Cancer-related fatigue is the most prevalent and distressing symptom experienced by adolescents and young adults (AYAs). An electronic survey was undertaken to ascertain current fatigue management and perceptions of its effectiveness. Eighty-five percent of responders (68/80) experienced fatigue, and it was worse more than 1 year after cancer treatment ended, compared to <1 year ($p = 0.007$). Forty-one percent received no fatigue management. Although advice to exercise was the most frequent intervention, the greatest impact of fatigue was on the ability to exercise and most did not find exercise advice helpful. Early intervention is warranted, supporting AYAs to persevere with increasing activity.

Keywords: fatigue, symptom control, quality of life, late effects

Introduction

Fatigue is one of the most prevalent, severe, and distressing symptoms experienced by adolescents and young adults (AYAs) with cancer.\(^1,2\) Healthy adolescents are inherently susceptible to fatigue. The developmental changes of adolescence mean that longer sleep times are needed, while accompanying shifts in circadian rhythm result in later bedtimes.\(^3\) Sleep is further hindered by use of light-producing electronic devices in the bedroom, social commitments, and alcohol consumption.\(^3,4\) AYA cancer patients experience even more severe fatigue than their healthy counterparts.\(^5–7\) Young age and a cancer diagnosis, dual risk factors for fatigue, combine to create a particularly significant problem in this patient group.

Despite this, fatigue is a neglected symptom in AYAs, with no interventional studies focusing on fatigue management at this young age. Research evaluating interventions in older adults has begun to accumulate over the last decade. The most promising approaches are exercise and psychosocial interventions such as education and treatment of concurrent symptoms, with little evidence for pharmacological management.\(^8–11\) AYAs have unique needs that differ from those of adults and children, and it is well recognized that research outcomes in older adults cannot be extrapolated to the younger population. This has led to an international drive to develop evidence-based age-appropriate care.\(^12,13\)

Preparatory research is needed to support the development of an AYA fatigue intervention that meets their needs. Although qualitative studies have provided insight into the unmet needs of this young group,\(^14–17\) patients’ needs have not been prioritized quantitatively, and there has been no evaluation of the impact on carers.\(^1\) Furthermore, current practice in fatigue management is unknown. The objectives of this multicenter survey were to quantify the impact of fatigue on young patients and their carers, to find out how fatigue is currently being managed in the United Kingdom, and to ascertain perceptions of the effectiveness of such management.
Methods

Procedure

Participants were recruited from three Teenage and Young Adult Principal Treatment Centres (TYA PTCs) in the United Kingdom, and were required to have been diagnosed or treated with cancer at age 13–24 and to be 16–27 at the time of the survey.18

Electronic survey methods were used, following advice from TYA PTC service users. After priming by email, poster, and social media, a personalized email of invitation was sent to potential participants, containing a unique token number and links to the Participant Information Sheet and survey. Nonresponders were ascertained by noting unused tokens, and one reminder email was sent to nonresponders after 7 days. Approval was given by the National Research Ethics Service Committee East of England—Cambridgeshire and Hertfordshire (ref. 15/EE/0037).

Outcome measures

Demographic data were collected from both responders and nonresponders. Participants completed two validated measures. The PedsQL Multidimensional Fatigue Scale (MFS) is a widely used 18-item scale validated in both adolescents and in young adults; scores are linearly transformed to a 0–100 scale, with lower scores representing worse fatigue.19 The Quality of Life Visual Analog Scale (QOL-VAS) is a single-item measure of quality of life that captures a clinically significant change in quality of life more readily than multiple-item measures.20,21

The severity of 11 symptoms and the impact of fatigue were quantified using a 5-item verbal rating scale: “not at all,” “a little bit,” “somewhat,” “quite a bit,” and “very much.” The successfullness of each treatment received for fatigue was rated using a 4-item scale: “entirely successful,” “somewhat successful,” “helped only a little,” and “no effect.” A “successfullness score” for fatigue treatment was generated by attributing 0, 1, 2, and 3 to the ratings “no effect,” “helped only a little,” “somewhat successful,” and “entirely successful,” respectively, for each treatment received and calculating a mean score. The former two ratings defined an unsuccessful treatment and the latter two a successful one. Selection of single or multiple best responses was used for a number of further questions, including defining the two symptoms most affecting daily life, views on the cause of the fatigue, and details of fatigue treatments received.

Participants indicating that fatigue had not affected them over the past month only provided demographic information and recorded the severity of the 11 symptoms, the 2 symptoms most affecting daily life, and the QOL-VAS. SPSS v22 was used for all analyses.

Results

The survey link was sent to 197 eligible patients and 80 responded, equating to a 41% response rate. Across all data entry points, 1.8% were missing.

Characteristics of participants

There were two main differences between survey responders and nonresponders. First, no patients currently receiving cancer treatment participated. Second, significantly more females than males responded, with 67% of responders and 47% of nonresponders being female. However, after excluding patients in the nonresponder group still receiving treatment, to compare responders and nonresponders off

<table>
<thead>
<tr>
<th>Table 1. Comparison of Responder and Nonresponder Demographic Data</th>
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<tbody>
<tr>
<td><strong>Responders</strong> (n = 80)</td>
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<tr>
<td>-------------------------</td>
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<tr>
<td>Off/on treatment, n (missing data)</td>
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<tr>
<td>Age at survey in years, mean (SD, range)</td>
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<td>Age at diagnosis in years, mean (SD, range)</td>
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<td>Months since diagnosis, median (IQR)</td>
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<td>Months since last treatment, median (IQR)</td>
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<td>Months treatment duration, median (IQR)</td>
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<td>Gender, n, male (%)</td>
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<tr>
<td>Diagnosis, n (%)</td>
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<tr>
<td>Leukemia</td>
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<td>Lymphoma</td>
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<tr>
<td>Osteosarcoma/Ewing’s</td>
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<td>Brain neoplasm</td>
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<tr>
<td>Other</td>
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<sup>a</sup> p-values comparing preceding column of nonresponders with responders and significant p-values in bold.
<sup>b</sup>Fisher’s Exact Test.
<sup>c</sup>Independent samples t-test.
<sup>d</sup>Mann–Whitney U test.
IQR, interquartile range; SD, standard deviation.
treatment, there was no longer evidence of a gender difference between the two groups (Table 1).

Impact of fatigue

Fatigue was the most prevalent symptom, experienced by 85% (68/80) during the preceding month (Fig. 1). Fatigue was ranked as more severe than each of the other symptoms (Wilcoxon signed-rank test; \( p < 0.001 \), except anxiety \( p = 0.007 \)). The mean fatigue severity of the fatigued participants was 44.3 (standard deviation [SD] 20.5). Fatigue severity was worse more than 1 year after the last cancer treatment (\( M = 39.0 \), SD = 19.7) compared to <1 year (\( M = 53.8 \), SD = 19.7; independent samples \( t \)-test, \( t(56) = 2.8, p = 0.007 \)). Fatigue was worse in females (\( M = 39.6 \), SD = 19.3) than males (\( M = 55.6 \), SD = 19.6; \( t(66) = 3.1, p = 0.003 \)), but was not associated with other demographic variables, including cancer type or treatment duration. In a linear regression analysis, anorexia and low mood explained 50% of the variation in fatigue severity (adjusted \( R^2 = 0.50 \), \( F \) change = 26.60, \( p < 0.001 \)). Gender no longer had a statistically significant relationship with fatigue after adjusting for anorexia and low mood. Fatigue (MFS) was an independent predictor of quality of life, after adjusting for the potential confounders, low mood, and anorexia (adjusted \( R^2 = 0.42 \), \( F \) change = 7.13, \( p = 0.01 \)).

All participants, including those not experiencing fatigue, were asked which 2 out of 11 symptoms most affected daily life. Fatigue was chosen more often than any other symptom (Wilcoxon signed-rank tests, \(-7.87 < Z < -4.99\), all \( p\)-values \( < 0.001\)). The greatest impact of fatigue on aspects of daily life was on the ability to exercise, with 74% (50/68) experiencing “somewhat,” “quite a bit,” or “very much” impact. Attending school or work was hindered for 65% and socializing for 56%. Forty-five percent of participants (29/64) felt that their fatigue made the person most involved in their care “somewhat,” “quite a bit,” or “very much” upset, 42% (27/64) felt it caused frustration, and 41% (26/64) believed it limited rest time. Only 13% (8/64) thought that there was no negative impact.

Fatigue management

A minority of fatigued participants (19%, 13/68) believed that “something could be done” about fatigue, with many (38%, 26/68) believing it to be a symptom one “has to live with.” Although 65% (44/68) had talked to a health carer about feeling fatigued; the most common reason for not talking was a belief that nothing could be done (60%, 9/15).

Forty-one percent (28/68) of the fatigued participants had not been recommended fatigue treatments. The remaining participants had been recommended a median of 2.5 treatments (range 1–7), most commonly advice to exercise (27/40) or to rest and relax (22/40). There was no difference in fatigue severity (MFS) between those recommended fatigue treatments and those not. Only blood transfusion was perceived as successful more often than unsuccessful (Fig. 2). Successfulness of fatigue treatment was lower in participants more than 1 year from last cancer treatment compared to those <1 year from treatment (\( t(30) = -2.93, p = 0.006 \)).

All participants, irrespective of whether they had received fatigue treatment, were asked what they personally thought was the single best treatment. Most chose exercise (24%, 14/58), followed by diet/nutrition (12%, 7/58), rest/relaxation (12%, 7/58), naps/sleeping (10%, 6/58), blood transfusion (7%, 4/58), complementary therapies (5%, 3/58), physiotherapy (3%, 2/58), and fatigue group attendance (3%, 2/58). Thirteen of the 16 participants who recommended an active treatment (exercise or physiotherapy) had received an active treatment themselves; those who had received an active treatment were more likely to recommend an active treatment as being better than a rest-focused treatment (Fishier’s Exact Test, \( p = 0.001 \)). However, no evidence was found that those who had received a rest-focused treatment (rest/relaxation, naps/sleeping, or complementary therapy) were more likely to recommend a rest-focused treatment (\( p = 0.62 \)).
Discussion

This survey confirms that fatigue is the single most prevalent and severe symptom experienced by AYAs with a cancer diagnosis and has the most negative impact on daily life. Fatigue persisted long after the end of cancer treatment, being perceived as worse more than 1 year after the end of treatment, compared to during the first year following treatment.

Research to date has focused on the burden of fatigue during cancer treatment. Only one study has shown worse fatigue in AYA cancer survivors than controls,7 and there have been no longitudinal studies evaluating fatigue changes over time.1 One small, cross-sectional observational study has evaluated fatigue according to the time after the end of cancer treatment; there was a suggestion that fatigue in adolescents more than 5 years after treatment (n = 8) could be worse than those 1–2 years after treatment (n = 6).14

There may be a number of reasons for the persistence of fatigue well beyond cancer treatment. Resting is counterproductive as the inactivity caused by fatigue leads to a destructive spiral of muscle deconditioning, loss of cardiorespiratory fitness, and worsening fatigue.8,22 Furthermore, AYA patients are at a formative age that puts them at particular risk of maladaptive behaviors such as activity avoidance.23 Unhelpful health behaviors, such as inactivity, tend to become lifelong habits, increasing the potential for long-term disability. The finding that fatigue severity was worse more than 1 year after the end of treatment may reflect this deconditioning spiral. It could also be that, further from the end of treatment, the expectation that fatigue should be resolving drives a perception of worse fatigue.

These data reveal concerning therapeutic nihilism and under-management of fatigue. Over a decade ago, Stone et al. undertook an influential multicenter survey of older adults with cancer-related fatigue.24 While the proportion of people believing that "something could be done" about fatigue remained comparably low (19% in this survey, 22% in Stone et al.), there is evidence of progress. Sixty-five percent fatigued participants in this survey had talked to a health carer about their fatigue; 59% had received some fatigue treatment and, of these, 68% had received advice to exercise (respective figures in older adults being 48%, 14%, and 4%).24 Exercise was viewed as the best fatigue treatment and was most recommended to others. The fact that most participants found exercise advice unhelpful could be explained by the finding that fatigue has a particularly negative impact on the ability to exercise.

The main limitation of this study is the risk of selection bias. Although a nonresponder analysis was undertaken, fatigue severity was not measured in nonresponders. It is possible that people experiencing fatigue may have been more likely to participate, leading to fatigue overestimation. Greater selection bias further from cancer treatment could have contributed to the findings of worse fatigue and less successful fatigue management in those more than a year after cancer treatment. A longitudinal cohort study, powered to detect changes in fatigue over time, is needed to confirm the study findings.

Nevertheless, this study has a number of implications for clinical practice. The persistence of fatigue, potentially sustained by a spiral of inactivity and deconditioning, suggests that early fatigue intervention is warranted, before maladaptive rest-based behaviors become entrenched at this formative age. Given the impact of fatigue on the ability to exercise, a fatigue intervention must find ways of engaging AYAs in persisting with increasing activity, despite the presence of fatigue. The negative consequences for carers suggest that it would be beneficial to incorporate family support into the intervention.

Considering the high prevalence and detrimental impact of fatigue, it is concerning that there has as yet been no research evaluating age-appropriate interventions. This survey provides evidence for the specific needs of fatigued AYAs and is supporting the design of future research aiming to develop an effective treatment for this important and debilitating symptom.

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