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The role of personality in the course of health-related quality of life and disease-specific health status among colorectal cancer survivors: A prospective population-based study from the PROFILES registry

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ABSTRACT

Background. Prospective studies in various cardiovascular populations show that Type D personality predicted impaired health-related quality of life (HRQoL) and disease-specific health status. We examined the effect of negative affectivity (NA), social inhibition (SI) and their combined effect (Type D personality) on HRQoL and disease-specific health status among colorectal cancer (CRC) patients.

Methods. CRC patients diagnosed between 2000 and 2009, as registered in the Dutch population-based Eindhoven Cancer Registry, received questionnaires on Type D personality (DS14), HRQoL (EORTC QLQ-C30) and disease-specific health status (EORTC QLQ-CR38) in 2010, 2011 and 2012.

Results. Response rates were 73% (n = 2625), 83% (n = 1643) and 82% (n = 1458), respectively. Analyses were done on those completing at least two questionnaires (n = 1735). Individuals with Type D (NA+ / SI+; 19%) and high NA (NA+ / SI-; 11%) reported a significantly worse HRQoL and disease-specific health status compared to NA-/SI+ and NA-/SI-. Differences were stable over time. Linear mixed effects models showed that Type Ds had a lower quality of life, cognitive and emotional functioning, more insomnia, diarrhea, gastrointestinal, defecation and stoma-related problems and poor body image and future perspective compared to the reference group (NA-/SI-), even after controlling for sociodemographic and clinical variables. High NA individuals (NA+ / SI-) reported similar poor health outcomes as Type Ds. However, they also reported lower social functioning and more fatigue, pain, micturition- and financial problems, while Type Ds reported more constipation, sexual problems and less sexual enjoyment.

Conclusions. Type D personality and high NA both have a significant negative stable impact on HRQoL and disease-specific health status among CRC patients.
beyond clinical characteristics [3–5]. Type D is defined by
the combination of two personality traits; the ten-
dency to experience negative emotions [negative affec-
tivity (NA)] and to inhibit self-expression in social in-
teraction [social inhibition (SI)] [6]. As such, Type D
has been associated with adverse health outcomes,
impaired health status and HRQoL, more serious ill-
ess perceptions and an increased health care utiliza-
tion [2,4,5,7,8]. Overall, the NA component of Type D
plays a more important role than the SI component in
terms of general HRQoL, while the specific combina-
tion of NA and SI in Type D has been related to symp-
toms of anhedonia, mental fatigue and decreased
motivation [9]. There are a number of potential path-
ways that could explain the relationship between per-
sonality and worse self-reported health and HRQoL.
For example, patients with Type D and high NA may
be more likely to perceive and attend to somatic symp-
toms and to interpret them as potentially pathological
[8]. In addition, Type D personality has been associated
with an unhealthy lifestyle and poor treatment adher-
ence, which, in turn, may have an adverse effect on
perceived health [8].

Research on the association of personality and
HRQoL among CRC cancer patients is scarce but rising.
A recent cross-sectional study among 162 CRC
patients reported that personality was associated with
HRQoL, independent of disease severity and psycho-
lological distress [10]. Furthermore, a prospective study
among 144 CRC patients stated that personality vari-
able can predict a decrease in HRQoL over a one-
year period [11]. Also, Type D personality was
associated with poor quality of life and mental health
among CRC and other cancer survivors in a cross-
sectional population-based study (n = 3080) [7].

A recent review stated that “more well-designed
prospective investigations are necessary to establish
the contributory role of personality dimensions for
the development of and protection from distress and impairment in the HRQoL of CRC patients”
[8]. In order to predict which CRC patients will
experience a self-reported healthy cancer survivor-
ship, we will investigate the degree to which HRQoL
and disease-specific health status can be explained
by individual differences in the personality (e.g.
NA, SI and Type D personality), while controlling
for clinical characteristics.

Methods

Setting and participants

Data from the first three waves (2010, 2011 and
2012) of a prospective population-based yearly
survey among CRC survivors from the Eindhoven
Cancer Registry (ECR) was used. The ECR
compiles data of all individuals newly diagnosed with
cancer in the southern part of the Netherlands, an
area with 10 hospitals serving 2.3 million inhabitants
[12]. Everyone diagnosed with CRC from 2000 to
2009 as registered in the ECR was eligible for par-
ticipation. Those with unverifiable addresses, with
cognitive impairment, who died prior to the start of
study or were terminally ill, with stage 0/carcinoma
in situ, and those already included in our 2009 study
or another study (n = 169) were excluded. A com-
plete overview of the selection of patients can be
found on our website http://www.profilesregistry.nl/
dataarchive/study_units/view/22 under ‘data & docu-
mentation’. This study was approved by the certified
Medical Ethics Committee of the Maxima Medical
Centre, Veldhoven, The Netherlands.

Data collection was done within PROFILES [13].
Data from the PROFILES registry is freely available
for non-commercial scientific research, subject to
study question, privacy and confidentiality restric-
tions, and registration (www.profilesregistry.nl).

Data collection

Survivors were informed of the study via a letter from
their (ex-)attending specialist. The letter included a
link to a secure website, a login name, and a pass-
word, so that interested patients could provide
informed consent and complete questionnaires
online. If the patient preferred written rather than
digital communication, (s)he could return our post-
card by mail after which (s)he received our paper-
and-pencil informed consent form and questionnaire.
Non-respondents were sent a reminder letter and
paper-and-pencil questionnaire within two months.

Survivors’ sociodemographic and clinical informa-
tion were available from the ECR. Comorbidity at
the time of the study was assessed with the adapted Self-
administered Comorbidity Questionnaire [14]. Ques-
tions on marital status, educational level, and current
occupation were included in the questionnaire.

NA, SI and Type D personality. NA, SI, and the Type
D personality construct were assessed with the Type
D Personality Scale (DS14) [6]. The 14 items are
answered on a five-point response scale ranging from
0 (false) to 4 (true). The DS14 has good measure-
ment properties; the subscales have high reliability
(Cronbach’s alpha 0.88/0.86) and good test-retest
reliability over a three-month period of r = 0.72/0.82
for the two subscales, respectively, and the DS14 dis-
criminates well between Type D and non-Type per-
sonality [6,15]. To compare the separate and
combined effects of high and low trait levels, the
standard cut-off score ≥ 10 on the NA and SI sub-
scales of the DS14 [6] was used to classify patients
Personality, quality of life and disease-specific health status

in four personality groups based on their scores at T1: NA ≥ 10 and SI ≥ 10 (NA+/SI+; the 'Type D' group), NA ≥ 10 but SI ≤ 9 (NA+/SI−; the 'NA only' group), SI ≥ 10 but NA ≤ 9 (NA−/SI+; the 'SI only' group), and both NA ≤ 9 and SI ≤ 9 (NA−/SI−; the 'reference' group).

Quality of life. The EORTC QLQ-C30 (Version 3.0) was used to assess cancer-specific QoL [16]. It contains five functional scales, a global health status/QoL scale, three symptom scales, and six single items. Each item is scored on a four-point Likert-scale, except the global QoL scale, which has a seven-point Likert-scale. Scores were linear transformed to a 0–100 scale [17]. A higher score on the functional scales and global QoL scale means better functioning and QoL. A higher score on the symptom scales mean more complaints.

Disease-specific health status. Disease-specific health status was assessed with the EORTC QLQ-CR38 [18]. It consists of two multi-item and two single-item scales, seven symptom scales, and an item on weight loss. Items were scored on a four-point Likert-scale. All scales were linearly converted to a 0–100 scale. A higher score on the EORTC QLQ-CR38 functional scales and single items (i.e. body image, sexual functioning, sexual enjoyment and future perspective) represent a higher level of function. For the symptom scales and single items, a higher score represents a higher level of symptoms.

Depression. Depressive symptoms were assessed with the depression subscale of the Hospital Anxiety and Depression Scale (HADS) comprising of seven items on a four-point Likert-scale [19]. It assesses levels of symptoms in the last week. The scale mainly covers anhedonia and loss of interest, which are core depressive symptoms.

Statistical analyses

Differences in sociodemographic and clinical characteristics between respondents, non-respondents or patients with unverifiable addresses at T1 and between patients who completed one or more questionnaires were compared with a χ², ANOVA or independent samples t-test where appropriate.

All other analyses are based on patients who completed at least two questionnaires. Differences in sociodemographic and clinical characteristics between patients with different personality types (NA−/SI−; NA−/SI+; NA+/SI−; Type D) were compared with a χ² or ANOVA while differences in HRQoL and disease-specific health status were determined by ANOVAs at each time point. As our results showed that NA had a large effect on our outcomes, the same analyses were performed for continuous NA scores divided into quartiles.

The course of HRQoL and disease-specific health status (separate models for each scale) according to the four personality groups was analyzed using linear mixed effects models (i.e. covariance pattern model with an unstructured error variance matrix and maximum likelihood estimation). This technique uses data efficiently by also including incomplete cases in the analyses. As a result of this, bias is limited and statistical power is preserved. Time was analyzed as a regular categorical predictor with three levels (i.e. three time points). Sex, age, time since diagnosis, treatment, disease stage, comorbid conditions, partnership and educational level and depression levels were entered as covariates into the models, based on a priori assumptions/hypotheses. Sociodemographic and clinical characteristics were analyzed as time-invariant predictors (i.e. baseline characteristics were used). Depression scores were analyzed as continuous time varying predictors. In order to correctly interpret all model parameters, all continuous variables have been grand-mean centered.

Finally, a sub analyses using linear mixed-effects models was performed on the general subscale global health status/QoL over time stratified by NA divided into quartiles.

All statistical tests were two-sided and considered significant if p < 0.05. All analyses were conducted using SPSS version 19.0 (Statistical Package for Social Sciences, Chicago, IL, USA). Missing items from multi-item scales of the EORTC QLQ-C30 and EORTC QLQ-CR38 were mean-imputed if at least half of the items from the scale were answered, according to the EORTC guideline. Missing items on the DS14 and HADS scales were mean imputed if only one item was missing, otherwise the scale score became missing [6,19].

Results

Sociodemographic and clinical characteristics

The questionnaire was completed by 73% (n = 2625) at T1, 83% (n = 1643) at T2 and 82% (n = 1458) at T3. Respondents were significantly younger (69.4 vs. 72.4; p < 0.01), more often male (55% vs. 48%; p < 0.01), were more often diagnosed with stage I disease (30% vs. 25%; p = 0.02) and were more often treated with radiotherapy (31% vs. 24%; p < 0.01) compared to non-respondents. Respondents significantly more often received radiotherapy (31% vs. 26%; p < 0.01) and surgery (99% vs. 96%; p = 0.01) and were more often male (55% vs. 48%; p < 0.01) compared to patients with unverifiable addresses.
Those who completed one versus those who completed ≥2 questionnaires differed with respect to gender (49% vs. 43% female; \( p < 0.01 \)), age (71.3 vs. 68.4 years; \( p < 0.001 \)), having a partner (71% vs. 79%; \( p < 0.001 \)), having a job (10% vs. 19%; \( p < 0.01 \)), and radiotherapy treatment (27% vs. 32%; \( p < 0.01 \)). Also, they more often had a lower educational level (\( p < 0.001 \)), were more often diagnosed with stage IV disease (\( p < 0.001 \)) and had worse scores on more than half of the subscales of the EORTC QLQ-C30 and EORTC QLQ-CR38 (global QoL, physical, role emotional and social functioning, fatigue, nausea, pain, dyspnea, appetite loss, sexual functioning, sexual enjoyment, future perspective, chemotherapy side effects and weight loss; data not shown). Those who completed ≥2 questionnaires had worse scores on the subscales gastrointestinal problems and defecation problems. From this point forward, only those who completed ≥2 questionnaires are described in the analyses.

At T1, 328 CRC survivors (19%) had a Type D personality (Table I). No statistically significant differences were observed between those with and without a Type D personality in age, years since diagnosis, primary treatment and tumor type. However, CRC survivors with a Type D personality were significantly more often female, more often diagnosed with stage II disease, had a lower educational level and had a partner less often compared to those without a Type D personality. Furthermore, those with NA only were younger compared to the other three groups. Finally, those with Type D and those with NA only had more comorbid conditions compared to the other two groups.

### HRQoL and disease-specific health status

At T1, patients with Type D and NA only reported a significantly worse HRQoL and more disease-specific symptoms compared to the other two groups except for sexual enjoyment which was only worse among those with a Type D personality, and female sexual problems which showed no differences between the groups (Table I). Furthermore, the differences in HRQoL scores between the Type D group and the reference group, and between the NA only group and the reference group, were clinically relevant for all subscales except for nausea (Type D group) and constipation (NA only group) [20].

These differences in HRQoL and disease-specific health status between Type D and NA only groups versus the SI only and reference groups were quite stable across the three time points, the most relevant EORTC QLQ-C30 and EORTC QLQ-CR38 subscales with respect to survivorship [21] are presented in Figure 1. However, problems with weight loss and female sexual functioning were more prevalent in the Type D group than in the other three groups at T2, and Type D patients also reported more problems regarding sexual enjoyment across all time points. Type D patients also reported more problems with sexual functioning and male sexual functioning at T1, but not at T2 or T3 and for female sexual functioning at T2 but not T1 or T3 than patients with NA only.

Linear mixed effects models showed that CRC survivors with a Type D personality were at a significantly increased risk of an impaired global quality of life, more insomnia, less sexual enjoyment, and a worse body image and future perspective compared to the reference group (NA-/SI-), even after controlling for sex, age, time since diagnosis, stage, chemotherapy, comorbidity, partner, education, time of questionnaire, and depression (Table II). Also, CRC survivors with a Type D personality were at a significantly increased risk of an impaired cognitive functioning (Beta -4.7, 95% CI -6.7--2.6), emotional functioning (Beta -10.3, 95% CI -11.8--8.7), more diarrhea (Beta 3.6, 95% CI 1.2--6.1), constipation (Beta 2.3, 95% CI 0.1--4.5), gastrointestinal problems (Beta 3.2, 95% CI 1.6--4.8), defecation problems (Beta 3.6, 95% CI 2.0--5.2), stoma-related problems (Beta 8.9, 95% CI 3.8--14.0), and female sexual problems (Beta 14.8, 95% CI 5.5--24.0) in comparison to the reference group. Other important determinants of HRQoL and disease-specific symptoms were sex, age, time since diagnosis, comorbid conditions and depression (data not shown).

Furthermore, linear mixed effects models showed that patients with NA only also reported an impaired global quality of life, more insomnia, a worse body image, and future perspective compared to the reference group but moreover, they also reported more fatigue and pain (Table II). Additionally, those who score high on NA reported more problems with cognitive functioning (Beta -5.4, 95% CI -7.8--2.9), social functioning (Beta -2.7, 95% CI -5.1--0.3), emotional functioning (Beta -12.6, 95% CI -14.4--10.7), and more diarrhea (Beta 3.7, 95% CI 0.8--6.6), gastrointestinal problems (Beta 4.4, 95% CI 2.5--6.5), defecation problems (Beta 2.9, 95% CI 1.0--4.8), stoma-related problems (Beta 8.6, 95% CI 2.7--14.5), micturition problems (Beta 3.1, 95% CI 0.8--5.4) and financial problems (Beta 3.2, 95% CI 0.7--5.6).

As patients with Type D and NA only reported very similar scores on most outcomes, we performed a sub analysis on the general and overarching “global health status/QoL” scale over time stratified by NA divided into quartiles (Figure 2). Results showed that all NA quartiles differed significantly from each other at each time point with respect to Global health status/QoL, except for quartile 1 (lowest) and 2 which did not differ.
Table I. Sociodemographic and clinical characteristics at T1 according to Type D personality.

| N (%) | All (n = 1735) | NA-SI- (n = 911) | NA-SI+ (n = 297) | NA+SI- (n = 199) | NA+SI+ (n = 328) | p-Value* |
|-------|----------------|------------------|------------------|------------------|------------------|---------|
| Sex (Female) | 743 (43%) | 383 (42%) | 110 (37%) | 92 (46%) | 158 (48%) | 0.03 |
| Age [mean (SD)] | 68.4 (9.4) | 68.8 (9.1) | 68.4 (9.2) | 66.6 (10.5) | 68.5 (9.8) | 0.03 |
| Years since diagnosis [mean (SD)] | 5.1 (2.8) | 5.1 (2.9) | 5.4 (2.8) | 5.1 (2.7) | 5.0 (2.7) | 0.21 |
| Tumor type | Colon 1031 (59%), Rectal 704 (41%) | 353 (34%) | 126 (42%) | 87 (44%) | 138 (42%) | 0.42 |
| Stage | I 522 (31%), II 610 (36%), III 503 (30%), IV 55 (3%) | 267 (30%) | 100 (35%) | 63 (32%) | 92 (29%) | 0.03 |
| Number of comorbid conditions | None 422 (25%), One 501 (30%), Two or more 736 (44%) | 239 (28%) | 95 (33%) | 33 (17%) | 55 (17%) | <0.001 |
| Quality of life [mean (SD)] | Global quality of life 78.9 (17.7), Physical functioning 82.3 (19.0), Role functioning 82.0 (26.0), Emotional functioning 86.8 (18.6), Cognitive functioning 85.3 (20.3), Social functioning 87.3 (16.8), Fatigue 20.1 (22.1), Nausea 3.2 (10.5), Pain 15.6 (23.8), Dyspnea 12.6 (23.3), Insomnia 20.3 (27.8), Appetite loss 4.2 (13.5), Constipation 8.6 (19.2), Diarrhea 10.6 (21.5), Financial problems 7.0 (19.1) | 83.3 (15.4), 84.7 (17.7), 85.3 (23.0), 94.0 (10.4), 90.1 (15.3), 91.3 (18.1), 15.6 (19.6), 2.3 (9.4), 12.5 (21.8), 9.9 (20.8), 15.9 (24.8), 2.5 (10.4), 6.7 (16.8), 8.5 (19.2), 5.3 (16.9), 89.0 (18.0), 79.5 (23.3), 26.0 (23.8), 19.6 (17.0), 8.3 (13.2), 19.6 (17.0), 8.3 (13.2), 38.9 (29.7), 19.6 (17.0), 8.3 (13.2), 38.9 (29.7), 19.6 (17.0), 8.3 (13.2), 38.9 (29.7), 19.6 (17.0), 8.3 (13.2), 38.9 (29.7), 19.6 (17.0), 8.3 (13.2), 38.9 (29.7), 19.6 (17.0), 8.3 (13.2), 38.9 (29.7), 19.6 (17.0), 8.3 (13.2), 38.9 (29.7), 19.6 (17.0), 8.3 (13.2) | 82.0 (15.4), 84.7 (17.4), 85.3 (23.0), 91.6 (13.1), 87.7 (17.6), 90.9 (16.7), 16.0 (18.7), 2.5 (8.9), 12.8 (20.6), 10.5 (21.1), 15.8 (24.0), 3.3 (13.5), 8.7 (18.7), 9.6 (19.9), 4.1 (15.0), 86.9 (19.3), 77.0 (20.7), 73.4 (29.0), 71.9 (22.3), 75.6 (25.9), 79.0 (24.2), 31.2 (25.0), 5.7 (14.1), 22.1 (27.1), 19.5 (27.7), 32.8 (34.1), 8.0 (19.0), 11.2 (22.7), 13.8 (24.6), 4.1 (15.0), 75.5 (27.3), 77.0 (21.1), 73.3 (29.2), 71.3 (23.3), 75.6 (25.1), 77.0 (27.4), 29.9 (24.4), 4.7 (11.8), 22.7 (27.3), 18.1 (26.9), 29.3 (30.6), 7.3 (16.1), 11.7 (22.8), 15.4 (25.5), 5.4 (23.4) | 70.5 (19.0), 77.0 (20.7), 73.4 (29.0), 71.9 (22.3), 75.6 (25.9), 79.0 (24.2), 31.2 (25.0), 5.7 (14.1), 22.1 (27.1), 19.5 (27.7), 32.8 (34.1), 8.0 (19.0), 11.2 (22.7), 13.8 (24.6), 4.1 (15.0), 75.5 (27.3), 77.0 (21.1), 73.3 (29.2), 71.3 (23.3), 75.6 (25.1), 77.0 (27.4), 29.9 (24.4), 4.7 (11.8), 22.7 (27.3), 18.1 (26.9), 29.3 (30.6), 7.3 (16.1), 11.7 (22.8), 15.4 (25.5), 5.4 (23.4) | 69.1 (19.3), 77.0 (21.1), 73.3 (29.2), 71.3 (23.3), 75.6 (25.1), 77.0 (27.4), 29.9 (24.4), 4.7 (11.8), 22.7 (27.3), 18.1 (26.9), 29.3 (30.6), 7.3 (16.1), 11.7 (22.8), 15.4 (25.5), 5.4 (23.4) | <0.001 |

Only those who completed two or more questionnaires were included in the analyses.

A higher score on the EORTC QLQ-C30 and EORTC QLQ-CR38 functional scales, global QOL scale and single items (i.e. body image, sexual functioning, sexual enjoyment and future perspective) represent a higher level of function. For the symptom scales and single items, a higher score represents a higher level of symptoms.

*p-Value represents the difference between the four personality groups; cClinically relevant differences between the Type D group (NA+SI+) and the reference group (NA-SI-) and between the NA only group (NA-SI-) and the reference group (NA-SI-) according the guidelines [20]; this question was filled out by a small number of patients.
they reported more insomnia, diarrhea, constipation, gastrointestinal-, defecation-, stoma-related-, and female sexual problems, and a worse body image and future perspective compared to the reference group, even after controlling for sociodemographic and clinical variables. High Na patients reported similar poor outcomes as Type Ds. However, they also reported a lower social functioning and more fatigue, pain, micturition- and financial problems.

**Discussion**

CRC survivors with a Type D personality and those with high Na reported a significantly worse HRQoL and disease-specific health status and these differences were quite stable over time. Furthermore, linear mixed effects models showed that Type D patients had a lower quality of life, cognitive and emotional functioning, and sexual enjoyment, and they reported more insomnia, diarrhea, constipation, gastrointestinal-, defecation-, stoma-related-, and female sexual problems, and a worse body image and future perspective compared to the reference group, even after controlling for sociodemographic and clinical variables. High Na patients reported similar poor outcomes as Type Ds. However, they also reported a lower social functioning and more fatigue, pain, micturition- and financial problems.
Continuous variables are grand-mean centered; Only those who completed two or more questionnaires were included in the analyses. Other studies also suggested that the NA component of Type D is the key predictor of subjective health outcomes in both healthy [27] and cardiac populations [28]. The specific combination of NA and SI in the Type D construct may be more important regarding the risk of adverse cardiac events. A recent study in 541 patients with coronary artery disease showed that Type D personality was associated with cardiac death and myocardial infarction, while patients with high NA or SI alone were not at an increased risk [29].

The present study has some limitations. Although we had sociodemographic and clinical information of non-respondents, it remains unknown whether they declined to participate because of poor health. Furthermore, personality traits like neuroticism or a low sense of coherence are also known to exert influence on HRQoL and we did not take them into account. Finally, although Type D personality is a stable construct [30], and although this is a prospective study, our analyses limit the determination of causal association between personality and patient-reported outcomes as baseline data on these outcomes are unknown. The strengths of this study are...
that we assessed HRQoL and disease-specific health status prospectively in a large population-based setting which provides information on the persistence of these constructs over time in a representative group of CRC patients in daily practice.

Type D personality and high NA were associated with poor HRQoL and disease-specific health status among survivors of CRC, even after controlling for sociodemographic and clinical variables. By taking a patient’s personality into account, this study offers a different view on personalized medicine. Evaluating HRQoL and disease-specific health status according to personality is of great value as this informs about the disease burden and treatment-related effects directly from the patients’ perspective. This information will help clinicians to inform CRC patients about potential late side effects. Furthermore, this can possibly lead to the development and evaluation of strategies for tailored long-term management and support for survivors on the basis of a more individualized approach, as a function of stable differences in coping with chronic medical conditions. For example, mindfulness-based stress reduction may reduce levels of negative affectivity and social inhibition [31]. Paying attention to the recognition of NA seems warranted as these patients reported a worse HRQoL and disease-specific health status.

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