Informal caregiver well-being during and after patients’ treatment with adjuvant chemotherapy for colon cancer: a prospective, exploratory study

S. M. C. H. Langenberg 1 · H. Poort 2,3 · A. N. M. Wymenga 4 · J. W. de Groot 5 · E. W. Muller 6 · W. T. A. van der Graaf 1,7 · J. B. Prins 8 · C. M. L. van Herpen 1

Received: 16 May 2020 / Accepted: 4 September 2020 / Published online: 15 September 2020
© The Author(s) 2020

Abstract
Introduction Caring for a significant other during cancer treatment can be demanding. Little is known about the well-being of informal caregivers of patients with colon cancer. This study aims to examine informal caregiver well-being during adjuvant chemotherapy for colon cancer.

Material and methods This exploratory longitudinal, prospective study measured the course of informal caregiver burden (Self-Perceived Pressure of Informal Care), distress (Hospital Anxiety and Depression Scale), health-related quality of life (RAND-36), marital satisfaction (Maudsley Marital Questionnaire), social support (Social Support List – Discrepancies), fatigue (Abbreviated Fatigue Questionnaire), and self-esteem (Caregiver Reaction Assessment) before (T0), during (T1), and after (T2) patients’ treatment.

Results Baseline data of 60 out of 76 eligible dyads (79%) were analyzed. Mean levels of informal caregiver burden and distress improved significantly over time, as did their health-related quality of life and perceived social support. At baseline, 30% and 26.7% of informal caregivers reported moderate-to-high levels of burden and clinically relevant levels of distress, respectively, which changed to 20% and 18.8% at T2. Informal caregiver burden and distress at baseline were the strongest predictors of informal caregiver burden and distress during and following patients’ treatment, respectively.

Conclusion When informal caregivers and patients experience problems before start of adjuvant chemotherapy, problems seem to improve over time. Approximately 20% of informal caregivers remain burdened and distressed after patients’ end of treatment. Paying attention to baseline distress and burden seems indicated, as these were strong predictors of informal caregivers’ well-being during and after treatment.

Keywords Caregiver · Colon cancer · Burden · Distress

Electronic supplementary material The online version of this article (https://doi.org/10.1007/s00520-020-05738-w) contains supplementary material, which is available to authorized users.
Introduction

Colon cancer is a common cancer among men and women [1]. In 2018, colon cancer accounted for 6% of all new cancer cases and 551,269 people died as a consequence of colon cancer worldwide [1]. In 2018, 9555 people were newly diagnosed with colon cancer in The Netherlands [2]. After diagnosis and staging, treatment plans are made according to (inter)national guidelines. When a patient is diagnosed with high-risk stage II or III colon cancer, surgical resection alone results in a 5-year survival rate of 60–80% for stage II and 30–60% for stage III disease [3]. To further improve survival, patients receive adjuvant chemotherapy containing fluoropyrimidines and oxaliplatin (i.e., 5FU/leucovorin with oxaliplatin (FOLFOX) or capecitabine with oxaliplatin (CAPOX)), or capecitabine monotherapy when oxaliplatin is contra-indicated [4, 5]. Common side effects of this regimen are nausea, vomiting, hand-foot syndrome, and sensory, peripheral neuropathy. Being diagnosed with cancer and receiving cancer treatment impact the patient and can cause psychological distress [6].

After a cancer diagnosis and during cancer treatment, support by a significant other is indispensable. Informal caregivers support patients both practically and emotionally [6, 7]. These tasks can cause burden and distress, which may impact an informal caregiver’s ability to support the patient during treatment [8, 9]. Informal caregiver burden can be experienced on several dimensions and be related to emotional, social, physical, spiritual functioning, and/or practical and financial problems [7]. The National Cancer Institute defined distress as “emotional, social, spiritual or physical pain or suffering that may cause a person to feel sad, afraid, depressed, anxious or lonely” [10]. Previous studies identified burden and distress as important problems for informal caregivers [6, 9]. Patients and informal caregivers cope with cancer as a dyad and informal caregivers’ distress may sometimes exceed patients’ distress [11, 12]. Informal caregiver burden and distress are interrelated and share risk factors, such as female gender, younger age, and perceived patient distress [7, 9]. Importantly, informal caregiver burden and distress may also negatively influence the informal caregivers’ physical health and social functioning [6–9]. Little is known about the course of well-being of informal caregivers of patients treated with adjuvant chemotherapy for colon cancer.

This prospective, observational study had four exploratory aims. First, we aimed to examine the course of informal caregiver well-being. Second, we intended to explore the clinically relevant levels of informal caregiver burden and distress. Third, we aimed to identify baseline risk factors for higher informal caregiver burden and distress during and after a patient’s treatment. Fourth, we explored the association between informal caregiver burden and distress and patient distress before, during, and after adjuvant chemotherapy.

Material and methods

Setting and participants

We conducted a prospective, observational study between October 2013 and September 2017 in four hospitals in The Netherlands. We recruited patients aged 18 or older, proficient in Dutch, who were scheduled to receive adjuvant chemotherapy (CAPOX or capecitabine monotherapy) after surgery for colon cancer, and their informal caregivers, for participation in this study.

Procedure

The attending physician or the nurse practitioner approached the patient and their informal caregiver for study participation after informing them about starting adjuvant chemotherapy. Written informed consents were obtained from both patient and informal caregiver. The study included completion of paper-and-pencil questionnaires at home at three time points: (1) baseline, before starting adjuvant chemotherapy (T0), (2) between the second and third cycle (T1), and (3) 3 months after ending adjuvant chemotherapy (T2). We asked patients and informal caregivers to complete questionnaires separately. The study was approved by the local medical ethics committee of the Radboud University Medical Center (registration number 2013/393).

Measures

Sociodemographic and clinical factors Patients and informal caregivers self-reported their age, sex, level of education (categorized using the International Standard Classification of Education), and employment (paid work, housekeeper, disablement insurance act, retired, volunteer, study) at baseline. Informal caregivers completed a general questionnaire on the nature of their relation to the patient (partner, child, sibling, friend), whether they lived together with the patient (yes/no) and the extent of their caregiving tasks (hours of caring, dependency of the patient, caring for more than patient alone), as well as their needs during caring for their significant other (practical and/or emotional support from social support system and/or professionals, information services from hospital and/or general practitioner, and/or better communication with physician/nurses/general practitioner). Furthermore, we inquired whether the treatment side effects of cancer treatment had a negative influence on informal caregiver well-being (no/yes, somewhat/yes).

Patients self-reported whether they had a colostomy after surgery, the total number of cycles of adjuvant chemotherapy, and experienced toxicity (T1 and T2). Patients also reported what sort of complications they had after surgery. The attending medical oncologist provided information which adjuvant chemotherapy...
chemotherapy (CAPOX versus capecitabine monotherapy) was prescribed.

We used the Self-administered Comorbidity Questionnaire (SCQ) to assess 14 common medical conditions in both patients and informal caregivers, and additional comorbidities could also be reported [13]. For each condition, patients and informal caregivers indicated whether it was present, being treated, or imposed functional limitations. For the present study, we used data on whether a comorbidity was present and causing functional limitations.

**Informal caregiver well-being** Informal caregivers completed the 9-item Self-Perceived Pressure from Informal Care (SPPIC) [14]. Items are scored on a 5-point scale ranging from 0 (no!) to 5 (yes!). We dichotomized scores to 0 ("no!") and "no") and 1 ("yes!", "yes,"
and more or less"). Total scores range from 0 to 9, and higher scores indicate higher levels of perceived informal caregiver burden. In accordance with other studies [15–17], we classified informal caregivers into low (0–3), moderate (4–6), and high levels of burden (7–9) based on their total score. Patients and informal caregivers each completed the Hospital Anxiety and Depression Scale (HADS) to measure distress [18]. The HADS consists of 14 items assessing symptoms of anxiety and depression during the past week. Items are scored on a 4-point Likert scale ranging from 0 to 3. Total scores range from 0 to 42, and higher scores indicate more distress. We used the 36-item RAND-36 Health Survey to assess functional status, well-being, and general health [19]. Scores on each of the eight subscales are transformed into a range from 0 to 100; higher scores indicate higher levels of functioning, well-being, and general health. The Social Support List – Discrepancies (SSL-D) is a 34-item questionnaire assessing discrepancies between an individual’s need for social support and their perceived social support [20, 21]. The questionnaire assesses six types of social support, namely emotional interactions, problem-focused emotional support, esteem support, instrumental interactions, social companionship, and informational support. The score on every item is transformed into a sum score ranging from 34 to 136. Higher scores indicate more unmet needs for social support. We used the “marital satisfaction” subscale of the Maudsley Marital Questionnaire (MMQ) to assess marital satisfaction [22]. This is a 10-item questionnaire, answered on a 9-point scale (0–8), ranging from 0 to 80. Higher scores indicate decreased marital satisfaction. We instructed participants to complete this questionnaire if they had a partner relationship. We administered the Abbreviated Fatigue Questionnaire (AFQ), a validated 4-item questionnaire to measure fatigue [23, 24]. The AFQ is an abbreviated version of the subscale fatigue severity of the Checklist Individual Strength. Items are scored on a 7-point scale ranging from “no, that’s not correct” (score 1) to “yes, that’s correct” (score 7). The total score on the 4-item questionnaire ranges from 4 to 28. Higher scores indicate greater levels of fatigue. Informal caregiver self-esteem was measured by the subscale caregiver self-esteem of the Caregiver Reaction Assessment Scale (CRA) [25]. The subscale consists of 7 items, scored on a 5-point Likert scale. The total score reflects a mean score of 7 items and ranges between 1 and 5; a higher score indicates more self-esteem.

**Statistical analysis**

Descriptive statistics were used for baseline characteristics of the patients and informal caregivers. Chi-square and Fisher’s exact tests were performed to examine the relation between categorical variables. Independent-samples t tests and Mann-Whitney U tests were used to compare continuous variables. We analyzed informal caregiver burden, distress, health-related quality of life, marital satisfaction, discrepancies in social needs, fatigue, and informal caregiver self-esteem with one-way ANOVA with repeated measures. Next, we applied the same statistical analyses for patient distress, health-related quality of life, marital satisfaction, discrepancies in social needs, and fatigue. We applied the Greenhouse-Geisser correction when the assumption of sphericity was violated. Post hoc tests with Bonferroni’s correction were used to identify which specific means differed. To identify cases of clinically significant distress, we applied a HADS total clinical cutoff score of ≥ 11 to identify patients with clinically significant distress [26] and a cutoff score of ≥ 12 for informal caregivers [27]. To identify possible cases of depression (HADS-D) and anxiety (HADS-A), we used the cutoff score of ≥ 8 [26, 27]. McNemar’s tests were carried out to determine whether the proportion of patients or informal caregivers exceeding the cutoff for clinically relevant distress differed between baseline and T2. We used multiple linear regression to predict informal caregiver burden and distress at T1 and T2 from informal caregiver gender, age, burden, distress, fatigue, and patient distress at baseline (method: enter). For the linear regression analyses, we used the continuous variables of burden, distress, and fatigue. All data analyses were performed using the statistical package for the social sciences (SPSS, version 24.0). Statistical significance was determined based on a two-sided alpha of 0.05.

**Results**

Of 76 eligible patients and their informal caregivers, 62 (82%) dyads enrolled in the study, and 14 (18%) declined participation, mostly because participation was considered too burdensome. Of the 62 dyads that provided consent, 2 dyads withdrew consent after enrollment. Thus, T0 was completed by 60 (79%), whereas T1 by 58 (76%) and T2 by 51 dyads (67%). Baseline, caregiving, and treatment characteristics are provided in Table 1. Table 2 displays an overview of what support
Table 1  Characteristics of informal caregivers and patients

| Characteristic                                      | Informal caregivers (n = 60) | Patients (n = 60) |
|-----------------------------------------------------|-----------------------------|-------------------|
| Age, years (SD)                                     | 59.88 (12.72)               | 63.83 (7.74)      |
| Sex                                                 |                             |                   |
| Male                                                | 25 (41.7%)                  | 34 (56.7%)        |
| Female                                              | 35 (58.3%)                  | 26 (43.3%)        |
| Education                                           |                             |                   |
| Lower education level (ISCED ≤ 4)                   | 35 (58.3%)                  | 41 (69.5%)        |
| Higher education level (ISCED 5–8)                  | 25 (41.6%)                  | 18 (30.5%)        |
| Missing                                             | 0                           | 1                 |
| Dependent children (lives with)                     |                             |                   |
| No                                                  | 43 (71.7%)                  |                   |
| Yes                                                 | 9 (15.0%)                   |                   |
| Missing                                             | 8 (13.3%)                   |                   |
| Employment                                          |                             |                   |
| Paid work                                           | 23 (38.3%)                  | 19 (31.7%)        |
| Housekeeper                                         | 9 (15%)                     | 7 (11.7%)         |
| Disablement insurance act                            | 0 (0%)                      | 7 (11.7%)         |
| Retired                                             | 25 (41.7%)                  | 28 (46.7%)        |
| Volunteer work                                      | 4 (6.7%)                    | 3 (5%)            |
| Study                                               | 2 (3.3%)                    | 0 (0%)            |
| Informal caregiver’s relationship to patient         |                             |                   |
| Partner                                             | 53 (91.4%)                  |                   |
| Child                                               | 2 (3.4%)                    |                   |
| Sibling                                             | 2 (3.4%)                    |                   |
| Friend                                              | 1 (0.8%)                    |                   |
| Missing                                             | 2                           |                   |
| Informal caregiver lives with patient                |                             |                   |
| Yes                                                 | 53 (89.8%)                  |                   |
| No                                                  | 6 (10.2%)                   |                   |
| Missing                                             | 1                           |                   |
| Providing > 8 h of care                             |                             |                   |
| T0                                                  | 19/60 (31.9%)               |                   |
| T1                                                  | 18/57 (31.6%)               |                   |
| T2                                                  | 5/51 (9.9%)                 |                   |
| Patients functioned independently or mostly independently |                   |                   |
| T0                                                  | 51/57 (89.5%)               |                   |
| T1                                                  | 51/57 (89.5%)               |                   |
| T2                                                  | 48/51 94.2%                 |                   |
| Providing care to more than patient alone           |                             |                   |
| T0                                                  | 7/60 (11.9%)                |                   |
| T1                                                  | 5/57 (8.8%)                 |                   |
| T2                                                  | 3/51 (5.9%)                 |                   |
| Negative influence of patients’ side effects on caregiver well-being (measured on T1) |                   |                   |
| No                                                  | 22/53 (41.5%)               |                   |
| Yes, somewhat                                       | 27/53 (50.9%)               |                   |
| Yes                                                 | 4/53 (7.5%)                 |                   |
| Colostomy after surgery                             |                             |                   |
| No                                                  |                             | 52 (86.7%)        |
| Yes                                                 |                             | 8 (13.3%)         |
| Complications after surgery                         |                             |                   |

* Springer
| Type of adjuvant chemotherapy | Informal caregivers (n = 60) | Patients (n = 60) |
|-------------------------------|-------------------------------|------------------|
| CAPOX                         | 53 (88.3%)                    | 7 (11.7%)        |
| Capecitabine monotherapy      |                              |                  |

| Number of cycles completed     | Informal caregivers (n = 60) | Patients (n = 60) |
|-------------------------------|-------------------------------|------------------|
| 8                             | 37 (72.5%)                    |                  |
| 7                             | 6 (7.8%)                      |                  |
| 6                             | 4 (7.8%)                      |                  |
| 5                             | 1 (2%)                        |                  |
| 4                             | 1 (2%)                        |                  |
| 3                             | 1 (2%)                        |                  |
| 2                             | 1 (2%)                        |                  |
| 1                             | 0 (0%)                        |                  |
| Missing                       | 9                             |                  |

| Reason for modification treatment (n = 26) | Informal caregivers (n = 60) | Patients (n = 60) |
|-------------------------------------------|-------------------------------|------------------|
| Discontinuation oxaliplatin only          | 13/26 (50%)                  |                  |
| Discontinuation oxaliplatin and capecitabine | 13/26 (50%)              |                  |

| Toxicity 3 months after ending adjuvant treatment | Informal caregivers (n = 60) | Patients (n = 60) |
|---------------------------------------------------|-------------------------------|------------------|
| No                                                | 1 (2%)                        |                  |
| Yes, suffered from side effect, but disappeared   | 15 (29.4%)                   |                  |
| Yes, suffered from side effects, now barely noticeable | 4 (7.8%)                  |                  |
| Yes, suffered from side effects and still do      | 31 (60.8%)                   |                  |
| Missing                                           | 9                             |                  |

| Comorbidities | Informal caregivers (n = 60) | Patients (n = 60) |
|---------------|-------------------------------|------------------|
| 0             | 16 (29.1%)                    | 11 (20.0%)       |
| 1–2           | 34 (61.8%)                    | 35 (63.7%)       |
| 3 or more     | 5 (9.1%)                      | 9 (16.4%)        |
| Missing       | 5                             | 5                |

| Most common comorbidities | Informal caregivers (n = 60) | Patients (n = 60) |
|---------------------------|-------------------------------|------------------|
| High blood pressure       | 15/59 (25.4%)                 | 22/58 (37.9%)    |
| Back pain                 | 15/57 (26.3%)                 | 14/58 (24.1%)    |
| Arthrosis                 | 12/57 (21.1%)                 | 8/58 (13.8%)     |
| Chronic obstructive pulmonary disease | 10/57 (17.5%) | 7/57 (16.7%) |
| Diabetes                  | 6/58 (10.3%)                  | 10/58 (17.2%)    |
| Hearth disease            | 2/57 (3.5%)                   | 11/58 (19%)      |
| Hindrance from comorbidity(ies) | 1/15 (6.7%)            | 0/22 (0%)        |
| High blood pressure       |                              |                  |
informal caregivers need before, during, and after a patients’ treatment with adjuvant chemotherapy. When informal caregivers reported they need support, it seemed most needed before starting chemotherapy and focused on practical support and receiving information.

Course of burden, distress, health-related quality of life, marital satisfaction, discrepancies in social needs, fatigue over time, and informal caregiver self-esteem

Table 3 shows mean scores for informal caregiver well-being over time. Mean scores for informal caregiver burden ($F[2,94] = 4.465; p = 0.014$), distress ($F[1.773,81.574] = 5.497; p = 0.008$), role emotional limitations ($F[2,94] = 8.814; p < 0.0001$), mental health ($F[2,94] = 4.949; p = 0.009$), social functioning ($F[2,94] = 3.985; p = 0.022$), and discrepancies in social support ($F[2,66] = 3.466; p = 0.037$) differed significantly between time points. Post hoc analyses indicated an improvement of all scores over time; i.e., burden, distress, and role emotional limitations decreased, whereas mental health and social functioning increased. Discrepancies in social support decreased during and increased after ending adjuvant chemotherapy. Post hoc analyses are shown in Supplementary Table 1.

Table 4 displays patients’ mean scores for health-related quality of life, marital satisfaction, discrepancies in social needs, and fatigue over time. Mean patient distress ($F[1.771,79.706] = 5.224; p = 0.010$), role physical limitations ($F[2,94] = 9.551; p < 0.0001$), vitality ($F[2,96] = 5.295; p = 0.007$), social functioning ($F[2,96] = 9.157; p < 0.0001$), general health ($F[2,92] = 6.672; p = 0.002$), marital satisfaction ($F[2,54] = 5.395; p = 0.007$), and fatigue ($F[2,92] = 11.393; p < 0.0001$) changed significantly over time. Post hoc analyses revealed an overall decrease of distress and increase in role physical limitations and social functioning. Vitality and general health decreased during and increased after ending
adjuvant chemotherapy. Marital satisfaction decreased over time. Fatigue increased during and decreased after ending adjuvant chemotherapy. Post hoc analyses are shown in Supplementary Table 2.

### Clinically relevant levels of informal caregiver burden and distress

Moderate or high levels of burden of informal caregivers were found in 17.2% (n = 10/58) and 12.1% (n = 7/58) at baseline, 19.3% (n = 11/57) and 10.5% (n = 6/57) at T1, and 12% (n = 6/50) and 8% (n = 4/50) at T2, respectively. A HADS total score exceeding the cutoff for clinically relevant distress was found in 26.7% (n = 16/60) at baseline, 22.8% (n = 13/57) at T1, and 18.8% (n = 9/48) at T2. Clinically relevant levels of informal caregiver depressive symptoms (HADS-D) were 15% at baseline, 10.5% at T1, and 12.2% at T2. Clinically relevant levels of informal caregiver anxiety (HADS-A) were 21.7% at baseline, 21.1% at T1, and 12.2% at T2. In patients, 22.4% (n = 13/58) exceeded the cutoff at baseline for clinically relevant levels of distress, 31.5% (n = 17/54) at T1, and 18.0% (n = 9/50) at T2. Clinically relevant levels of patient depression were 12.1% at baseline, 16.1% at T1, and 7.8% at T2. Clinically relevant levels of patient anxiety were 17.2% at baseline, 13.0% at T1, and 7.8% at T2. There was no significant difference between the proportions of informal caregivers (p = 0.289) or patients (p = 0.508) with clinically relevant levels of distress at baseline and T2.

### Predictors of informal caregiver distress and burden at T1 and T2

Informal caregiver gender, age, burden, distress, fatigue, and patient distress at baseline predicted informal caregiver self-perceived burden at T1 (F[6,53] = 4.493, p = 0.001, R² = 0.365) and T2 (F[6,46] = 4.523, p < 0.001, R² = 0.404). Only informal caregiver burden at baseline added significantly to the prediction at T1 (p = 0.002) and T2 (p = 0.002). The multivariate regression model was also used to predict distress at T1 (F[6,53] = 12.305, p < 0.0001, R² = 0.611) and T2 (F[6,44] = 7.204, p < 0.0001, R² = 0.532). Only baseline informal caregiver distress (p < 0.001) added significantly to the prediction at T1 (p < 0.001) and T2 (p = 0.001). Details are displayed in Table 5.

### Interaction between informal caregivers and patients

Generally, informal caregivers reported higher distress levels (T0 mean 8.47 (SD 6.83); T1 mean 7.42 (SD 6.34); T2 mean 6.50 (SD 7.14)) compared with patients (T0 mean 7.02 (SD 5.80); T1 mean 7.20 (SD 5.93); T2 mean 5.56 (SD 4.69)), but these differences did not reach statistical significance (T0

---

### Table 3: Informal caregivers’ course of burden, distress, health-related quality of life, marital satisfaction, fatigue, and self-esteem

|                           | Population norm scores | T0 Mean (SD) | T1 Mean (SD) | T2 Mean (SD) | Sign p value |
|---------------------------|------------------------|--------------|--------------|--------------|--------------|
| Burden (SPPIC) [14, 15]   | a                      | 2.90 (2.44)  | 2.82 (2.38)  | 2.04 (2.25)  | 0.014*       |
| Distress (HADS) [27]      | 8.4 (6.3)              | 8.96 (6.91)  | 7.57 (6.32)  | 6.64 (7.15)  | 0.008*       |
| Health-related quality of life (RAND 36) [19] | | | | | |
| Physical functioning      | 81.90 (23.20)          | 91.04 (10.47) | 90.94 (13.90) | 87.81 (16.24) | 0.153 |
| Role limitations due to physical health | 79.40 (35.50) | 79.89 (23.20) | 87.50 (22.82) | 84.78 (30.95) | 0.253 |
| Role limitations due to emotional problems | 84.10 (32.30) | 63.88 (41.73) | 81.25 (34.32) | 86.11 (30.62) | < 0.0001* |
| Vitality                  | 67.40 (19.90)          | 67.29 (14.73) | 71.98 (15.04) | 71.88 (20.39) | 0.067 |
| Mental health             | 76.80 (18.40)          | 73.00 (14.16) | 77.08 (12.83) | 73.83 (14.41) | 0.015*       |
| Social functioning        | 86.90 (20.50)          | 80.00 (20.52) | 84.75 (18.09) | 87.00 (15.14) | 0.022*       |
| Pain                      | 79.50 (25.60)          | 92.00 (12.98) | 89.63 (15.63) | 88.30 (17.17) | 0.173 |
| General health            | 72.70 (22.70)          | 74.56 (16.68) | 75.56 (16.42) | 73.33 (20.83) | 0.430 |
| Marital satisfaction (MMQ) [22, 28] | 13.58 (10.79) | 9.76 (9.94)  | 10.37 (10.80) | 11.66 (11.53) | 0.189 |
| Social support discrepancies (SSL-D) [21] | 43.6 (10.3) | 41.18 (9.09) | 38.38 (5.68) | 39.06 (6.63) | 0.037* |
| Fatigue (AFQ) [24]        | b                      | 10.02 (2.65) | 10.49 (6.30) | 10.55 (7.26) | 0.748 |
| Self-esteem (CRA) [29]    | 4.19 (0.41)            | 4.32 (0.54)  | 4.32 (0.49)  | 4.23 (0.48)  | 0.181 |

*p values represent significant changes in mean scores over time; references given in this table refer to population norm scores. SPPIC Self-Perceived Pressure of Informal Care, HADS Hospital Anxiety and Depression Scale, MMQ Maudsley Marital Questionnaire, SSL-D Social Support List – Discrepancies, AFQ Abbreviated Fatigue Questionnaire, CRA Caregiver Reaction Assessment, T0 baseline, T1 between 2nd and 3rd cycles of adjuvant chemotherapy, T2 3 months after ending adjuvant chemotherapy

a Scores for levels of burden: 0–3 low, 4–6 moderate, 7–9 high
b Scores for level of fatigue: low = 4, below average = 4, average = 5–8, above average = 9–14, high ≥ 15
Table 4  Patient course of distress, health-related quality of life, marital satisfaction, fatigue, and self-esteem

|                          | Population norm scores | T0 Mean (SD) | T1 Mean (SD) | T2 Mean (SD) | Significant difference over time p value |
|--------------------------|------------------------|-------------|-------------|-------------|----------------------------------------|
| Distress (HADS) [27]     | 8.4 (6.3)              | 7.22 (6.04) | 7.37 (5.41) | 5.54 (4.31) | 0.010*                                 |
| Health-related quality of life (RAND 36) [19] |                        |             |             |             |                                        |
| Physical functioning     | 81.90 (23.20)          | 76.35 (19.76) | 76.15 (20.78) | 79.52 (20.85) | 0.168                                  |
| Role limitations due to physical health | 79.40 (35.50)          | 31.77 (40.52) | 33.33 (38.71) | 59.36 (45.44) | 0.0001*                                |
| Role limitations due to emotional problems | 84.10 (32.30)          | 65.93 (44.66) | 74.81 (40.92) | 85.19 (33.75) | 0.061                                  |
| Vitality                 | 67.40 (19.90)          | 64.59 (17.61) | 58.16 (18.33) | 64.90 (19.59) | 0.007*                                 |
| Mental health            | 76.80 (18.40)          | 79.92 (14.64) | 70.49 (14.33) | 82.45 (11.07) | 0.344                                  |
| Social functioning       | 86.90 (20.50)          | 68.37 (19.45) | 72.45 (20.41) | 82.14 (19.76) | 0.0001*                                |
| Pain                     | 79.50 (25.60)          | 74.90 (23.06) | 74.82 (22.63) | 80.42 (20.56) | 0.285                                  |
| General health           | 72.70 (22.70)          | 63.51 (19.33) | 60.00 (18.44) | 68.62 (20.50) | 0.002*                                 |
| Marital satisfaction (MMQ) [28] | 13.58 (10.79)          | 5.79 (6.85)  | 6.86 (8.37)  | 9.00 (9.58)  | 0.007*                                 |
| Social support discrepancies (SSL-D) [21] | 43.2 (10.7)            | 36.86 (4.90) | 36.19 (3.03) | 36.06 (3.18) | 0.445                                  |
| Fatigue (AFQ) [24]       |                        | 12.36 (6.61) | 16.21 (7.23) | 12.85 (6.37) | 0.0001*                                |

*p values represent changes in mean scores over time; references given in this table refer to population norm scores. HADS Hospital Anxiety and Depression Scale, MMQ Maudsley Marital Questionnaire, SSL-D Social Support List – Discrepancies, AFQ Abbreviated Fatigue Questionnaire, T0 baseline, T1 between 2nd and 3rd cycles of adjuvant chemotherapy, T2 3 months after ending adjuvant chemotherapy

*Scores for level of fatigue: low = 4, below average = 5–12, average = 13–21, above average = 22–27, high ≥28

Discussion

The present longitudinal study explored the course of informal caregiver well-being, clinically relevant levels of informal caregiver burden and distress, and baseline risk factors for higher levels of burden and distress during and after a patients’ adjuvant chemotherapy for colon cancer. We found that informal caregivers seem to report more mental problems before and during chemotherapy, whereas patients report more physical problems. Additionally, 20–30% of informal caregivers report relevant levels of burden and distress, between, before, during, and after a patients’ treatment. Baseline burden and distress are risk factors for burden and distress during and after a patients’ adjuvant treatment.

Informal caregivers reported more mental than physical problems after a patient’s cancer diagnosis and during treatment, and social functioning seemed to be influenced negatively. This was also found in another study on the impact of colorectal cancer on patients and their partners [30]. This study of Traa et al. showed that both partners and patients suffer mentally and in social functioning, which is in line with our findings. Furthermore, a study of Law et al. showed that informal caregivers’ social functioning changed due to fear of burdening others, and when informal caregivers do get support, this support is perceived insufficient and not what they need on that moment [31]. Also, informal caregivers report that family and friends become avoidant in their contact with the patient and informal caregiver, which challenges their social interactions [32]. In contrast to the findings of Traa et al. who found a stabilization in mental well-being and social functioning, we observed an improvement over time. A possible explanation could be that in the study of Traa et al., 60% had an colostomy after surgery compared with only 13% in our study. Having a colostomy is known to cause distress for the patient and their informal caregiver and may impact social activities and increase social isolation [32–35].

Before starting adjuvant chemotherapy, almost 30% of informal caregivers reported moderate-to-high levels of burden and clinically relevant distress, which decreased to 20% 3 months after the end of adjuvant treatment. Ohlsson-Nevo et al. pointed out that partners’ lives are turned upside down after colorectal cancer diagnosis, being confronted with how fragile life is [33]. They had to deal with new and other unwanted responsibilities at home that they felt compelled to fulfill [33]. Northouse et al. found that during patients’ treatment for cancer, informal caregivers experienced worries about the effectiveness of the treatment, accompanied by difficulties managing side effects [6]. For informal caregivers...
who still experienced burden and distress after the patient’s completion of adjuvant chemotherapy, a possible contributing factor could be the ongoing treatment toxicity, which was reported by 61% of the patients in our study. Although toxicity can diminish over time, there is a group of patients for which the toxicity, in particular peripheral neuropathy, remains a limiting factor in a patient’s life [36], and thereby also influences the life of their significant others. Fortunately, there are new insights for treating stage III colon cancer, namely a shift from 6 to 3 months CAPOX, which is non-inferior in terms of survival and induces less toxicity [36]. Our study was performed when these data were not available yet. Since the number of cycles will be reduced and thereby also the associated cumulative toxicity, this may ultimately also positively impact the informal caregiver [36]. Another contributing factor for ongoing burden and distress can be fear of cancer recurrence. This is found in patients, and van de Wal et al. found that partners report the same levels of fear of recurrence as well [37].

When thinking of ways to support informal caregivers in need, it could be particularly helpful to predict who is in need of support and when. In our study, we found that informal caregiver burden and distress at baseline predicted informal caregiver burden and distress both during adjuvant chemotherapy and 3 months after completion of treatment. Jansen et al. reported that informal caregivers of patients with different types of cancer with higher baseline levels of burden remained burdened over the following years [38]. It is possible that informal caregivers who remain burdened and distressed have more difficulties coping, as negative coping skills are associated with higher levels of burden and distress [39].

Future research with ongoing assessment of burden and distress due to the dynamic nature of these constructs, with a focus on what causes and maintains burden and distress, is recommended. Especially it is known that long-term burden and distress cause serious general health problems [7]. Additionally, our study shows that informal caregivers’ distress exceeds patients’ distress. This finding is consistent with those of other studies [6, 30]. Based on these findings, we recommend to pay attention and offer support when informal caregiver burden and distress are observed before starting adjuvant chemotherapy, or even earlier, shortly after diagnosis. More specifically, repeated assessment of informal caregiver needs for more practical support or other informational services from care professionals is recommended. Importantly, the management of treatment side effects deserves special attention as 60% of informal caregivers report that their well-being is negatively influenced by patients’ side effects during treatment. Additionally, based on our inquiry of informal caregiver needs, the general practitioner may be particularly well-suited for providing support to the informal caregiver, as leading practitioner in informal caregiver care.

Our exploratory longitudinal study adds significantly to the scarce literature on informal caregiver burden and distress during adjuvant treatment of colon cancer. However, it is important to take into account several limitations. First, the sample size is relatively small which prevented us from studying additional predictors of informal caregiver burden and distress, such as the influence of a patient’s colostomy. Second, patients and informal caregivers who declined participation in this study often refused because participation was perceived too burdensome. This may limit the generalizability of our findings and our conclusion on burden and distress to the larger population of informal caregivers, and our findings may be an underestimation. Third, although we assessed statistical significance of changes in informal caregiver well-being over time, we were unable to determine clinical relevance due to the lack of established minimally clinically important differences (MCID) for the measures that we used. Future research assessing MCIDs for among informal caregiver population is warranted.

In conclusion, before and during adjuvant chemotherapy, informal caregivers report more mental problems whereas patients report more physical problems. When informal caregivers and patients experience problems before start of adjuvant chemotherapy, problems seem to improve over time.
Nevertheless, approximately 20% of informal caregivers remain burdened and distressed after patients’ end of treatment, and remarkably informal caregivers’ distress exceeds patients’ distress. Additionally, informal caregivers’ baseline burden and distress seem to be risk factors for ongoing burden and distress after treatment. Therefore, it is of great importance to identify burden and distress among informal caregivers of patients treated with adjuvant chemotherapy for colon cancer and offer them support according to their needs.

Acknowledgments The authors want to thank the patients and informal caregivers who participated in this study. The authors want to thank M. Hijmans–van der Vegt, C. Leverink, B. Heck, I. Schoenacker, and E. Kist for assisting with the recruitment of participants, and M. Cox for assisting with data entrance.

Authors’ contribution All authors contributed to the study conception and design. Material preparation, data collection, and analysis were performed by Simone Langenberg and Hanneke Poort. The first draft of the manuscript was written by Simone Langenberg and Hanneke Poort, and all authors commented on previous versions of the manuscript. All authors read and approved the final manuscript.

Compliance with ethical standards
Conflict of interest The authors declare that they have no conflict of interest.

Informed consent Written informed consent was obtained from all participants.

Ethical approval The local medical ethical committee gave permission for the study. All procedures performed in accordance with the ethical standards of the institutional research committee and with the 1964 Helsinki Declaration and its later amendments or comparable ethical standards.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence, and is not used by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

References
1. Union for International Cancer Control - GLOBOCAN (2018) https://www.uicc.org/news/new-global-cancer-data-globocan-2018
2. Incidentie darmkanker www.cijfersoverkanker.nl
3. Labianca R, Nordlinger B, Beretta GD, Mosconi S, Mandala M, Cervantes A, Arnold D, Group EGW (2013) Early colon cancer: ESMO Clinical Practice Guidelines for diagnosis, treatment and follow-up. Ann Oncol 24(Suppl 6):vi64–vi72. https://doi.org/10.1093/annonc/mdt354
4. Shah MA, Renfro LA, Allegra CJ, Andre T, de Gramont A, Schmoll HJ, Haller DG, Alberts SR, Yothers G, Sargent DJ (2016) Impact of patient factors on recurrence risk and time dependency of oxaliplatin benefit in patients with colon cancer: analysis from modern-era adjuvant studies in the Adjuvant Colon Cancer End Points (ACCENT) database. J Clin Oncol 34(8):843–853. https://doi.org/10.1200/JCO.2015.63.0558
5. Andre T, Boni C, Navarro M, Tabernero J, Hickish T, Topcham C, Bonetti A, Clingan P, Bridgewater J, Rivera F, de Gramont A (2009) Improved overall survival with oxaliplatin, fluorouracil, and leucovorin as adjuvant treatment in stage II or III colon cancer in the MOSAIC trial. J Clin Oncol 27(19):3109–3116. https://doi.org/10.1200/JCO.2008.20.6771
6. Northouse LL, Katapodi MC, Schafmancker AM, Weiss D (2012) The impact of caregiving on the psychological well-being of family caregivers and cancer patients. Semin Oncol Nurs 28(4):236–245. https://doi.org/10.1016/j.socn.2012.09.006
7. Northouse L, Williams AL, Given B, McCorkle R (2012) Psychosocial care for family caregivers of patients with cancer. J Clin Oncol 30(11):1227–1234. https://doi.org/10.1200/JCO.2011.39.5798
8. Applebaum AJ, Breitbart W (2012) Care for the cancer caregiver: a systematic review. Palliat Support Care 11:1–22. https://doi.org/10.1017/S1478951512000594
9. Adelman RD, Tmanova LL, Delgado D, Dion S, Lachs MS (2014) Caregiver burden: a clinical review. JAMA 311(10):1052–1060. https://doi.org/10.1001/jama.2014.304
10. National Cancer Institute - definition of distress. https://www.cancer.gov/publications/dictionaries/cancer-terms/def/distress
11. Kim Y, van Ryn M, Jensen RE, Griffin JM, Potosky A, Rowland J (2014) Effects of gender and depressive symptoms on quality of life among colorectal and lung cancer patients and their family caregivers. Psycho-oncology. 24:95–105. https://doi.org/10.1002/pon.3580
12. Northhouse LL, Mood D, Templin T, Mellon S, George T (2000) Couples’ patterns of adjustment to colon cancer. Soc Sci Med 50(2):271–284
13. Sangha O, Stucki G, Liang MH, Fossel AH, Katz JN (2003) The Self- Administered Comorbidity Questionnaire: a new method to assess comorbidity for clinical and health services research. Arthritis Rheum 49(2):156–163. https://doi.org/10.1002/art.10993
14. Pot AM, van Dyck R, Deeg DJ (1995) Perceived stress caused by informal caregiving. Construction of a scale. Tijdschr Gerontol Geriatr 26(5):214–219
15. Haj Mohammad N, Walter AW, van Oijen MG, Hulshof MC, Bergman JJ, Anderegg MC, van Berge Henegouwen MI, Henselmans I, Sprangers MA, van Laarhoven HW (2015) Burden of spousal caregivers of stage II and III esophageal cancer survivors 3 years after treatment with curative intent. Support Care Cancer 23(12):3589–3598. https://doi.org/10.1007/s00520-015-2727-4
16. Langenberg S, Reyner K, Wymenga ANM, Sieleng GCM, Veldhoven CMM, van Herpen CML, Prins JB, van der Graaf WTA (2019) Caregivers of patients receiving long-term treatment with a tyrosine kinase inhibitor (TKI) for gastrointestinal stromal tumour (GIST): a cross-sectional assessment of their distress and burden. Acta Oncol 58(2):191–199. https://doi.org/10.1080/0284186X.2018.1518592
17. Langenberg S, van Herpen CML, van Opstal CCM, Wymenga ANM, van der Graaf WTA, Prins JB (2019) Caregivers’ burden and fatigue during and after patients’ treatment with concomitant chemoradiotherapy for locally advanced head and neck cancer: a prospective, observational pilot study. Support Care Cancer 27(11):4145–4154. https://doi.org/10.1007/s00520-019-04700-9
