Cancer sleep symptom-related phenotypic clustering differs across three cancer specific patient cohorts

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Summary
Specific sleep disorders have been linked to disease progression in different cancers. We hypothesised sleep symptom clusters would differ between cancer types. The aim of this study was to compare sleep symptom clusters in post-treatment melanoma, breast and endometrial cancer patients. Data were collected from 124 breast cancer patients (1 male, 60 ± 15 years, 28.1 ± 6.6 kg/m\textsuperscript{2}), 82 endometrial cancer patients (64.0 ± 12.5 years, 33.5 ± 10.4 kg/m\textsuperscript{2}) and 112 melanoma patients (59 male, 65.0 ± 18.0 years, 29.1 ± 6.6 kg/m\textsuperscript{2}). All patients completed validated questionnaires to assess sleep symptoms, including the Epworth Sleepiness Scale (ESS), Pittsburgh Sleep Quality Index (PSQI), Insomnia Severity Index (ISI), and Functional Outcomes of Sleep Questionnaire-10 (FOSQ-10). Snoring, tiredness, observed apneas, age, BMI, and gender data were also collected. Binary values (PSQI, ISI, FOSQ), or continuous variables for sleepiness (ESS) and perceived sleep quality (PSQI), were created and sleep symptom clusters were identified and compared across cancer cohorts. Four distinct sleep symptom clusters were identified: minimally symptomatic (n = 152, 47.7%); insomnia-predominant (n = 87, 24.9%); very sleepy with upper airway symptoms (n = 51, 16.3%), and severely symptomatic with severe dysfunction (n = 34, 11.1%). Breast cancer patients were significantly more likely to be in the insomnia predominant or severely symptomatic with severe dysfunction clusters, whereas
melanoma patients were more likely to be minimally symptomatic or sleepy with upper airway symptoms \( (p < 0.0001) \). Endometrial cancer patients were equally distributed across symptom clusters. Sleep symptom clusters vary across cancer patients. A more personalised approach to the management of sleep-related symptoms in these patients may improve the long term quality of life and survival.

**KEYWORDS**

breast cancer; endometrial cancer; melanoma; sleep symptom clusters

## 1 | INTRODUCTION

Cancer patients frequently complain of poor sleep across all stages of their cancer journey. Poor sleep is a broad, poorly defined term, reflecting patient perceptions of difficulty falling asleep, insufficient sleep duration, and interrupted and unrefreshing sleep; all accompanied by a range of daytime symptoms, including fatigue, tiredness, daytime sleepiness, low daytime activity, and depressive symptomology (Colagiuri et al., 2011). Indeed, elements of these symptoms are reported by 35–75% of all cancer patients (Palesh et al., 2013).

Assessing the clinical significance of sleep-related symptoms in cancer cohorts is challenging. While they may arise from the holistic impacts of specific cancer pathologies (e.g., pain), treatment regime side effects, and patient anxiety, there is also the potential that they may reflect a co-morbid specific sleep disorder (e.g., obstructive sleep apnea, insomnia). The differential diagnosis of sleep-related symptoms in cancer patients is fundamental to understanding direct and indirect medical consequences that may arise from a specific sleep disorder afflicting a specific cancer patient.

There is now accumulating evidence suggesting that sleep disorders can influence both the risk and outcomes of cancer. For example, obstructive sleep apnea (OSA), has been linked to histologically more aggressive melanoma subtypes (Martinez-Garcia et al., 2018) and an increase in circulating biomarkers associated with tumour growth (Santamaria-Martos et al., 2018). Other cancers have also been associated with OSA, including lung cancer (Cabezas et al., 2019), brain tumours (Cho et al., 2020), breast cancer (Chang et al., 2014), colorectal cancer (Lee et al., 2017), and prostate and head and neck cancers (Fang et al., 2015). It is thought that these associations may be a consequence of intermittent hypoxia associated with upper airway obstruction during sleep (Almendros & Gozal, 2018). Insomnia, which is prevalent in breast cancer survivors (Lowery-Allison et al., 2017), is thought to raise the risk of breast cancer recurrence (Collins et al., 2017). Circadian rhythm disruption (e.g., shift work) increases cancer risk, especially for breast (Szkiela et al., 2020) (Ball et al., 2016) and endometrial cancers (Viswanathan et al., 2007), although a recent meta-analysis does not support this association (Travis et al., 2016). In addition, poor sleep, of any aetiology, is a predictor of poorer outcomes for cancer patients, with sleep disruption and duration, both short and long, being linked to reduced survival rates for advanced cancer patients (Collins et al., 2017). Given these two-way interactions, there is an obvious need to define and assess sleep symptoms in cancer patients.

Sleep related symptoms are commonly assessed through a range of validated questionnaires. Typically these questionnaires are designed and validated for their ability to provide a risk assessment for a specific sleep disorder (e.g., OSA-STOP BANG or insomnia -Insomnia Severity Index, ISI), or attempt to quantify the level and nature of sleep disturbance (e.g., Pittsburgh Sleep Quality Index, PSQI), or score severity for specific symptoms (e.g., Epworth Sleepiness Scale), or rate the impact of sleep disturbance on daytime function (e.g., Functional Outcomes of Sleep Questionnaire, FOSQ). Conventionally, an overall risk or impact score is calculated and then compared with a validated threshold score to distinguish from population values. While the score-based outcomes have been validated for each specific questionnaire (e.g., risk of OSA), the individual question responses themselves provide a more detailed picture of sleep symptom phenotypes.

Sleep-related symptom cluster analyses have been applied to sleep clinic databases from Iceland (Ye et al., 2014), an international collaboration (Keenan et al., 2018), France (Gagnadoux et al., 2016), Italy (Lacedonia et al., 2016), Greece (Vavougios et al., 2016), and Europe (Saaresranta et al., 2016), and have identified different sleep symptom phenotypes related to specific sleep disorders. Depending

**TABLE 1** Patient characteristics

| Cohort          | Breast cancer | Endometrial cancer | Melanoma | Total | \( p \) |
|-----------------|---------------|-------------------|----------|-------|-------|
| Total N (%)     | 124           | 82                | 112      | 318   |       |
| Gender          | F 123 (99.2) M 1 (0.8) | F 82 (100.0) M 0 (0.0) | F 53 (47.3) M 59 (52.7) | F 258 (81.1) M 59 (18.6) | <0.001 |
| Age (years)     | 60.0 (15.0)   | 64.0 (12.5)       | 65.0 (18.0) | 63.0 (15.0) | 0.004 |
| Systolic blood pressure (mmHg) | 125.0 (23.0) | 131.0 (16.5) | 135.0 (23.0) | 129.0 (23.5) | <0.001 |
| Diastolic blood pressure (mmHg) | 77.0 (12.5) | 79.0 (12.0) | 80.0 (11.5) | 79.0 (11.8) | 0.099 |
| Body mass index (kg/m\(^2\)) | 28.1 (6.6) | 33.5 (10.4) | 29.1 (6.6) | 29.7 (7.5) | <0.001 |
TABLE 2  Responses to symptom-variables in breast cancer, endometrial cancer and melanoma patients

| Symptom variable                          | Level                    | Breast cancer | Endometrial cancer | Melanoma | *p  |
|------------------------------------------|--------------------------|---------------|-------------------|----------|-----|
| ESS Total N (%)                          |                          |               |                   |          |     |
| Not sleepy                               | 69 (55.6)                | 49 (59.8)     | 54 (48.2)         | 0.261    |     |
| Mildly sleepy                            | 41 (33.1)                | 20 (24.4)     | 35 (31.2)         |          |     |
| Moderately sleepy                        | 9 (7.3)                  | 11 (13.4)     | 19 (17.0)         |          |     |
| Severely sleepy                          | 5 (4.0)                  | 2 (2.4)       | 3 (2.7)           |          |     |
| FOSQ Difficulty concentrating on things  | 11 (8.9)                 | 7 (8.5)       | 5 (4.5)           | 0.371    |     |
| Difficulty remembering things           | 20 (16.1)                | 5 (6.1)       | 7 (6.2)           | 0.016    |     |
| Difficulty operating a motor vehicle for short distances | 3 (2.4)   | 2 (2.4)       | 0 (0.0)           | 0.251    |     |
| Difficulty operating a motor vehicle for long distances | 8 (6.5)   | 4 (4.9)       | 8 (7.1)           | 0.821    |     |
| Difficulty visiting family or friends   | 5 (4.0)                  | 4 (4.9)       | 3 (2.7)           | 0.716    |     |
| Relationship with family, friends, or colleagues affected | 3 (2.4) | 2 (2.4)       | 3 (2.7)           | 0.992    |     |
| Difficulty watching a movie             | 10 (8.1)                 | 10 (12.2)     | 10 (8.9)          | 0.618    |     |
| Difficulty being active in the evening  | 18 (14.5)                | 12 (14.6)     | 8 (7.1)           | 0.140    |     |
| Difficulty being active in the morning  | 14 (11.3)                | 9 (11.0)      | 9 (8.0)           | 0.669    |     |
| Desire for intimacy or sex affected      | 18 (14.5)                | 8 (9.8)       | 9 (8.0)           | 0.219    |     |
| ISI Difficulty falling asleep           | 62 (50.0)                | 22 (26.8)     | 28 (25.0)         | <0.001   |     |
| Difficulty staying asleep               | 72 (58.1)                | 31 (37.8)     | 42 (37.5)         | 0.002    |     |
| Waking up too early                     | 59 (47.6)                | 28 (34.1)     | 39 (34.8)         | 0.088    |     |
| Dissatisfied with current sleep pattern  | 93 (75.0)                | 45 (54.9)     | 59 (52.7)         | 0.001    |     |
| Sleep problems noticeable to others    | 38 (30.6)                | 19 (23.2)     | 19 (17.0)         | 0.048    |     |
| Worried or stressed about sleep problems | 44 (35.5)               | 21 (25.6)     | 19 (17.0)         | 0.005    |     |
| Interferes with daily functioning       | 41 (33.1)                | 25 (30.5)     | 26 (23.2)         | 0.234    |     |
| PSQI Cannot get to sleep                | 80 (64.5)                | 37 (45.1)     | 36 (32.1)         | <0.001   |     |
| Wake up in the middle of the night      | 107 (86.3)               | 65 (79.3)     | 88 (78.6)         | 0.245    |     |
| Get up to use the bathroom              | 104 (83.9)               | 63 (76.8)     | 83 (74.1)         | 0.170    |     |
| Cannot breathe comfortably              | 18 (14.5)                | 16 (19.5)     | 7 (6.2)           | 0.019    |     |
| Cough or snore loudly                   | 53 (42.7)                | 28 (34.1)     | 46 (41.1)         | 0.442    |     |
| Feel too cold                           | 35 (28.2)                | 16 (19.5)     | 16 (14.3)         | 0.030    |     |
| Feel too hot                            | 83 (66.9)                | 41 (50.0)     | 37 (33.0)         | <0.001   |     |
| Have bad dreams                         | 25 (20.2)                | 18 (22.0)     | 17 (15.2)         | 0.440    |     |
| Have pain                               | 52 (41.9)                | 25 (30.5)     | 24 (21.4)         | 0.003    |     |
| Overall sleep quality                   | Very good                | 14 (11.3)     | 16 (19.5)         | 0.208    |     |
|                                          | Fairly good              | 63 (50.8)     | 39 (47.6)         | 61 (54.5) |     |
|                                          | Fairly bad               | 37 (29.8)     | 22 (26.8)         | 23 (20.5) |     |
|                                          | Very bad                 | 10 (8.1)      | 5 (6.1)           | 4 (3.6)  |     |
| Sleep medication                        | 17 (13.7)                | 6 (7.3)       | 13 (11.6)         | 0.348    |     |
| Trouble staying awake                   | 10 (8.1)                 | 7 (8.5)       | 6 (5.4)           | 0.626    |     |
| SB Loud snoring                         | 39 (31.5)                | 30 (36.6)     | 41 (36.6)         | 0.597    |     |
| Feel tired, fatigued, or sleepy during daytime | 78 (62.9) | 55 (67.1)     | 55 (49.1)         | 0.023    |     |
| Stop breathing during sleep             | 14 (11.3)                | 8 (9.8)       | 10 (8.9)          | 0.834    |     |

Note: N = number of patients (% of the total number of patients within each cancer cohort), with high severity symptoms in each of the cancer cohort. *p-values not adjusted for multiple comparisons. Bold font indicates statistically significant differences between cohort (p <0.05).

Abbreviations: ESS, Epworth sleepiness scale total scores; FOSQ, functional outcomes of sleep questionnaire; ISI, insomnia severity index; PSQI, Pittsburgh sleep quality index; SB, STOP-BANG.
on the methodology and cluster variables used, these analyses have identified 3–6 different sleep symptom and comorbidity clusters (Bailly et al., 2016; Gagnadoux et al., 2016; Keenan et al., 2018; Lacedonia et al., 2016; Saaresranta et al., 2016; Ye et al., 2014). These phenotypic classifications have then been used to define sub-populations within a specific sleep disorder cohort (Bailly et al., 2016; Keenan et al., 2018; Ye et al., 2014), to explore demographic and comorbidity interactions (Anttalainen et al., 2019; Lacedonia et al., 2016; Saaresranta et al., 2016; Vavougios et al., 2016) and to evaluate treatment response variation (Gagnadoux et al., 2016). This approach has not previously been applied to a cancer patient cohort. Although poor sleep is a common complaint across cancer types, it is not known if sleep-related symptomology differs between or within these specific cancer cohorts. This may occur as a consequence of different interactions between specific cancers and sleep disorders. A recent review has underlined the bi-directional interconnections between sleep and cancer, focusing on associations between specific sleep disorders (according to the International Classifications of Sleep Disorders 3rd Edition (ICSD-3 [American Academy of Sleep Medicine, 2014]) and specific tumours (Mogavero et al., 2021). Understanding individual patient symptomology has the potential to inform personalised therapeutic intervention strategies aimed at alleviating symptoms, improving quality of life and helping to contribute to better outcomes.

The aim of the present study was to measure sleep symptom phenotypes in patients with a history of breast, endometrial, and melanoma cancer. These cancer types were selected based on their putative links with sleep disorders (Martinez-Garcia et al., 2018; Palesh et al., 2013; Viswanathan et al., 2007) and the magnitude of the potential impact. Breast cancer is the leading cause of cancer globally with over 2,000,000 cases annually, endometrial cancer is the most common gynaecological cancer, and melanoma is the deadliest skin cancer, with over 400,000 and 300,000 cases diagnosed per year, respectively (Sung et al., 2021).

2 | METHODS

2.1 | Patients

The study was approved by the Western Sydney Local Health District ethics committee. Written informed consent was obtained from all patients.

Patients were recruited from either breast, gynaecological, or melanoma outpatient cancer clinics at Westmead Hospital, between 2017 and 2019. Patients were eligible if they were > 18 years of age, had a confirmed diagnosis of breast cancer, endometrial cancer, or melanoma and had completed corresponding treatment (e.g., chemotherapy, radiotherapy, or surgery) either 2 months (endometrial cancer patients), 6 months (melanoma patients), or 12 months (breast cancer patients) prior to study participation. Data from some patients have been reported elsewhere (Madut et al., 2021). The timing of recruitment was chosen to mitigate against the acute impacts of a recent diagnosis or treatment regime on study results. Patients were excluded if they were pregnant or had any serious or active medical or psychiatric comorbidities that would interfere with the subjects’ compliance with the protocol or their assessment.

Age, gender, body mass index (BMI), cancer history (obtained through hospital medical records), systolic and diastolic blood pressure, and current medications at baseline were collected for each patient.

2.2 | Symptom questionnaires and variable selection

All patients completed standardised, validated questionnaires to assess subjective sleep symptoms (hereafter known as “symptoms”). These included: Pittsburgh Sleep Quality Index (PSQI), Epworth Sleepiness Scale (ESS), Insomnia Severity Index (ISI), Functional Outcomes of Sleep Questionnaire (FOSQ-10), and STOP BANG (Snoring, Tiredness, Observed apnea, High blood pressure [do you have or are you being treated for high blood pressure?], BMI, age, neck circumference, and gender) questionnaires.

Each of the above questionnaires has an associated and validated scoring system that is used to quantify an overall outcome based on

![FIGURE 1 Estimated response probabilities of binary symptom-variable for each symptom-cluster (minimally symptomatic, cluster 1 = orange; insomnia predominant, cluster 2 = green; very sleepy with upper airway symptoms cluster 3 = purple; and severely symptomatic with severe dysfunction cluster 4 = blue). Sleep quality and sleepiness are not shown as they were not binary variables](image-url)
The individual question responses. The overall questionnaire score data for the patients were not included in the current analysis, although results for some have been reported separately (Madut et al., 2021; Trivedi et al., 2021). For the purposes of the present study, demographic data and individual question response data were extracted from each completed questionnaire, in addition to systolic and diastolic blood pressure, with the resulting database serving as the data set for cluster analysis.

To achieve efficient clustering, based on reported symptoms, responses to all individual questions from ISI and FOSQ-10, questions 5–8 (except 5j) of PSQI (see Appendix S1 for questionnaires), questions 1–3 of STOP-BANG, and ESS total scores (0–24) were used to create individual “symptom-variables”; each symptom-variable representing a specific sleep symptom (i.e. symptoms related to insomnia, sleep-related functional outcomes, sleep quality, obstructed nocturnal breathing, and daytime sleepiness). Symptom variables were coded into binary values (see Appendix Table 1). STOP-BANG variables are binary. The ESS total scores and question 6 of PSQI (sleep quality-very good to very bad), were converted into ordinal variables, by categorising responses into four “bands”· each representing increasing degrees of daytime sleepiness and poorer sleep quality, respectively. This resulted in 31 symptom-variables being generated, with 31 binary and 2 ordinal variables (ESS total scores and PSQI question 6). Patient characteristics included in the cluster analysis were cancer type, gender, age, BMI, and systolic and diastolic blood pressure.
Statistical analyses were conducted using R (version 4.0.2). Descriptive statistics for participant characteristics and symptoms were generated for each cancer type, and differences between cohorts were assessed using separate ANOVA analyses and chi-square tests for continuous and categorical variables, respectively.

Clusters were estimated using patients with five or fewer missing responses on the symptom-variables. Patients were clustered using latent class analysis (LCA), a statistical technique allowing for separation response patterns into mutually exclusive clusters, based on response patterns to questionnaires. Latent class analysis models were fitted using the poLCA package for R (version 1.4.1), using a maximum of 100,000 iterations. Models were fitted with one cluster.
Comparisons of patient characteristics between clusters in the four cluster LCA model

| Patient characteristic     | D      | df                | p*   |
|----------------------------|--------|-------------------|------|
| Cancer cohort              | 9.35   | 2,3905239.7       | <0.001 |
| Gender                     | 7.78   | 2,5747920.6       | <0.0013 |
| Age (years)                | 6.33   | 1,8661680.6       | <0.024 |
| Systolic blood pressure (mmHg) | 0.13   | 1,24919626.4     | 0.72 |
| Diastolic blood pressure (mmHg) | 0.77   | 1,18545045      | 0.57 |
| Body mass index            | 0.16   | 1,16713576        | 0.72 |

Note: *p*-values adjusted for multiple comparisons using the Benjamini-Hochberg procedure for relationship between assigned cluster and patient characteristics. Bold font indicates variables with statistically significant differences between cancer groups.

Abbreviations: D, D statistic from combining test statistics from multiply imputed datasets; df, degrees of freedom obtained from the multiple imputations.

to six clusters. To find the global maximum likelihood solution, each model was estimated 100 times with random starting parameters, and the model with the greatest likelihood was used. Fit statistics including the sample size adjusted Bayesian information criterion (aBIC), Bayesian information criterion (BIC), and Akaike information criterion (AIC) were compared to identify the best model. Model selection was also informed by the interpretability of each model.

Confidence intervals for estimated response probabilities (conditional probability of reporting a symptom, provided being allocated into a certain cluster), and differences between clusters in those probabilities were generated through non-parametric bootstrapping. m = 10,000 bootstrapped datasets with the same size as the original data were generated, by sampling patients with replacement from the full dataset. Latent class models were fitted to each bootstrapped sample, using the same randomly generated initial probabilities used for the original dataset to keep the order of clusters consistent.

Relationships between cluster membership and other variables were assessed using chi-square tests for categorical variables, and ANOVA analyses for continuous variables. Since cluster assignments in latent class analyses are probabilistic, pseudo-class draws were used in which each patient's cluster allocation was sampled 10,000 times, using their posterior probability of cluster assignment. Analyses were repeated in each sample, and the results were combined using rules for combining statistics in multiple imputation.

Multiple comparisons were controlled for using the Benjamini-Hochberg method. A significance level of (p <0.05) was used for all analyses, after adjusting for multiple comparisons.

3 | RESULTS

3.1 Patient characteristics

There were 318 patients included in the study, comprising 124 breast cancer patients, 82 endometrial cancer patients, and 112 melanoma patients. Patient demographic, anthropometric and blood pressure data, and cancer histopathology are summarised in Table 1. Since two of the cancers included occur predominantly or exclusively in women, all but one of the male patients were in the melanoma group. Melanoma patients were older (p <0.004) with comparatively higher systolic blood pressures (p <0.001). Endometrial cancer patients had a significantly higher BMI (p <0.001) compared with the other cancer patient groups.

3.2 Symptoms within cancer cohorts

Compared with the other cancer patients, breast cancer patients had significantly more symptoms in the ISI and PSQI questionnaires (Table 2). Breast cancer patients also had subjective difficulties with remembering things; however, no other differences in functional questions from the FOSQ-10 questions were observed between cancer cohorts (Table 2). There were also no differences in the distribution of sleepiness from the ESS between cancer groups, with few patients complaining of this symptom within any cancer cohort; although breast cancer patients did complain more of daytime tiredness in the STOP-BANG questionnaire (Table 2).

3.3 Symptom clusters

A model of four clusters was the optimal model, based on fit statistics and interpretability of the cluster models tested (See Appendix Table 2).

Based on patterns of estimated response probabilities in each symptom-cluster (Figure 1), cluster 1 (n = 152; cohort proportion = 47.7%), was labelled “minimally symptomatic” as it had the lowest probability of most symptoms (Figure 1) and the highest sleep quality (Figure 2). The majority of patients sorted into this cluster (Figure 3).

Cluster 2 (n = 87; cohort proportion = 24.9%) was labelled “insomnia predominant”, as it had a higher probability of all symptoms than Cluster 1 and had more pronounced insomnia symptoms. Cluster 2 also had a higher probability of patients taking sleep medication (Figure 4). Clusters 3 (n = 51; cohort proportion = 16.3%) and 4 (n = 34; cohort proportion = 11.1%) were labelled “very sleepy with upper airway difficulty” and “severely symptomatic with dysfunction” respectively, as both clusters displayed a high probability of most symptoms in comparison with the other clusters. Cluster 3 had a higher probability of most symptoms and relatively more reports of breathing difficulties during sleep, compared with other
symptom-clusters (Figure 4). Cluster 4 had a high symptom burden for most symptom-variables, with patients reporting more issues with functional outcomes (Figures 1, 4) and “very bad” sleep quality (Figure 2), compared with other symptom-clusters.

While all four symptom-clusters had patients reporting little to no daytime somnolence; Clusters 3 and 4 had a higher proportion of patients with moderate and severe daytime somnolence, respectively (Figure 2).
3.4 | Comparing symptom response patterns between clusters

Despite a sizeable difference in estimated response probabilities for many symptom-variables between the four clusters (Figures 1, 4), there were no significant differences between the response probabilities between the different classes for individual symptoms (all $p \geq 0.05$) (see Appendix Table 3).

3.5 | Patient characteristics within symptom clusters

Descriptive statistics of participant characteristics for each estimated symptom-cluster are shown in Tables 3 and 4. Cancer patients were significantly differently distributed between the four clusters ($p = <0.001$). Overall, breast cancer patients had more severe symptoms and were more likely to be included in Cluster 2 and 4 (Figure 5a). Melanoma patients tended to fall within clusters with low or moderate symptoms (Figure 5a). More males were present in the low or moderate symptom clusters ($p = 0.0013$) (Figure 5b), consistent with the higher proportion of melanoma patients in these clusters. Severely symptomatic patients with severe dysfunction were lower in age ($p = 0.024$), (Figure 5c), compared with other clusters.

There were no differences in BMI, systolic and diastolic blood pressures between the four symptom-clusters (Table 4).

4 | DISCUSSION

This is the first study to implement clustering analysis to quantify and characterise phenotypes of poor sleep within cancer patients. In this study, four distinct sleep symptom clusters were identified: a minimally symptomatic group, an insomnia predominant group, a very sleepy with upper airway symptoms group and a group who were severely symptomatic with dysfunction. Sleep symptom clusters differed between different cancer subgroups, with breast cancer patients more likely to have a phenotype characterised by insomnia or fatigue, with additional impacts on functional outcomes. Melanoma patients were less likely to have any symptoms at all, or have sleepiness with upper airway symptoms, and patients with endometrial cancer were evenly distributed between the different sleep symptom phenotypes.

Cluster analyses facilitate an understanding in the heterogeneity of clinical presentations of a disease, allowing for its improved recognition and understanding of symptom presentations. Variation in clinical presentations has been studied in sleep clinic patients with a sleep study confirmed OSA (Anttalainen et al., 2019; Bailly et al., 2016; Gagnadoux et al., 2016; Keenan et al., 2018; Lacedonia et al., 2016; Saarersanta et al., 2016; Vavougios et al., 2016; Ye et al., 2014). This includes patients with moderate–severe OSA ($AHI \geq 15$ events/hr) in Iceland (Ye et al., 2014), in an international multicentre consortium (Keenan et al., 2018), and in subjects with at least mild sleep-apnea ($AHI \geq 5$ events/hr) in another international multicentre consortium (Saarersanta et al., 2016). Cluster analyses of sleep clinic patients, including comorbidities, demographics, OSA severity and symptoms, have also been performed on a database in Greece (Vavougios et al., 2016) and France (Bailly et al., 2016; Gagnadoux et al., 2016). Each of these OSA-related cluster analyses had produced 3–6 different clusters of sleep symptoms using various combination of questionnaires, demographics, co-morbidities, and sleep studies. These clusters are disturbed sleep with insomnia symptoms (Keenan et al., 2018; Ye et al., 2014), insomnia phenotype (Saarersanta et al., 2016), minimally symptomatic (Keenan et al., 2018; Ye et al., 2014), non-sleepy, non-insomnia phenotype (Saarersanta et al., 2016), and excessive daytime somnolence (Keenan et al., 2018; Saarersanta et al., 2016; Ye et al., 2014). These clusters were expanded to upper airway symptoms with sleepiness (Keenan et al., 2018) and upper airway symptoms dominant (Keenan et al., 2018), and a group with both insomnia and excessive daytime sleepiness (Saarersanta et al., 2016). Other groups have further expanded these clusters using demographic data and comorbidities (Bailly et al., 2016; Vavougios et al., 2016).

We have used validated questionnaires to assess insomnia symptoms, sleep quality (PSQI), sleepiness (ESS), OSA risk (STOP-BANG), and the short form functional outcomes of sleep (FOSQ-10), as well as demographics and blood pressure to describe sleep symptom clusters in a cancer cohort. Although these questionnaires are not identical to symptom assessments in other cluster studies, and were performed in a cancer cohort rather than in patients with a diagnosis of OSA, the sleep symptom clusters identified in this study of cancer groups are similar to those identified in OSA symptom cluster studies (Keenan et al., 2018; Saarersanta et al., 2016; Ye et al., 2014).

We have recently demonstrated that the prevalence of moderate–severe OSA in women with breast or endometrial cancer ($AHI > 15$ events/hr) is more than 57% in both groups (Madut et al., 2021). A high prevalence of at least moderate OSA in two of the included cancer cohorts, as a consequence of shared risk factors in an obese, older population, helps to explain similarities between OSA symptom cluster studies (Keenan et al., 2018; Saarersanta et al., 2016; Ye et al., 2014) and the sleep symptom clusters described in this study, despite the use of different inclusion requirements and methodology.

We also found demographic differences in the clusters, as have others (Bailly et al., 2016; Vavougios et al., 2016). Younger patients were more likely to have severe symptoms with severe dysfunction, and men were more likely to have minimal symptoms. Unlike previous OSA cluster analyses, which are up to 89% male (Bailly et al., 2016; Keenan et al., 2018; Lacedonia et al., 2016; Saarersanta et al., 2016; Vavougios et al., 2016; Ye et al., 2014), our cohort was composed predominantly (81%) of women. However, men were more likely to have minimal symptoms in our study, similar to the previously described non-insomnia, non-sleepy cohort (Saarersanta et al., 2016). Previous reports have suggested that younger OSA patients are more likely to report daytime sleepiness (Saarersanta et al., 2016), in contrast to our findings. The majority of participants in the cancer cohort are older, likely explaining the difference in findings. We also found no
differences in systolic or diastolic blood pressure, or BMI between sleep symptom clusters, in contrast to others who have demonstrated variations in co-morbidities across different clinical OSA phenotypes (Saaresranta et al., 2016). Thus, although we have performed a cluster analysis in a very different clinical population (mostly female cancer patients), we demonstrated similar sleep clusters to those described in OSA clinic populations. This is not surprising as we have used sleep symptom questionnaires in a group with a likely high prevalence of OSA (Madut et al., 2021).

Different sleep symptom clusters were present among the cancer cohorts, apart from the endometrial cancer cohort where distribution was equal across the symptom clusters. The reasons for this variation in distribution are not clear, however, interactions between specific sleep disorders and cancer may differ, and these differences may be explained by the presence of sleep disorders other than OSA. Epidemiological studies have linked night shift work, which results in exposure to light at night, and short sleep and sleep disruption as a risk factor for breast cancer and endometrial cancer (Grundy et al., 2013; Kakizaki et al., 2008; Viswanathan et al., 2007). OSA and fluctuations in oxygen related to recurrent upper airway obstruction are associated with an increased risk of cancer mortality and incidence (Marshall et al., 2014; Nieto et al., 2012). Associations between OSA severity and melanoma have been shown in prospective studies (Martinez-Garcia et al., 2018), however, these associations have not been found for breast cancer (Campos-Rodriguez et al., 2018; Madut et al., 2021) or endometrial cancer (Madut et al., 2021). Short sleep duration and insomnia symptoms reportedly pre-date the development of breast cancer in women (Phipps et al., 2016). The different distribution of sleep symptom clusters across cancer cohorts may be a consequence of different interactions between sleep disorders and cancer biology.

Previous sleep symptom cluster studies have demonstrated differences in treatment responses between clusters for patients with the same sleep disorder. For example, OSA patients with insomnia symptoms were less likely to be compliant with positive airway pressure treatment (CPAP) in one study (Gagnadoux et al., 2016), but both insomnia and sleepiness symptom clusters, although not minimally symptomatic clusters, were likely to have a symptomatic response to CPAP therapy in another study (Pien et al., 2018). The identification of an insomnia symptom cluster in the present study of cancer patients, which was particularly prevalent in the breast cancer cohort, but also prevalent in the endometrial cohort, highlights the importance of recognising that these women may have OSA as an explanation for insomnia symptoms, and that investigation and treatment may improve these symptoms.

Different sleep symptom clusters may also have implications for cancer outcomes. Insomnia symptoms have been found to be associated with increased cardiovascular mortality in the general population, however, a recent meta-analysis concluded there was no association between insomnia symptoms and cancer mortality (Ge et al., 2019). Short self-reported sleep duration, as well as increased snoring after cancer diagnosis, has been shown to be associated with reduced cancer survival in post-menopausal women (Phipps et al., 2016). Similarly, sleepiness may also be a risk for poor outcomes in cancer, and a sleep duration of more than 9 hours is associated with increased risk of breast cancer mortality (Trudel-Fitzgerald et al., 2017). Severe OSA has been linked to more aggressive melanomas (Martinez-Garcia et al., 2018), however, it is unknown whether treatment of OSA will improve the outcome for these melanomas. Identification of different sleep symptom clusters in patients with cancer, and addressing underlying cause of these symptoms, may be important to improving cancer outcomes.

4.1 | Limitations

Unlike previous sleep symptom cluster studies, this study was performed in 318, mostly female, patients. Previous cluster studies have used OSA databases of 800–6500 sleep clinic patients (Bailly et al., 2016; Keenan et al., 2018; Saaresranta et al., 2016; Vavougios et al., 2016; Ye et al., 2014), although one study was in 198 subjects only (Lacedonia et al., 2016), and these were predominantly male. We also studied two cancers which are either exclusively female (endometrial cancer), or predominantly female (breast cancer), which may have influenced the outcome, although there were no differences in the number of males and females across the clusters. In addition, patients were studied at different times after their initial treatment, which may also have influenced the distribution within clusters, although this seems unlikely as patients with endometrial cancer, who were studied earlier after diagnosis, were evenly distributed across the symptom clusters, whereas breast cancer patients who were studied later were unevenly distributed. In addition, within each cancer cohort there are variations within each of the cancers in terms of molecular and histopathological subtypes in addition to the cancer stage and grade which may also interact with sleep symptoms, although we do not have sufficient numbers to determine this. Similar to previous studies, this study is cross-sectional, descriptive, and does not contain longitudinal or outcome data. This study only examined symptoms of sleep disorders, and there is no objective assessment or clinical diagnosis of sleep disorders. We also do not have the prevalence of OSA in these cancer cohorts, although it is likely that many of the subjects have at least moderate OSA based on our previous study (Madut et al., 2021), which includes a subgroup of the current study. In addition, it is possible that some of the symptoms may be due to other medical diagnoses associated with cancer, such as anaemia. Despite these limitations, we were able to describe four distinct sleep symptom clusters in cancer patients, a similar outcome to those described in much larger and predominantly male sleep clinic groups.

5 | CONCLUSION

In a group of 318 patients with a history of melanoma, breast and endometrial cancer we described four distinct sleep symptom clusters, which were differently distributed between the cancer groups. Identification of each of these sleep symptom clusters allows for a more personalised approach to the management of sleep in these cancer
patients, which may then impact the long term quality of life and survival outcomes on an individual patient basis.

AUTHOR CONTRIBUTION
KK, Study design, Data Collection, Data Analysis, manuscript preparation, manuscript review. ASM: Data Collection, Manuscript review. No conflicts to declare. HS: Data Collection, manuscript preparation, manuscript review, no conflicts to declare. MM: Data analysis, manuscript preparation, manuscript review. AB: Study design, manuscript review. EE: Study design, Manuscript review. JH. Study design, Manuscript review GJM: Study design, Manuscript review. TCA. Study design, Data analysis, Manuscript preparation, manuscript review. ADF: Study design, manuscript review.

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CONFICT OF INTEREST STATEMENT
No authors have any conflict of interest to declare.

DATA AVAILABILITY STATEMENT
The data in this study will be available in a freely accessible repository on acceptance of the publication.

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