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The applicability of a weight loss grading system in cancer cachexia: a longitudinal analysis

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Abstract

Background A body mass index (BMI) adjusted weight loss grading system (WLGS) is related to survival in patients with cancer. The aim of this study was to examine the applicability of the WLGS by confirming its prognostic validity, evaluating its relationship to cachexia domains, and exploring its ability to predict cachexia progression.

Methods An international, prospective observational study of patients with incurable cancer was conducted. For each patient, weight loss grade was scored 0–4. Weight loss grade 0 represents a high BMI with limited weight loss, progressing through to weight loss grade 4 representing low BMI and a high degree of weight loss. Survival analyses were used to confirm prognostic validity. Analyses of variance were used to evaluate the relationship between the WLGS and cachexia domains [anorexia, dietary intake, Karnofsky performance status (KPS), and physical and emotional functioning]. Cox regression was used to evaluate if the addition of cachexia domains to the WLGS improved prognostic accuracy. Predictive ability of cachexia progression was assessed by estimating proportion of patients progressing to a more advanced weight loss grade.

Results One thousand four hundred six patients were analysed (median age 66 years; 50% female, 63% KPS ≤ 70). The overall effect of the WLGS on survival was significant as expressed by change in −2 log likelihood (P < 0.001) and persisted after adjustment for age, sex, and cancer type and stage (P < 0.001). Median survival decreased across the weight loss grades ranging from 407 days (95% CI 312–502)—weight loss grade 0 to 119 days (95% CI 93–145)—weight loss grade 4. All cachexia domains significantly deteriorated with increasing weight loss grade, and deterioration was greatest for dietary intake, with a difference corresponding to 0.87 standard deviations between weight loss grades 0 and 4. The addition of KPS, anorexia, and physical and emotional functioning improved the prognostic accuracy of the WLGS. Likelihood of cachexia progression was greater in patients with weight loss grade 2 (39%) than that with weight loss grade 0 (19%) or 1 (22%).

Conclusions The WLGS is related to survival, cachexia domains, and the likelihood of progression. Adding certain cachexia domains to the WLGS improves prognostic accuracy.

Keywords Neoplasms; Cachexia; Classification; Weight loss; Survival; Nutritional status
Introduction

Cancer cachexia was first described by Hippocrates as severe weight loss and a sign of impending death. It is considered to be one of the most distressing aspects of advanced cancer, resulting in progressive functional impairment and psychological distress, and it impedes the delivery of anticancer treatment. About 80% of patients with advanced cancer experience weight loss, which is a key component of cancer cachexia. Cachexia contributes to about 20% of cancer-related deaths. Despite this, there has been relatively little research in this area, resulting in a failure to advance treatment and a therapeutic nihilism that cachexia is an inevitable and untreatable consequence of advanced cancer.

One of the barriers to cachexia research has been the lack of an agreed definition and classification system. The latter is fundamental as cancer cachexia is not a single entity but a syndrome with various stages. Various definitions have been used, resulting in heterogeneous research populations, making comparison of findings difficult.

A major step in advancing the research agenda in cachexia was taken in 2011 when a consensus-based cachexia definition was published. Cachexia was defined as a ‘multifactorial syndrome characterized by an ongoing muscle loss (with or without fat loss) that cannot be fully reversed by nutritional support and leads to progressive functional impairment’. Integral was the idea that cachexia is a progressive process from an early to a late stage, and the stages of pre-cachexia, cachexia, and refractory cachexia were proposed (Figure 1). Diagnostic criteria, based predominantly on weight loss and body mass index (BMI), were assigned to the cachexia stage, whereas for the other stages, only suggestive characteristics were presented. Further work was recommended to validate the definition and classification.

Another important step forward was made when Martin et al. confirmed that severity of weight loss depends on the concurrent depletion of fat and muscle reserves. They showed that weight loss, adjusted for concurrent BMI, predicted survival; patients with a low degree of weight loss and a high BMI had the best prognosis, and those with a high degree of weight loss and a low BMI had the worst prognosis. By combining weight loss and BMI, they produced and validated a weight loss grading system (WLGS) ranging from 0 to 4, with each weight loss grade predicting survival independently of cancer type and stage, age, sex, and performance status.

The WLGS was not intended as a classification system of cancer cachexia. However, because weight loss is a key component of the syndrome, the WLGS could still potentially be used to classify cancer cachexia. In order to test this hypothesis, concurrent validity in relation to established cachexia domains needs to be demonstrated. Further, because cachexia is a trajectory from pre-cachexia to refractory cachexia, a classification system should be able to predict which patients are at risk of having cachexia progression. This is of particular interest early in the cachexia trajectory.

Thus, the present study had three aims: (i) to confirm the prognostic validity of the WLGS in an independent, prospective cohort of patients with incurable cancer; (ii) to evaluate the concurrent validity of the WLGS in relation to cachexia domains (anorexia, dietary intake, performance status, and physical and emotional functioning) and to explore if adding these domains to the WLGS improves prognostic accuracy; and (iii) to evaluate if the WLGS predicts cachexia progression.

Methods

Patients and study design

Between April 2011 and October 2013, 1739 patients from 30 centres across Europe (27), Canada (2), and Australia (1) were included in the European Palliative Care Cancer Symptom...
study. This was a prospective observational study by the European Palliative Care Research Centre (PRC) and the European Association for Palliative Care (EAPC) Research Network with the aim of improving the understanding of the development of symptoms and how these symptoms may best be assessed and classified in order to improve symptom management. Eligible patients met the following key criteria: ≥18 years of age, with incurable cancer, enrolled in a palliative care programme, and available for at least one follow-up registration. All patients provided written informed consent.

**Data collection and weight loss grading**

Patients were assessed at baseline and approximately every 4 weeks for at least three follow-up visits or until death. The following information was collected: patient demographics, height, current body weight, and patient-reported weight loss in the 6 months prior to inclusion. Weight loss at subsequent visits was computed by adding measured weight change to baseline-reported weight loss. BMI was recalculated at every visit based on current body weight. Weight loss grade was assessed and given a score of 0–4 by combining weight loss and BMI (Table 1).12

### Table 1. Grading of weight loss (0–4) based on percentage weight loss and current body mass index

<table>
<thead>
<tr>
<th>Weight loss (%)</th>
<th>&gt;=28</th>
<th>25–27.9</th>
<th>22–24.9</th>
<th>20–21.9</th>
<th>&lt;20</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;2.5</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>1</td>
<td>3</td>
</tr>
<tr>
<td>2.5–5.9</td>
<td>1</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>6–10.9</td>
<td>2</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>11–14.9</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td>&gt;15</td>
<td>3</td>
<td>4</td>
<td>4</td>
<td>4</td>
<td>4</td>
</tr>
</tbody>
</table>

**Table 2 includes the items from the Patient-Generated Subjective Global Assessment (PG-SGA)14 and the European Organization for Research and Treatment of Cancer Quality of Life Questionnaire (EORTC-QLQ) C15 PAL15 that were used to collect data pertaining to the cachexia domains for nutrition and functional and psychosocial effects. Both instruments are commonly used and well validated patient reported outcome measurements. The clinical meaning of PG-SGA scores is listed in Table 2. Regarding the EORTC-QLQ, a difference in score of ≥20 was considered a definite clinical significant difference, a difference in score of 10–20 was considered a moderate difference, and a difference in score of 5–10 was considered a small difference. In addition, health care personnel-reported performance status (Karnofsky scale) was assessed due to its long standing importance in cancer prognostication. The Karnofsky scale ranges from 0 (Dead) to 100 (normal, no complaints, and no evidence of disease).

### Statistical considerations

To evaluate the prognostic validity of the WLGS, time to death or last known date to be alive was calculated and Kaplan–Meier curves were plotted for each grade. Hazard
ratios (HRs) were calculated by using Cox proportional hazard methods. Adjustments for possible differences in age, sex, and cancer type (digestive organ cancer, respiratory organ cancer, breast cancer, cancer of urinary tract and male genitalia, gynaecological cancer, haematological cancer, and others) and stage (local, locally advanced, or metastatic/disseminated) between weight loss grades were made. Adjustments for differences in performance status between grades were not performed as a worsening in this parameter is an expected consequence of cachexia progression.

To investigate the baseline differences between weight loss grades 0 and 4 in terms of severity of appetite loss, dietary intake, performance status, fatigue, and physical and emotional functioning, one-way analysis of variance was used. A linear test for trend was applied to confirm if severity increased with grade. Where normality could not be assumed, the non-parametric Kruskal–Wallis test and Jonckheere–Terpstra test were used instead. To illustrate the magnitude of change over the entire spectrum of the WLGS, absolute change in mean values between grade 0 and grade 4 was calculated. To rank which domain had the greatest relative change, the absolute change in mean values of each domain was divided by its standard deviation.

To examine if the cachexia domains improved the prognostic validity of the WLGS, a survival prediction model was built by using Cox proportional hazard methods. The general prognostic factors age, sex, and cancer type and stage were added first, and then a forward stepwise (likelihood ratio) method was used to add the cachexia domains (appetite loss, dietary intake, performance status, fatigue, and physical and emotional functioning) and the WLGS.

To assess the likelihood of progression of cachexia during follow-up, a longitudinal analysis was performed. Weight loss grade was assessed at every follow-up visit and the proportion of patients progressing to higher grades, improving to lower grades, or dying after 1, 2, and 3 months was calculated.

High attrition resulting in missing data was expected because of patient deterioration. Therefore, a sensitivity analysis replacing missing values with extreme values was performed to assess the robustness of the longitudinal analysis. This significantly altered the results (data not shown), and thus, for patients alive but unable to attend follow-up, imputations of likely values for missing data on body weight were performed by using the iterative estimation and maximization algorithm with auxiliary

<table>
<thead>
<tr>
<th>Table 3. Baseline characteristics of the total population and by weight loss grade</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total</td>
</tr>
<tr>
<td>n</td>
</tr>
<tr>
<td>Median age [years] (IQR)</td>
</tr>
<tr>
<td>Sex</td>
</tr>
<tr>
<td>Female</td>
</tr>
<tr>
<td>Male</td>
</tr>
<tr>
<td>KPS ≤70</td>
</tr>
<tr>
<td>&gt;70</td>
</tr>
<tr>
<td>Principal cancer diagnosis</td>
</tr>
<tr>
<td>Cancer of the digestive organs</td>
</tr>
<tr>
<td>Cancer of the respiratory organs</td>
</tr>
<tr>
<td>Breast cancer</td>
</tr>
<tr>
<td>Other cancers</td>
</tr>
<tr>
<td>Urinary cancer or cancer of the male genitalia</td>
</tr>
<tr>
<td>Gynaecological cancer</td>
</tr>
<tr>
<td>Haematological cancer</td>
</tr>
<tr>
<td>Stage</td>
</tr>
<tr>
<td>Local</td>
</tr>
<tr>
<td>Locally advanced</td>
</tr>
<tr>
<td>Metastatic</td>
</tr>
<tr>
<td>Median time since diagnosis [months] (IQR)</td>
</tr>
<tr>
<td>Current oncolgic treatment</td>
</tr>
<tr>
<td>No treatment</td>
</tr>
<tr>
<td>Chemotherapy</td>
</tr>
<tr>
<td>Radiotherapy</td>
</tr>
<tr>
<td>Hormonal therapy</td>
</tr>
<tr>
<td>Other</td>
</tr>
<tr>
<td>Mean weight loss (6 months)[%] (SD)</td>
</tr>
<tr>
<td>BMI [kg/m²] (SD)</td>
</tr>
</tbody>
</table>

Abbreviations: BMI, body mass index; IQR, interquartile range; KPS, Karnofsky performance status; SD, standard deviation.
variables sex, height, body weight, weight loss, and appetite loss. This was done in order to minimize potential bias that could arise by simply ignoring missing assessments. A moderate effect of imputations was observed, with the largest difference seen in patients with weight loss grade 2, where it led to a computational increase in risk of cachexia progression of 9 percentage points compared with the non-imputed dataset.

Statistical analyses were performed with IBM SPSS Statistics for Windows, version 21, IBM Corp, Armonk, NY, USA.

Results

A total of 1406 patients had data available on BMI and weight loss at baseline; those who did not were excluded (n = 333). Patient characteristics are shown in Table 3. Median [IQR] age was 66 [57–74] years, 705 (50%) were female, and most were outpatients (1136, 81%) and had a KPS ≤ 70 (886, 63%). The 333 patients who were excluded were significantly older (median age 70 vs. 66), had a poorer performance status (KPS ≤ 70, 83% vs. 63%) and a shorter median time since diagnosis (13 vs. 19 months), and were more likely to be hospice or nursing home inpatients (25% vs. 6%).

Of the 1406 patients included, 574 (41%) patients completed 3 months of follow-up, while 259 (18%) died. A total of 573 (41%) were lost to follow-up: 239 (17%), 126 (9%), and 208 (15%) after the first, second, and third visits respectively. Although several patients were lost to follow-up after the baseline visit, survival data was still available for a total of 1327 patients.

Prognostic validity

Survival worsened with increasing weight loss grade (Figure 2). The overall effect of the WLGS on survival was significant as expressed by change in −2 log likelihood (P < 0.001) and persisted after adjustment for age, sex, and cancer type and stage (P < 0.001).

Median (95% CI) survival ranged from 407 (312–502) days in weight loss grade 0 to 119 (93–145) days in weight loss grade 4 (Table 4). Adjusted HR ranged from 1.2 (P = 0.20) in weight loss grade 1 to 2.2 (P < 0.001) in weight loss grade 4 (Table 4).

Concurrent validity in relation to other cachexia domains and impact on prognostic accuracy

There was a worsening of all cachexia domains with increasing weight loss grade (P < 0.001) (Figure 3). Between grades 0 and 1, the severity of the cachexia domains was similar with overlapping confidence intervals.

The magnitude of the differences between grades 0 and 4 was greatest for food intake. The mean score worsened from 0.4 to 1.4 (0.87 SD), followed by appetite loss 18 to 48 (0.86 SD), fatigue 42 to 62 (0.69 SD), physical functioning 72 to 54 (0.64 SD), KPS 71 to 63 (0.46 SD), and emotional functioning 69 to 63 (0.29 SD).

When performance status (P < 0.001), physical functioning (P < 0.001), emotional functioning (P = 0.004), and appetite loss (P = 0.005) were added to the WLGS (P < 0.001), the accuracy of survival prediction improved. Of note was that the magnitude of improvement due to performance status (Karnofsky) or physical functioning (EORTC-QLQ C15 PAL) depended on which of the two factors were added first (data not shown). This indicates some collinearity between the two. However, they were both highly significant when present together, indicating some degree of independent contribution, so both were kept in the model.

Predicting the progression of cachexia

Figure 4 is based on the imputed dataset and presents the likelihood of surviving patients progressing or improving according to the WLGS dependent on their baseline weight loss grade. There was a slightly higher tendency of progression of cachexia in grade 1 compared with grade 0, while the risk was considerably higher in grade 2. The tendency towards improvement declined over the last three grades.
After 3 months, the proportions of patients with weight loss grades 0 to 3 progressing to a more advanced weight loss grade were 19%, 22%, 39%, and 19%, respectively. Conversely, those with weight loss grade 1 to 4 improving to a lower weight loss grade were 5%, 13%, 6%, and 4%, respectively.

**Discussion**

The present study has confirmed the prognostic validity of the WLGS in an independent, prospective cohort of patients with incurable cancer. Moreover, the WLGS also has concurrent validity in relation to established cachexia domains and predicts cachexia progression. It has also been demonstrated that the addition of cachexia domains to the WLGS serves to improve its prognostic accuracy.

The present study has several implications for the management of cancer cachexia. Firstly, the findings support the observation made previously by Martin et al. that survival significantly worsens with increasing weight loss grade and are consistent with several other publications showing that patients with involuntary weight loss have a poor prognosis. Looking at weight loss grades 1 and 2 individually, survival did worsen compared with that at weight loss grade 0. However, the difference did not reach...
statistical significance, although weight loss grade 2 showed a tendency towards significance. Failure to reach significance may be due to the smaller sample in the present study compared with the study by Martin et al.12

Secondly, the WLGS not only predicts survival but the present findings suggest that it may also be useful in cachexia classification as it has concurrent validity in relation to established cachexia domains. Of these, dietary intake and appetite loss were the factors most strongly related to increasing grade, which is not surprising given that these factors are central in the pathophysiology of cancer cachexia.20 From grades 0 to 4, dietary intake changed from a mean value of 0.4, representing a near normal intake of nutrients (normal intake is score 0), to 1.4, which represents a reduced intake of normal food (or feeding by nasogastric tube or vein) (score 1) or intake of little solid food (score 2).14 KPS changed from a mean value close to 70 to close to 60, the difference being the ability to fully care for oneself vs. requiring occasional assistance.17 Appetite loss and fatigue changed by a score of 20, which is considered to be a definite clinical significant change. Physical functioning changed by a score of 18, which is considered to be a moderate change, and emotional functioning changed by a score of 6, which is considered to be a small change.16

Performance status, appetite loss, and physical and emotional functioning improve prognostication of survival independently of the WLGS. This suggests that they can be used in combination with the WLGS to further improve the prognostic ability of the system. Future research should examine their full potential to classify cancer cachexia.

The third and, arguably, the finding with the greatest potential clinical relevance was that weight loss grade was predictive of the likelihood of cachexia progression. The risk of progression was considerably higher in weight loss grade 2 compared with that in weight loss grade 0 or 1. This suggests that more patients have started on the cachexia trajectory in weight loss grade 2 than in weight loss grade 0 or 1. Notably, the percentage of patients receiving chemotherapy was similar in patients with weight loss grades 0–2 (47%, 55%, and 50%, respectively), so less anti-cancer treatment does not seem to explain the increased risk of cachexia progression in weight loss grade 2. Regarding the two most severe grades of the WLGS (grades 3 and 4), the probability of improvement decreased, reflecting that irreversibility increases as cachexia becomes more advanced. This is consistent with the findings of Prado et al. who found that weight gain is unlikely to occur in the last 90 days of advanced cancer patients’ lives.21 Overall, the risk of progression to more severe grades supersedes the rate of improvement at any grade (with the obvious exception of grade 4), confirming the progressive nature of cachexia.

So how might these findings influence practice? One such way is the early identification of patients where cachexia interventions should be implemented. As less than a quarter of patients with grades 0 and 1 have cachexia progression, it would seem sensible to start cachexia intervention for patients with grade 2 where the likelihood of progression is greater. That way, one might avoid over-treating many patients. This strategy is supported by the finding that patients with grade 2 have a higher symptom load (appetite loss and fatigue), lower dietary intake, and poorer function than patients with lower weight loss grade. Thus, grade 2 seems to fit the description of pre-cachexia in the consensus definition.11 At the other end of the scale, patients with grade 4 had a median survival of a little over 3 months and a mean KPS close to 60. This is in accordance with the
description of refractory cachexia in the consensus definition, and aggressive treatment attempts against cachexia, as well as chemotherapy, should be avoided in this group of patients. Instead, the attention should be given to palliation and immediate symptom relief.

The main limitation of this study was the expected, but considerable, attrition that may have affected the longitudinal analyses. Presumably, the high dropout rate was caused by patients deteriorating and becoming too weak to continue participation. This was addressed by performing imputations of missing values aiming to reduce the risk of bias. Another limitation was the lack of information on important cachexia domains such as systemic inflammation and objective measures of muscle and fat mass. Systemic inflammation is interesting because it is considered a driver of cancer cachexia, and markers of inflammation could potentially be important in diagnosing cachexia. Measures of muscle and fat mass could have explained whether observed weight gain was due to improvement of cachexia or due to accumulation of fluids or shifts in body composition. Nevertheless, concurrent validity could be evaluated with regard to important cachexia domains such as appetite, food intake, and physical function. Furthermore, because cachexia development is likely to be affected by the response to chemotherapy, nutrition, and other treatments of cachexia, one cannot claim that this study describes the natural development of cachexia in an untreated population. Nevertheless, it describes the development of cachexia in a population without systematic intervention against cachexia. The strengths of this study are the longitudinal design and the large number of patients included from several countries, both rare in palliative research.

**Conclusion**

Our findings support that the WLGS predicts survival and is in keeping with the cancer cachexia phenotype. Furthermore, the addition of key cachexia domains improves prognostic accuracy of the WLGS. Prospective clinical trials should examine the WLGS’s ability to stratify cachexia treatments. This could have significant implications for clinical practice and challenge the widely accepted paradigm that cancer cachexia is an inevitable consequence of advanced disease.

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Conflicts of interest

F.S. has had punctual advisorships for Acacia, ACRAF, Amgen, Baxter, Celgene, Danone, Fresenius, GlaxoSmithKline, Grünenthal, Helsinn, ISIS Global, Millennium/Takeda, Mundipharma, Novartis, NovoPharm, Nycomed, Obezia, Otsuka, Ono, Pharm-Olam, Pfizer, Psioxus, PrIIME, Santhera, Sunstone, Teva, and Vifor. He has received unrestricted industry grants for clinical research from Celgene, Fresenius, and Helsinn. He has participated in a clinical cachexia trial lead by Novartis. O.M.V., D.B., A.W., P.F., V.E.B., M.J.H., S.K., B.L., and T.S.S. declare that they have no conflicts of interest.

References