and functional assessment

Somatic and functional assessment was performed in three studies, resulting in 39 associations [23.1% of all associations]. Of these, 32 associations reported on malnutrition and seven reported on physical capacity [all handgrip strength]. None of the studies reported on functional performance using questionnaires such as ADL or IADL.

The different studies used a variety of screening tools to measure malnutrition. These were the malnutrition universal screening tool,\[42\] malnutrition inflammation risk tool,\[27\] subjective global assessment,\[7,42\] nutrition risk screening 2002,\[42\] Onodera’s prognostic nutritional index,\[42\] controlling nutritional status,\[42\] bioelectrical impedance analysis measuring phase angle,\[7\] and malnutrition diagnosis code.\[32\]

Malnutrition or high risk of malnutrition was highly prevalent in the included studies, with a range between 10.6% and 72.5%. Takaoka et al., using the Onodera’s prognostic nutritional index, reported that up to 72.5% of included hospitalised patients were at high risk of malnutrition.\[42\] Micic et al. analysed hospital discharges in 53,942 patients and found that, when using the ICD-9 malnutrition code, 10.6% of patients were diagnosed with malnutrition.\[32\] In their study, malnutrition was an independent predictor of 30-day readmission (aOR 1.37, 95% confidence interval [CI] 1.22–1.54).\[32\]

Jansen et al. reported on physical capacity using handgrip strength, resulting in seven associations. A mean handgrip strength at baseline of 38.2 kg (standard deviation [SD] 9.9) was reported, but no data on the prevalence of impaired handgrip strength were presented. Handgrip strength did not predict different measures of disease activity, disease-related complications, or a composite endpoint in this study.\[27\]

A total of 97.4% of the associations on somatic or physical capacity included only CD patients. Malnutrition or impaired physical capacity was a significant predictor of adverse health outcomes in 10 out of 39 associations [25.6%] [Figure 5].

### 3.7. Mental assessment

Mental assessment was evaluated in 24 studies, resulting in 117 associations [69.2% of all associations]. Of these, all associations reported on depression and/or anxiety. None of the studies reported on cognitive status.
| Publication characteristics | Study population | Study characteristics |
|-------------------------------|------------------|-----------------------|
| **Author**                   | **Year**         | **Country** | **Number of patients at baseline** | **Age, years [mean]** | **Disease type** | **Inclusion criteria** | **Follow-up duration** |
| Allegretti                   | 2015             | USA         | 324                                   | 41.7                  | CD/UC               | ≥18 years, hospital admission for non-elective IBD-related reason | 90 days |
| Ananthakrishnan             | 2013             | USA         | 10834                                 | 49.5 & 44.7; 52.7 & 47.9 | CD/UC               | Exclusion of patients with anxiety or depression date of diagnosis code before surgery | CD 11.5 years & 7.7 years; UC 12.5 years & 9.1 years |
| Banovic                      | 2010             | France      | 57                                    | 41.2                  | CD                  | Outpatients complaining of fatigue, no steroid dependence, no rheumatoid or peripheral arthritis | 1 year |
| Barnes                       | 2017             | USA         | 52498                                 | 20.2% ≥60; 31.2% ≥60 | CD/UC               | ≥18 years, excluding patients with discharge codes for both CD and UC | 90 days |
| Bernstein                   | 2010             | USA         | 704                                   | 52.1                  | CD                  | >18 years >18 years and ≤80 years, clinical remission for ≥1 month, endoscopic remission at baseline, p.o. or rectal mesalamine dose stable for 1 month or 6-mercaptopurine and azathioprine dose stable for 3 months, no use of p.o. or rectal corticosteroids within the past 30 days | 1 year |
| Bitton                      | 2008             | Canada      | 101                                   | 33.6                  | CD                  | 18–65 years, clinical remission for ≥1 month, p.o. or rectal mesalamine dose stable for 1 month or 6-mercaptopurine and azathioprine dose stable for 3 months, no use of p.o. or rectal corticosteroids within the past 30 days, no current complications, no previous extensive small bowel resection, no presence of ileostomy or colostomy, no antibiotic use at baseline | 1 year |
| Boer, de                     | 1998             | The Netherlands | 271                                   | 42                    | CD/UC               | Attending IBD outpatinet clinic in year preceding study, completion of follow-up | 1 year |
| Cámara                       | 2011             | Switzerland | 467                                   | 41.6                  | CD                  | Complete/returned questionnaires at follow-up | 1.5 years |
| Cámara                       | 2011             | Switzerland | 476                                   | 41.8                  | CD                  | Adult patients with recurrence of CD symptoms, no missing or invalid information on important control variables, returning baseline questionnaires within 6 months of inclusion | 1.5 years |
| Deter                        | 2008             | Germany     | 108                                   | 52.9% <30, 47.1% >30 | CD                  | 18–55 years, at least one active disease episode [defined as requiring drug treatment] in past 2 years, no psychotherapy, no resection for CD within past 2 years and no further relapse thereafter, no ongoing immunosuppressive therapy or resection in the near future, no colostomy or ileostomy | 2 years |
| Gaines                       | 2016             | USA         | 5707                                  | 43                    | CD                  | ≥18 years, internet access | 1 year |
| Jansen                       | 2016             | Germany     | 55                                    | 40                    | CD                  | 18–75 years, CDAI <200, occurrence of relapse/flare-ups, intestinal complication or hospitalisation within the past 2 years, known disease location and behaviour within the past 2 years, absence of cancer or other severe disease, no pregnancy or lactation, no high-dose systemic corticosteroid treatment within 3 months before study entry, absence of stoma or short bowel syndrome, and no BMI <17.5 or severe weight loss | 6 months |
| Kocher                       | 2018             | USA         | 2798                                  | 41                    | CD/UC               | ≥18 years, excluding patients without follow-up | Mean |
| Langhorst                    | 2013             | Germany     | 80                                    | 45.1 & 48.7 | UC                  | 18–75 years, self-reported clinical remission for ≥1 week and <12 months, an interval of 4 weeks in remission for 4 weeks at the beginning of the 12-months interval, absence of clinically active disease, no infectious or chronic active colitis, no current use of antibiotics or corticosteroids, no treatment within the past 3 months with immunosuppressive drugs, no complete colectomy, no relevant somatic comorbidities, no pregnancy | 22 months, 1 year |
| Publication characteristics | Study population | Study characteristics |
|-----------------------------|------------------|----------------------|
| **Author**                  | **Year**         | **Country**          | **Number of patients at baseline** | **Age, years [mean]** | **Disease type** | **Inclusion criteria** | **Follow-up duration** |
| Levenstein                   | 2000             | Italy                | 63                              | 38.8                | UC              | Clinical remission, for at least 2 months off systemic or local steroids, using oral and rectal 5-aminosalicylate or oral azulfidine in maintenance doses as sole therapy, completion of follow-up | 68 months |
| Mardini                      | 2004             | USA                  | 18                              | 31                  | CD              | No history of psychosis and/or clinical depression that required hospitalisation, no structuring disease and/or history of ileostomy, total colectomy, or short gut syndrome ≥18 years, no colectomy or indications of cardiovascular illness | 2 years |
| Maunder                      | 2005             | Canada               | 146                             | 42.7                | UC              | ≥18 years, return of questionnaires <1 month of administration | 7–37 months [median 686 days] |
| McCombie                     | 2015             | New Zealand          | 54                              | 33.5                | CD/UC/IBD-U     | ≥18 years, primary discharge diagnosis of CD or UC, or primary diagnosis of an IBD-related complication and a secondary diagnosis of CD or UC, no death during index admission, exclusion of elective admissions | 6 months |
| Micic                        | 2017             | USA                  | 43680                           | 47.8                | CD/UC           | Diagnosis established ≥4 months before inclusion or at least one recurrence of symptoms, completion of baseline and follow-up visit, no pregnancy, no missing data on depression and anxiety scores | 30 days |
| Mikocka-Walus                | 2008             | Australia            | 139                             | 50                  | IBD/IBS/HCV     | Diagnosis established ≥4 months before inclusion or at least one recurrence of symptoms, completion of baseline and follow-up visit, no pregnancy, no missing data on depression and anxiety scores | 1 year |
| Mikocka-Walus                | 2016             | Switzerland          | 2289                            | 40.5                | CD/UC/IC        | Diagnosis established ≥4 months before inclusion or at least one recurrence of symptoms, completion of baseline and follow-up visit, no pregnancy, no missing data on depression and anxiety scores | 8 years |
| Mittermaier                  | 2004             | Austria              | 60                              | 31                  | CD/UC           | 18–65 years, in remission 8 to 12 weeks after a flare defined as CDAI/CAI, and in remission at baseline for at least 4 weeks [CDAI <150 or CAI <5], sufficient knowledge of German, no known or evident psychiatric diseases, no psychopharmacotherapy use, absence of stoma | 1.5 years |
| North                        | 1991             | USA                  | 33                              | 39.8                | CD/UC           | At least one gastrointestinal exacerbation during study period, occurring no earlier than 4 months after date of enrolment | 2 years |
| Persoons                     | 2005             | Belgium              | 100                             | 34                  | CD              | ≥18 years, refractory, active [CDAI >150] luminal disease treated with infliximab [5 or 10 mg/kg], no short bowel syndrome, absence of stoma, no participation in clinical trial | 10 months |
| Riley                        | 1990             | UK                   | 100                             | Range 20–78         | UC              | ≥18 years, maintenance sulphasalazine [2–4 g daily] or delayed release mesalazine [800–1600 mg daily]. Clinical remission [absence of blood in stool and macroscopic appearance of normal mucosa or erythema only on sigmoidoscopy], absence of oral or rectal steroids within 1 month of study entry | 48 weeks |
| Takaoka                      | 2017             | Japan                | 40                              | 32.4                | CD              | Hospitalised at gastroenterology unit during inclusion | A median of 25.5 days [IQR 13.5–45.0] |

CD, Crohn's disease; UC, ulcerative colitis; IBD, inflammatory bowel disease; p.o., by mouth; CDAI, Crohn's Disease Activity Index; BMI, body mass index; IBD-U, inflammatory bowel disease-unclassified; IBS, inflammatory bowel syndrome; HCV, hepatitis C virus; IC, indeterminate colitis; CAI, Clinical Activity Index; IQR, interquartile range.

1Age only mentioned separately for CD or UC and with or without psychiatric comorbidity.
2Age and follow-up duration only mentioned separately for patients with CD or UC and with or without psychiatric comorbidity.
3Mean age only mentioned separately for relapse group and continued remission group.
4Number of patients stated in table is number of IBD patients, and only associations regarding IBD patients from this study are included.
533 patients out of 85 patients developed the endpoint [a flare] in the 2-year study period and were included in analyses.
Depressive and/or anxiety symptoms were mostly measured with the Hospital Anxiety and Depression Scale (HADS), in eight studies [33 associations]. The presence of a depression diagnosis code, anxiety diagnosis code, or a combination of these, was used in four studies, resulting in 30 associations. Depressive symptoms or a diagnosis of depression were present in between 2.3% and 32.0% of patients. Anxiety symptoms or diagnosis were present in between 9.4% and 39.0% of patients.

Ananthakrishnan et al. combined a diagnosis of a depressive disorder and/or a generalised anxiety diagnosis into one ‘psychiatric comorbidity’ factor. In their cohort of 10 834 patients, this factor was associated with the occurrence of several adverse health outcomes. For instance, psychiatric comorbidity was associated with an increased risk of IBD-related surgery in CD (aOR 1.22 [95% CI 1.01–1.47]) and the use of steroids in CD patients (aOR 1.83 [95% CI 1.58–2.13]). A study by Mikocka-Walus et al., assessing depressive and anxiety symptoms with the Hospital Anxiety and Depression Scale (HADS), found that depression was associated with their composite outcome measure ‘clinical recurrence’ in both CD and UC. Anxiety was associated with clinical recurrence in CD, but not in UC.

Half of all associations with depressive and/or anxiety symptoms [49.5%] described CD patients; the other associations described UC patients [26.4%] or no distinction was made in IBD type [23.9%]. Depressive and/or anxiety symptoms were predictive of adverse health outcomes in 50 out of 117 associations [42.7%]. Of the associations reporting on patients with CD, 57.9% were statistically significant, and 42.9% of associations reporting on UC patients were statistically significant.

### 3.8. Social assessment

Social assessment was evaluated in five studies and this resulted in 13 associations [7.7% of all associations]. Three associations reported on marital status or living situation [living together versus alone]. The other 10 associations assessed social support or social functioning using the Social Network Index, Social Support Inventories (medical outcomes study [MOS] and enhancing recovery in coronary heart disease [ENRICHD]) or Social Support List.

Allegretti et al. reported the highest percentage, of 57.9% patients being single, divorced, or widowed. A study by De Boer et al. reported 12% of patients as living alone.

Bernstein et al. assessed the amount of high-contact roles using the Social Network Index. This index calculates the number of different social roles in which the patient participates at least once every 2 weeks, involving contact with a familiar person. The study reported a presence of 19.9% of ≤ 4 high-contact roles in the ‘flare’ group compared with 17.4% in the ‘non-flare’ group; this difference was not significant. In a study by Camara et al., mean social support was 24.26, SD 5.50, measured with the ENRICHD Social Support Inventory on a scale from 6 [low social support] to 30 [high social support]. Better social support was an independent predictor of less adverse events [aOR 0.666, 95% CI 0.516–0.839, p = 0.002].

Four associations [30.8%] described CD patients; in other associations with social functioning no distinction was made regarding IBD type. There were no associations with social functioning in UC patients alone. In two out of the 13 associations [15.4%], a significant relationship between lower social functioning and adverse health outcomes was reported [Figure 3].

### 3.9. Supplementary analysis

The overall sample size of the included studies was relatively low, which causes a low power to detect statistical significance. Hence, to enhance statistical power, we selected the six studies with the largest sample size. These studies accounted for 56 associations [33.1% of total associations] with a minimum sample size of 2289 patients and a maximum of 52 498 patients. The associations described in these studies mostly [98.2%] assessed depressive and/or anxiety symptoms; 62.5% of these associations showed a statistically significant relationship between a component of a CGA and a higher risk for adverse health outcomes [Figure 3].

### 3.10. Quality assessment

The overall study quality assessed by the modified Newcastle-Ottawa scale was moderate to low [Table 3]. There were concerns about the representativeness of the cohorts, the duration of follow-up, and the adequacy of follow-up.

None of the studies focused on older patients or performed separate analyses on a subgroup of older patients. Six studies [partly] excluded older patients, with the lowest maximum age of exclusion of 55 years reported in the study by Deter et al.

Ten studies had a questionable duration of follow-up which was only partly or not enough for investigated outcomes to occur, and six studies did not report on patients lost to follow-up.

### 4. Discussion

This systematic review aimed to identify longitudinal studies describing components of a CGA in IBD patients and their associations with adverse health outcomes. There were three main findings. First, components of a CGA were used in 27 studies and none...
| Author               | Number of patients | Component of comprehensive geriatric assessment and measured method                                                                 | Outcome                              | Association                                                                 |
|---------------------|--------------------|-------------------------------------------------------------------------------------------------------------------------------------|--------------------------------------|-------------------------------|
| Allegretti**16**    | 324                | Mental assessment by depression diagnosis<br>Mental assessment by anxiety diagnosis<br>Social assessment by marital status     | 90-days readmission                  | Depression: readmission + Anxiety: readmission ns<br>Marital status: readmission ns |
| Ananthakrishnan**17** | 10834             | Mental assessment by psychiatric comorbidity [depressive disorder diagnosis and/or generalised anxiety diagnosis]                  | IBD-related surgery, IBD-related hospitalisation, all-cause hospitalisation, anti-TNF use, immunomodulator use, steroid use, outpatient visits, GE visits, abdominal CT/MRI scan, lower GI endoscopies | IBD-related surgery: CD + Anxiety+, depression ns<br>UC ns, IBD-related hospitalisation: CD ns UC ns, all cause hospitalisations: CD + UC+, anti-TNF use: CD ns UC ns, immunomodulator use: CD + UC ns, steroid use: CD + UC+, outpatient visits: CD + UC ns, GE visits: CD ns UC ns, abdominal CT/MRI scan: CD ns UC ns, GI endoscopies: CD - UC-            |
| Banovic**18**       | 52                 | Mental assessment by depression [HADS-D]                                                                                        | QoL: SF36 Vitality, mental health and general health, CDAI score       | IBD-related surgery: CD + UC+ Anxiety: readmission CD + UC+ |
| Barnes**19**        | 52498              | Mental assessment by depression diagnosis                                                                                          | 90-days readmission                  | Depression: vitality+, mental health+, general health+, CDAI ns, Anxiety: vitality ns, mental health+, general |
| Bernstein**20**     | 704                | Mental assessment by Positive and Negative Affect Schedule<br>Social assessment by Social Network Index<br>Social assessment by married/not married | Exacerbation                       | Depression: readmission CD + UC+ Anxiety: readmission CD + UC+ |
| Bitton**21**        | 60                 | Mental assessment by depression [SCL-90-R]                                                                                        | Exacerbation                        | Depression: ns<br> Anxiety: ns |
| Bitton**22**        | 101                | Mental assessment by depression [SCL-90-R]                                                                                        | Exacerbation                        | Depression: ns Anxiety: ns |
| Roet, de**23**      | 222                | Mental assessment by depression [CES-D]<br>Mental assessment by emotional functioning [IBDQ]<br>Social assessment by IBIDQ<br>Social assessment by living alone yes/no<br>Social assessment by MOS Social Support Survey<br>Social assessment by ENRICHD Social Support Inventory | GE and GP visits                    | Depression: GE ns GP ns Emotional functioning: GE ns GP ns |
| Câmara**24**        | 458                | Social assessment by ENRICHD Social Support Inventory                                                                             | IBD-related adverse event          | Social functioning [IBDQ]: GE + GP ns Social functioning [living alone]: GE ns GP ns Social functioning [social support]: GE ns GP ns Social functioning: |
| Câmara**25**        | 461                | Mental assessment by depression [HADS-D]<br>Mental assessment by anxiety [HADS-A]                                                | Exacerbation                        | Depression+, anxiety+ |
| Deter**26**         | 87                 | Mental assessment by depression [BDI]<br>Mental assessment by anxiety [STAI]                                                        | Combined measurement of health care utilization, HRQoL, and the somatic course of disease | Depression: combined outcome ns, health care use ns, HRQoL ns, somatic course ns Anxiety: combined outcome ns, health care use ns, HRQoL ns, somatic course ns |
| Gaines**27**        | 2144               | Mental assessment by PROMIS depression questionnaire                                                                             | SCDAl >150, any abdominal surgeries, any hospitalisations, use of anti-TNF therapy | Depression: SCDAl+, any abdominal surgeries ns, any hospitalisations+, use of anti-TNF ns |
| Jansen**27**        | 55                 | Somatic assessment by malnutrition [SGA]<br>Somatic assessment by malnutrition [MIRT]<br>Somatic assessment by malnutrition [BIA Phase angle]<br>Functional assessment by handgrip strength | CDAI, HBI, CD-related hospitalisations, flares, complications, CD-related surgeries, composite assessment of CD-related doctor visits, complications, CD-associated hospitalisation, exacerbation, CD-related surgery and changes in CD medication | Malnutrition [SGA]: all seven outcome parameters ns Malnutrition [MIRT]: hospitalisations+, surgeries+, complications+, exacerbation+, composite assessment+, CDAI: ns, HBI ns Malnutrition [BIA phase angle] and handgrip strength: for both all seven outcome parameters ns |
| Author          | Number of patients | Component of comprehensive geriatric assessment and measured method | Outcome                                                                 | Association                                                                 |
|-----------------|--------------------|---------------------------------------------------------------------|---------------------------------------------------------------------------|----------------------------------------------------------------------------|
| Kochar          | 2798               | Mental assessment by depression [PHQ-8]                             | Exacerbation, new biologic prescription, new steroid prescription, any hospitalisation, IBD-related surgery | Depression: exacerbation CD + UC ns, new biologic prescription CD + UC, new steroid prescription CD + UC ns, hospitalisation CD + UC, IBD-related surgery CD + UC+ |
| Langhorst       | 75                 | Mental assessment by depression [HADS-D]                            | Exacerbation                                                              | Depression: exacerbation ns                                                  |
| Levenstein      | 62                 | Mental assessment by depression [CES-D]                             | Exacerbation                                                              | Depression: exacerbation ns; short term [<8 months] exacerbation ns           |
| Mardini          | 18                 | Mental assessment by depression [BDI]                                | CDAI score                                                                | Depression: CDAI+, anxiety CDAI+                                              |
| Maunder         | 99                 | Mental assessment by depression [CES-D]                             | Disease activity                                                          | Depression: disease activity ns                                              |
| McCombie        | 54                 | Mental assessment by anxiety [HADS-D]                               | SIBDQ score                                                              | Depression: SIBDQ ns, anxiety: SIBDQ ns                                      |
| Micic           | 43680              | Somatic assessment by Malnutrition diagnosis                        | All-cause hospital readmission within 30 days                              | Malnutrition: readmission+, depression: readmission ns, anxiety: readmission+ |
| Mikocka-Walus    | 59                 | Mental assessment by depression [HADS-D]                            | Exacerbation                                                              | Depression [HADS-D]: exacerbation ns, depression [SCL90]: exacerbation ns, anxiety [HADS-A]: exacerbation ns, anxiety [SCL90]: exacerbation ns |
| Mikocka-Walus    | 2007               | Mental assessment by depression [HADS-D]                            | Clinical recurrence, fistulas, exacerbation, IBD surgery, biologic use, steroid use | Depression: clinical recurrence CD + UC+, fistula CD+, exacerbation UC+, IBD surgery CD+, biologic use CD+, steroid use CD+ |
| Mittermaier     | 60                 | Mental assessment by depression [BDI]                                | Exacerbation                                                              | Depression: exacerbation at 12 months+, exacerbation at 18 months+, Anxiety: exacerbation at 12 months ns, Anxiety: exacerbation at 18 months ns, Anxiety: exacerbation at 12 months ns |
| North           | 32                 | Mental assessment by depression [BDI]                                | Change in disease activity and exacerbation                               | Depression [BDI]: gastrointestinal scale score 1-month lag ns, gastrointestinal scale score 2-month lag ns, 1 month before exacerbation ns, 2 months before |
| Persoons        | 100                | Mental assessment by MDD presence [PHQ-9]                           | Response to infliximab, achievement of remission, time to re-treatment     | Depression: response to infliximab ns, failure to achieve remission+, time to re-treatment+, Anxiety: response to infliximab ns, failure to achieve remission ns, time to retreatment ns |
| Riley           | 92                 | Mental assessment by depression [HADS-D]                            | Exacerbation                                                              | Depression: exacerbation ns                                                  |

Table 2. Continued
Table 2. Continued

| Author | Number of patients | Component of comprehensive geriatric assessment and measured method | Outcome |
|--------|-------------------|---------------------------------------------------------------|---------|
| Takaoka | 42                | Somatic assessment by malnutrition [SGA], intestinal resection | LOS    |
|        |                   | Somatic assessment by malnutrition [MUST], intestinal resection | LOS    |
|        |                   | Somatic assessment by malnutrition [NRS 2002], intestinal resection | LOS    |
|        |                   | Somatic assessment by malnutrition [O-PNI], intestinal resection | LOS    |
|        |                   | Somatic assessment by malnutrition [CONUT], intestinal resection | LOS    |

+ : a significant association between a component of a geriatric assessment and outcome of interest in which more geriatric impairment leads to worse outcome; - : a significant association between a component of a geriatric assessment and outcome of interest in which more geriatric impairment leads to better outcome. When both univariate and multivariate analyses were performed, only results from multivariate analyses [most corrected model] are tabulated. For an extended version of Table 2 including association measures, see Supplementary Table 1, available as Supplementary data at ECCO-JCC online.

For corrected confounders see supplementary Table online.

- ns, non-significant; HR[QoL], health-related quality of life; IBD, inflammatory bowel disease; anti-TNF, anti-tumour necrosis factor; GE, gastroenterologist; CT, computerised tomography; MRI, magnetic resonance imaging; GI, gastrointestinal; CD, Crohn’s disease; UC, ulcerative colitis; HADS-D, Hospital Anxiety and Depression Scale Depression; St-ID, Short Form 36; SCL, Short Form 36; EORTC, European Organisation for Research and Treatment of Cancer; GIQLI, Gastrointestinal Quality of Life Index; SF-36, Short Form 36; SF-12, Short Form 12; PSQI, Pittsburgh Sleep Quality Index; PGI, Psychological General Well-being Index; SES, Social Support Life Interactions; MUST, malnutrition universal screening tool; NRS, nutrition risk screening; O-PNI, Onodera’s Prognostic Nutritional Index; CONUT, controlling nutritional status; LOS, length of hospital stay.

a Total of 124 patients, 59 IBD patients.
objective measurement of physical capacity (handgrip strength). Handgrip strength is often used to diagnose sarcopenia, and there is a growing evidence for the association between sarcopenia and adverse health outcomes. However, the study by Jansen et al. did not find a significant association between reduced handgrip strength and adverse health outcomes. This could be explained by the short follow-up duration (6 months), low mean age (40 years), and small sample size (55 patients). Thus, in older patients with IBD, evidence regarding impairments in physical capacity, cognitive status, or frailty, and its associations with adverse health outcomes, is lacking.

Despite the relatively young population included and the lack of a CGA, we found impairments in the components of a CGA to be prevalent and, in more than one-third of the associations, significantly associated with adverse health outcomes. Older patients are more susceptible to geriatric impairments such as depression, low physical capacity, and malnutrition, compared with younger patients. It is very likely that prevalence of geriatric impairment and its association with adverse health outcomes have been underestimated in this review when applied to older patients. Evidence on the underlying pathophysiological relationship between components of a CGA and adverse health outcomes is still scarce, especially in IBD.

In Figure 6 we present a summary of the potential pathophysiological interactions between these components and IBD disease outcomes. In IBD patients, depression contributes to lower pain thresholds, more reported symptoms, and a poorer well-being. This could contribute to the relationship between depressive symptoms and adverse health outcomes found in this systematic review. In the Health Aging and Body Composition [Health ABC] study on 3075 individuals aged 70–79 years, a significant and independent

\[\text{Figure 3. Visual representation of significant associations. Positive significant associations are associations in which more geriatric impairment led to more adverse outcomes, negative significant associations in which more geriatric impairment led to less adverse outcomes. A: percentage of significant associations in associations of all included studies. B: percentage of significant associations in associations of the six largest studies.}\]

\[\text{Figure 4. The detected effect of geriatric impairments on adverse health outcomes in inflammatory bowel disease patients. No studies on functional or cognitive impairment were found.}\]
Association between physical capacity, measured with both low quadriceps muscle strength and low handgrip strength, and serum levels of the inflammatory markers tumour necrosis factor [TNF] and interleukin [IL]-6 was found. High levels of inflammatory markers are associated with increased morbidity and mortality in older persons. In IBD patients, these pro-inflammatory markers correlate with disease activity, and it has been shown that patients with IL-6 serum levels >20 picograms per millilitre have a 17-fold increased risk of relapse over a 1-year period compared with patients with a lower level. The latter could also contribute to the relationship between low physical capacity and [IBD-related] adverse health outcomes. Malnutrition, besides being one of the principal mechanisms involved in the genesis of sarcopenia, is also a well-known risk factor for poor prognosis in IBD, especially postoperative complications.

The disease course of IBD could also influence several components of a CGA, in this way causing a bidirectional relationship between geriatric impairment and adverse health outcomes. For example, IBD patients experiencing an exacerbation of disease express a higher risk of malnutrition, due to the decrease of oral food intake or increased gastrointestinal nutrient loss. While studying the predictive role of a CGA in IBD, relationships should therefore be interpreted with caution, and associations should be corrected for disease activity to take into account this possible bidirectional relationship. The multimodal effects of IBD and its treatment may very well be an important cause of geriatric impairments or frailty.

To enhance statistical power, as sample sizes in the included studies were small, a sensitivity analysis was performed by selecting six studies with the largest patient population. The percentage of statistically significant associations increased from 36.7% to 62.5%. This suggests that most included studies lacked statistical power to detect significant differences. However, 98.2% of the

Figure 5. Graphic representation of associations of somatic or physical impairment, depressive and/or anxiety symptoms, and social impairment with adverse health outcomes in inflammatory bowel disease patients. No studies reported on cognitive impairment, functional impairment, or frailty.

Table 3. Quality assessment of the included studies.

| Author          | Year | Representativeness of the exposed cohort | Ascertainment of exposure [geriatric measure] | Assessment of outcome | Sufficient duration of follow-up | Adequacy of follow-up |
|-----------------|------|-----------------------------------------|---------------------------------------------|-----------------------|----------------------------------|----------------------|
| Allegratti      | 2015 | ±                                       | +                                           | +                     | +                               | +                    |
| Ananthakrishnan | 2013 | +                                       | +                                           | +                     | ±                               | ±                    |
| Banovic         | 2010 | −                                       | +                                           | ±                     | +                               | ±                    |
| Barnes          | 2017 | +                                       | +                                           | +                     | ±                               | ±                    |
| Bernstein       | 2010 | +                                       | +                                           | +                     | ±                               | ±                    |
| Bitton          | 2003 | −                                       | +                                           | ±                     | +                               | ±                    |
| Bitton          | 2008 | −                                       | +                                           | +                     | ±                               | −                    |
| Boer, de        | 1998 | +                                       | +                                           | ±                     | ±                               | −                    |
| Câmara          | 2011 | ±                                       | +                                           | +                     | ±                               | ±                    |
| Câmara          | 2011 | ±                                       | +                                           | +                     | ±                               | ±                    |
| Deter           | 2008 | −                                       | +                                           | ±                     | ±                               | ±                    |
| Gaines          | 2016 | +                                       | +                                           | ±                     | ±                               | ±                    |
| Jansen          | 2016 | −                                       | +                                           | ±                     | −                               | ±                    |
| Kochar          | 2018 | ±                                       | +                                           | +                     | ±                               | ±                    |
| Langhorst       | 2013 | −                                       | +                                           | +                     | −                               | −                    |
| Levenstein      | 2000 | ±                                       | +                                           | +                     | +                               | +                    |
| Mardini         | 2004 | −                                       | +                                           | +                     | +                               | +                    |
| Maunder         | 2005 | −                                       | +                                           | +                     | ±                               | ±                    |
| McCombie        | 2015 | ±                                       | +                                           | +                     | ±                               | ±                    |
| Metc            | 2017 | +                                       | +                                           | +                     | ±                               | ±                    |
| Mikocka-Walus   | 2008 | +                                       | +                                           | ±                     | ±                               | ±                    |
| Mikocka-Walus   | 2016 | +                                       | +                                           | ±                     | ±                               | ±                    |
| Mintermaier     | 2004 | −                                       | +                                           | +                     | +                               | +                    |
| North           | 1991 | ±                                       | +                                           | ±                     | +                               | ±                    |
| Persoons        | 2005 | −                                       | +                                           | +                     | ±                               | ±                    |
| Riley           | 1990 | +                                       | +                                           | ±                     | +                               | +                    |
| Takaoka         | 2017 | −                                       | +                                           | +                     | ±                               | ±                    |
Associations described by the six largest studies reported on depression and/or anxiety and therefore are not representative of a full CGA. We were unable to perform sensitivity analyses separately for each component of a CGA due to the few studies included per factor: three-quarters of the associations included in our systematic review described depressive and/or anxiety symptoms. Alexakis et al. performed a systematic review and meta-analysis on the relationship between depressive state and disease course in adults with IBD but, due to a lack of randomised controlled studies, small study populations, and a large variety in depression symptom scores, it was inconclusive. Therefore, even though the majority of associations found in our systematic review concerned depression and/or anxiety, no firm conclusion can be drawn regarding the influence of depression and/or anxiety on adverse health outcomes.

A limitation of this review is the fact that, because of the heterogeneity of the components of a CGA, outcome measures, and reported measures of association, we could not perform a formal meta-analysis. Furthermore, interpretation of the results regarding the number of significant associations has to be viewed with caution due to a possibility of publication bias, as negative or non-significant associations may not have been reported in included studies. Besides this, the number of older patients in the included studies is low, and therefore it is not guaranteed that results of these studies can be extrapolated to the population of older patients with IBD.

The strengths of our review include the extended systematic search performed in seven online databases without a restriction on publication date. In this way all potentially relevant studies, concerning associations of components of a CGA with adverse health outcomes in IBD patients, were assessed. In addition, a quality assessment using the adapted Newcastle-Ottowa Scale was performed. One of the greatest strengths of this review is the important addition to existing literature regarding IBD in the older patient, because little research has been performed on this subject so far.

The findings of this study imply that more research regarding geriatric assessment in older IBD patients is needed. With more evidence available on the association between components of a CGA and adverse health outcomes, a risk stratification could be performed regarding geriatric impairment. In this way, older patients at high risk for adverse health outcomes could be selected in time and monitored closely or an alternative treatment regimen might be selected according to their risk profile.

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Conflict of Interest
No conflicts of interest.

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Author Contributions
VA, FvD, SM, and PM contributed to the study concept and design, VA, FLK, and EK assessed eligibility of studies, VA and FLK performed data extraction and quality control of included studies, VA performed data analysis, interpretation of data, and drafting of the manuscript, FvD, SM, and PM contributed to critical revision of the manuscript, and PM supervised the study.

Supplementary Data
Supplementary data are available at ECCO-JCC online.

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