



Citation for published version:
Košir, U, Loades, M, Wild, J, Wiedemann, M, Krajnc, A, Roškar, S & Bowes, L 2020, 'The impact of COVID-19 on the cancer care of adolescents and young adults and their well-being: Results from an online survey conducted in the early stages of the pandemic', Cancer. https://doi.org/10.1002/cncr.33098

DOI:

10.1002/cncr.33098

Publication date: 2020

Document Version Peer reviewed version

Link to publication

University of Bath

Copyright and moral rights for the publications made accessible in the public portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy
If you believe that this document breaches copyright please contact us providing details, and we will remove access to the work immediately and investigate your claim.

Download date: 22. Sep. 2020

The impact of COVID-19 on adolescents and young adults' cancer care and their wellbeing: Results from an online survey conducted in the early stages of the pandemic

Urška Košir¹, Maria Loades², Jennifer Wild¹, Milan Wiedemann¹, Alen Krajnc³, Sanja Roškar⁴, Lucy Bowes¹

*Corresponding Author: Urška Košir, Department of Experimental Psychology, University of Oxford, Anna Watts Building, Radcliffe Observatory Quarter, Woodstock Rd, Oxford, OX2 6HG, United Kingdom. Email: urska.kosir@psy.ox.ac.uk.

Acknowledgment: Authors would like to thank all participants who took the time to respond during the stressful times. Urška Košir would like to acknowledge the support of the Economic and Social Research Council (ESRC), Ad Futura (Slovenia), and the following patient organisations: EuropaDonna Slovenia, Lymphoma and Leukemia Society, OnkoMan, OnkoNet, and Elephants and Tea for help with recruitment and continued support of research endeavours. Dr Loades is funded by the National Institute for Health Research (NIHR Doctoral Research Fellowship, DRF-2016-09-021). Dr Wild's research is supported by MQ and the Oxford Health NIHR Biomedical Research Centre. This report is independent research. The views expressed in this publication are those of the authors(s) and not necessarily those of the NHS, NIHR or the Department of Health and Social Care.

¹ Department of Experimental Psychology, University of Oxford, Oxford, United Kingdom; ² Department of Psychology, University of Bath, Bath, United Kingdom;

³ Department of Chemistry, Chemistry Research Lab, University of Oxford, Oxford, United Kingdom;

⁴ Centre for Hearing and Speech Maribor, Maribor, Slovenia

Abstract

Background: Due to the global spread of COVID-19, oncology departments across the world

rapidly adapted their cancer care protocols, balancing the risk of delaying cancer treatments and

risk of COVID-19 exposure. COVID-19 and associated changes may have an impact on psycho-

social functioning of cancer patients and survivors. We wanted to understand the impact of

COVID-19 pandemic on young people living with and beyond cancer.

Method: In this cross-sectional study, we surveyed 177 individuals, aged 18 - 39 years, about

the impact of COVID-19 on their cancer care and psychological wellbeing. Participants also

reported their information needs relating to COVID-19. Responses were summarized using

content analysis approach.

Results: This is the first study to examine psychological functioning of young patients and

survivors during the first weeks of the COVID-19 pandemic. Close to a third of respondents

reported increased levels of psychological distress, and as many as 60% reported feeling more

anxious than they did prior to COVID-19. More than half also wanted more information tailored

to them as young cancer patients.

Conclusions: The COVID-19 pandemic is rapidly evolving and changing the landscape of

cancer care. Young people living with cancer are a unique population and might be more

vulnerable during this time compared to their healthy peers. There is a need to screen for

psychological distress and attend to young people whose cancer care has been delayed. As the

lockdown begins to ease, the guidelines about cancer care should be updated according to this

population's needs.

Keywords: COVID-19; survey; depression; anxiety; information needs; adolescent cancer;

young adult cancer

2

Introduction

COVID-19 was first identified in China, in December 2019¹ and is an infectious disease caused by the novel SARS-CoV-2 virus.² In most patients, COVID-19 infection results in mild symptoms such as a fever and a dry cough, but in some cases, and particularly in those with underlying health conditions and those who are immune compromised, COVID-19 can be more severe and can result in serious complications such as Acute Respiratory Distress Syndrome or even death.³ To date, findings suggest that COVID-19 poses little threat to children and young people,⁴ however, cancer patients and survivors may be a vulnerable population. Individuals undergoing active treatment or taking immunosuppressive medication are known to be at higher risk for viral infections,⁵ while those in survivorship might experience distress due to changes in their follow-up care. Most young people living beyond cancer worry about cancer recurrence^{6,7} and resultantly the disrupted follow-up care, which includes physical exams and screening may lead to heightened anxiety. As such, adolescent and young adult (AYA) cancer patients and survivors remain a group with unique needs during a pandemic outbreak like the current COVID-19, even though the relative risk of infection and mortality for this population have not yet been established.

Adolescents and Young Adult (AYA) cancer patients and survivors have been recognized as a unique group of patients who face different challenges and needs than their younger peers, or older adults.^{8,9} In response to the global spread of the novel SARS-CoV-2, oncology departments quickly acted and adapted measures and protocols for cancer care. Most centers sought to balance the risk of delaying cancer treatment with that of exposing a vulnerable cancer patient to the virus, which resulted in a crucial reprioritization; the most essential and necessary treatments remained on schedule, while early preventative screenings, follow-up appointments, and non-essential surgeries were often delayed.¹⁰ These changes were made rapidly and the newly devised guidelines were operationalized with increased survival in

mind, and in hopes to render the oncology centers 'pandemic proof.' While taking such measures is necessary for successful infection control, the subsequent steps will have to involve consideration of the psychological impact of COVID-19 and related changes on cancer patients, survivors and their families.

Even though the scientific, as well as patient support communities were quick to recognize the need to provide information about COVID-19 to cancer patients, ¹³ the sudden increase and a plethora of resources available was not necessarily tailored to AYA patients' and survivors' needs. We do not know if the information available is adequate, and if AYAs living with or beyond cancer feel confident about making sense of and applying it.

As after any major global disaster, the COVID-19 pandemic is also likely to result in increased mental health symptomatology. ^{14,15} Considering that a subset of AYA cancer patients experiences significant levels of psychological distress even under normal circumstances, ^{16,17} it is important to recognize that worries associated with the pandemic, and any delays in life-saving treatments are likely to contribute to the anxiety in this population. Furthermore, the social distancing measures imposed to contain the spread of the infection may have unintended consequences for mental health via changes in social support, economic consequences and changes to daily routines. ¹⁸ In AYAs who are already vulnerable to experiencing distress, this may further compound their anxiety. To ensure that the needs of AYAs continue to be met during and after the COVID-19 outbreak, developing new strategies that will help manage, and attenuate worries about the COVID-19 exposure should be a priority. This effectively requires an understanding of what the needs of AYAs are in this unprecedented context.

To adapt and better prepare for any future outbreaks, it is important to understand the impact of the novel SARS-CoV-2 on AYAs cancer care and their wellbeing. The aim of this study was 1) to gather initial evidence of the impact of COVID-19 on AYA cancer patients' and survivors' psychological well-being and cancer care, and 2) to understand where they

received the information about the pandemic and how satisfied they were with the resources on COVID-19.

In order to provide the future readership with context, **Figure 1** provides an overview with all the relevant dates and facts known to the authors at the time of writing this article (May 12, 2020).

[INSERT FIGURE 1: Global and UK COVID-19 context at the time of this study.]

Method

This anonymous cross-sectional survey was an extension of an ongoing longitudinal study exploring the predictors of psychological well-being and holistic recovery in AYA cancer patients. The original study protocol and COVID-19 survey material can be viewed at the following link: https://osf.io/ncpv8/. This study has been approved by the Medical Sciences Inter-Divisional Research Ethics Committee at the University of Oxford (R61437/RE005).

Data were collected using Qualtrics, a secure online survey software, via an anonymous link distributed on social media outlets and online via patient organizations' web posts or newsletters. Participants were provided brief information about the study, eligibility criteria, and the current aims; understanding what was the COVID-19 impact on their cancer care and wellbeing. The survey had two equal arms: a Slovenian and an English, accessible to anyone who was able to read and respond in English. Participants were all cancer patients or survivors aged between 18 and 39 years, diagnosed with any form of malignant cancer at age 10 or later.

Measures and survey description

Demographic and Medical information

Participants reported their age, gender, geographical region, and current living arrangement. They also reported their current health status, treatment status, whether or not

they had a pre-existing mental health condition or other chronic illness, or if they were taking any immunosuppressant medication (i.e. steroids).

Psychological distress

Psychological distress was measured using the Patient Health Questionnaire for Depression and Anxiety (PHQ-4), a 4-item measure of the frequency of two anxiety and two depressive symptoms on a 4-point Likert scale from 0 (not at all) to 3 (nearly every day). Reliability and validity of the brief PHQ-4 scale have been established in general and clinical populations, and a score of 3 or above on either sub-scale, and a total score of 6 or above is indicative of clinically relevant distress. 19,20 Cronbach's α for PHQ-4 in our study was 0.84, and 0.83 and 0.78 for anxiety and depression subscales, respectively.

COVID-19: impact on care and information resources

Respondents self-reported if and how the pandemic related events disrupted their cancer treatment and care, what the pandemic impact was on their psychological well-being, where they received the information about the novel SARS-CoV-2, and how satisfied they were with the information available on a scale from 0 (least satisfied/trustworthy) to 10 (most satisfied/trustworthy). (See https://osf.io/ncpv8/)

Analysis

The analyses were conducted in R (Version 3.6.3)²¹ using the following packages: ggplot2 (Version 3.3.0),²² dplyr (Version 0.8.5),²³ psych (Version 1.9.12.31)²⁴ and papaja (Version 0.1.0.984).²⁵

Survey data were analysed in an exploratory and descriptive manner in order to provide preliminary evidence of the impact of the COVID-19 pandemic on the subjective well-being

of cancer patients. We also reported self-reported COVID-19 impact on individuals' cancer treatment protocols.

Qualitative responses were analysed using content analysis approach.²⁶ The approach has been described as suitable for mixed method research, allowing for a more positivistic approach to analysis and phenomena description.²⁷ Two researchers (UK, SR) independently read and coded all of the responses to open-ended questions about COVID-19 impact on cancer patients and caretakers' psychological well-being. The researchers coded responses, both deductively by organizing them around the present study aims and survey questions, as well as inductively by devising a list of categories to describe impact on cancer care and the most common worries from participants' descriptions. The formulated categories were refined and operationalized into a coherent list upon which all qualitative responses were mapped. The inter-rater reliability for this procedure was 96%. Lastly, the classification of responses was checked by a third author, and discrepancies were resolved in a consensus meeting. All illustrative quotes in the main body are from different participants.

Results

Participants

During the period between April 6 and May 12, 2020, 177 AYA cancer patients and survivors participated in the study and completed at least 80% of the survey. Of those, 39 were from the Slovenian arm and 138 from the English arm. Participants' demographic and medical information is presented in **Table 1**. On average, cancer patients were 29.33 (SD = 6.17) years old. A third [57 out of 177] of responders were in active treatment at the time of the study, 14% [24 out of 177] completed treatment within the previous 6 months, and 54% [96 out of 177] completed their cancer treatment more than 6 months ago. Of the latter, 14 (15%) individuals were still taking immunosuppressive medication.

[INSERT TABLE 1: Demographic and Medical information of participants]

COVID-19 impact on cancer care

Forty-five percent [79 out of 177] of AYA cancer patients reported that COVID-19 had an impact on their cancer treatment or care, and 21% [38 out of 177] reported no impact at the time, but worried that there might be.

According to the short answers, the analyses revealed six categories of most commonly reported changes in cancer care. These categories were: follow-up appointments being postponed or cancelled, appointments carried out over the phone or virtually, cancer treatment and/or surgery being postponed, having to be alone during the treatment, reduced access to medicine, and changes in treatment protocols, such as reduced blood-work or oral instead of intravenous chemotherapy.

COVID-19 impact on psychological well-being

Over a third of AYA participant scored in the clinical range for psychological distress. Individuals undergoing treatment or within 6 months of treatment completion, on average, reported higher levels of psychological distress. Symptoms of anxiety were more common than depressive symptomatology. The mean scores and proportions of participants scoring in the clinical range of psychological distress are reported in **Table 2**.

Additionally, participants were asked how they felt compared to before the pandemic. Sixty-two percent [109 out of 177] of respondents reported feeling more anxious than before, 34% [52 out of 177] reported feeling the same, and 9% [16 out of 177] reported feeling less anxious than before. The proportion of individuals feeling more anxious was the same for those currently in treatment, as well as those who have completed treatment over 6 months ago (see Supplementary material).

[INSERT TABLE 2: Psychological distress among AYAs]

The short responses revealed that the most common concerns among the AYAs related to COVID-19 and their health status were not being sure how big of a risk COVID-19 posed for them, how their body would react to the infection, anxiety about experiencing serious complications if they were to catch the virus, and having their cancer care delayed. However, individuals also reported impact on mood, as well as worries about their family members, returning to "the normal" and finances. The categories and examples of participants' responses can be viewed in Table 3.

[INSERT TABLE 3: Categories of content on COVID-19 impact on psychological wellbeing]

Moreover, physical distancing measures imposed by many authorities resulted in social isolation with 52% [90 out of 173] of AYAs reported feeling more isolated than prior to the pandemic. On the other hand, 39% [68 out of 173] reported feeling about the same, and 9% [15 out of 173] reported feeling less isolated than before, and better understood by their friends and family.

"Even though I am still as isolated as I was when going through treatment and recovery, I find I feel less isolated than before because now my friends and family understand a lot better how it feels."

The qualitative responses highlighted that despite an increase in social interactions online, many individuals missed a more intimate human connection with others. While a few individuals reported having more time for themselves and their families, others reported that the isolation made them more anxious, depressed, and claustrophobic. Their needs as cancer patients were felt to have become secondary:

"People are so concerned with COVID-19 that they don't care about my cancer anymore."

For some, isolation also acted as a trigger, reminding them of their cancer:

"It makes me feel like I am in the first steps of my treatment all over again."

COVID-19 information resources and satisfaction

Participants were asked what sources they relied on for obtaining information related to the COVID-19 pandemic. The majority of respondents [159 out of 172] relied on a minimum of 2 different sources, with the general news being most read [135 out of 172], followed by social media [used by 109 out of 172]. Less than a third [46 out of 172] relied on cancer support groups and patient organizations for COVID-19 specific information. Individuals in different illness phases reported relying on the same information resources. Satisfaction and trustworthiness of the available information related to COVID-19 were perceived to be moderate, 6.2/10 and 6.1/10 respectively, with individuals currently undergoing treatment reporting least satisfaction and trustworthiness for the available information (see Supplementary material for breakdown by treatment status).

Only 24% [42 out of 173] of respondents reported receiving direct communication regarding COVID-19 from their oncologists or health care teams, majority (64%) of whom was undergoing treatment at the time of this study. Over half of the participants, 56% [96 out of 170] reported wanting more information about how to cope with the pandemic (see Supplementary material for breakdown by treatment status). Furthermore, 35% [60 out of 172]

of AYAs would have preferred information about COVID-19 tailored to their needs as young cancer patients and survivors.

"It would be nice to have something to read designed with me in mind." (Female, 32)

On the other hand, a few individuals reported that having more information would be overwhelming, and some reported the need for information specifically for mental health resources for cancer patients during a pandemic.

Discussion

Evidence to-date suggests that young people are at lower risk for contracting the novel SARS-CoV-2 virus or developing serious complications from it, 4,28 however, youth living with and beyond cancer might be at an increased risk because due to the compromised immunity resulting from illness and/or cancer late-effects. 5,29 Moreover, changes in cancer care protocols may increase the levels of anxiety and uncertainty not only for those undergoing active treatment, but also for individuals in the long-term care. Our cross-sectional study of youth living with and beyond cancer, conducted in the initial weeks of the COVID-19 pandemic, found that 3 in 10 AYAs experienced increased levels of psychological distress, with anxiety being more prominent than depressive symptomatology. As many as 6 in 10 AYAs reported feeling more anxious during the COVID-19 pandemic than they had been previously.

Higher levels of distress during a pandemic outbreak might be expected¹⁴ and a variety of psychological problems such as panic disorder, anxiety, and depression due to COVID-19 pandemic have already been observed in general population.³⁰ It may be that this is compounded for cancer patients, given the salience of risks to their health and their

vulnerability due to immunosuppression. The themes driving anxiety and worry for our participants were predominantly related to physical health and cancer-related concerns. A recent study of Italian adolescent cancer patients and survivors revealed that both groups worried about contracting the virus, and felt personally at risk for severe complications, with survivors reporting more worry than those in follow up care. When stratified by treatment status, our results revealed the highest levels of anxiety in those who have completed treatment within the past 6 months, followed by those in treatment at the time of the study. We speculate that this may in part be due to the uncertainty often experienced upon treatment completion, when individuals often have to wait for scans before remission can be confirmed. These treatment transitions can be burdensome for patients. The first few months after treatment are also often marked by frequent follow up appointments, all of which were likely postponed or cancelled due to the current pandemic. The added uncertainty is likely to contribute to anxiety symptoms.

The lowest levels of clinically elevated psychological distress were present in those off treatment, for whom the care was also least impacted by the pandemic and associated events. Nevertheless, a high proportion of those off treatment also reported feeling more anxious now than compared to the times prior the pandemic.

The social impact of the disease containment measures has been considerable. Most governments and authorities across the world have imposed physical distancing as a way to slow the spread of the virus within communities. For many, these measures resulted in social isolation, which can result in increased feelings of loneliness and further contribute to the feelings of low mood and anxiety.³³ The majority of our participants recounted being familiar with isolation measures due to their cancer experience. For some AYAs, de novo isolation acted as a negative reminder of their illness, while on the other hand, almost 10% reported feeling less anxious and isolated compared to the time prior the pandemic. The qualitative

responses revealed that this may be due to the fact that they felt better understood, spent more time with their families, or even found more time for themselves.

Research to date suggests that AYAs with cancer like to be involved in their care and need tailored, age-appropriate information about their cancer, care, late-effects, and resources.^{34,35} Only a quarter of our participants reported receiving direct communication related to COVID-19 from their health care providers and we demonstrated that more than half of AYAs would like to have received more information on how to cope during the COVID-19 pandemic, specifically, information tailored to them as young cancer patients and survivors. Regardless of the time passed since diagnosis, the most common worry was knowing the extent of the risk for contracting the virus and how their body would react if they were to contract the infection.

Unlike Italian adolescents,³¹ our AYA participants consumed most of the news related to the pandemic via general news sources, followed by social media outlets. This difference may be due to the older age of our sample. The fact that AYAs received most information from general news sources could also explain the reported need for more tailored information. The utility of social media as a way to deliver reliable information, as well as interventions to AYA cancer communities is gaining traction and could pave the way for an eHealth approach to oncological AYA care.³⁶ Preliminary evidence supports the feasibility and acceptability of online psychosocial interventions for AYAs in treatment³⁷ or early post-treatment phases,³⁸ however, future research should carefully evaluate if online support and digital approaches to care respond equally well to the needs of AYAs living with and beyond cancer.

Strengths and Limitations

While the global reach and early data collection add the largest contribution to the growing body of knowledge about the COVID-19 pandemic, there are several limitations that limit the generalizability of our work. Our sample size was relatively small, highly heterogeneous,

mostly female, and with most responses submitted from North America and Europe. While a geographically diverse sample offers broad and preliminary insights into COVID-19 related concerns of AYAs globally, it also limits generalizability due to the diverse health care systems, time-lags between peaks of the pandemic across different countries, and measures taken to prevent the spread of COVID-19.

Participants were a self-selected group who responded to the online study post, and we collected only minor, though relevant details relating to their medical and cancer history. Moreover, the survey was cross-sectional in nature and psychological distress was assessed using a very brief measure, and relied on short answer questions, to which many individuals provided responses with limited context and detail. We relied on retrospective self-report of changes in psychological distress since the advent of COVID-19, which are subject to reporting biases. Nevertheless, have provided important preliminary evidence, which offers invaluable directions for the future research on the impact of COVID-19, and also captured the experiences of this vulnerable population in the early weeks of the COVID-19 pandemic. Our sample included respondents at different stages of cancer treatment and beyond.

Clinical implications

The COVID-19 pandemic is rapidly evolving, and we learn new details every day. Even though the disruptions in our daily routines and health care systems pose many challenges, therein lie many opportunities for improvement of AYA cancer care in the future. The current situation highlights the continued need to screen for psychological distress and attend to those who might be experiencing distress even under normal circumstances. Finding ways of doing this whilst providing follow-up care remotely is particularly important within the current ways of working, and the brief PHQ-4 used in this study may offer a promising way to monitor psychological distress in clinical contexts.

The health care teams should actively engage and provide relevant and reliable information about COVID-19 for cancer patients and survivors, as well as information about any mental health resources. This will be important as the lockdown measures begin to ease in the coming months. Individuals whose care was delayed should receive adequate support so that they follow through with their appointments and continue the necessary treatment. While attending to those who are undergoing treatment or just recently completed it has been a priority, our results emphasize that the needs of long-term survivors should not be overlooked, as many may be experiencing higher levels of distress due to their illness history.

Telemedicine and eHealth might offer a favorable approach for some AYAs, however, in scaling them up, we have to ensure we do not leave behind those with limited access to digital technologies.

References:

- Lu H, Stratton CW, Tang YW. Outbreak of pneumonia of unknown etiology in Wuhan, China: The mystery and the miracle. *J Med Virol*. 2020;92(4):401-402. doi:10.1002/jmv.25678
- WHO. Clinical management of severe acute respiratory infection when Novel coronavirus (nCoV) infection is suspected: interim guidance.
 https://www.who.int/publications-detail/clinical-management-of-severe-acute-respiratory-infection-when-novel-coronavirus-(ncov)-infection-is-suspected. Published 2020. Accessed May 11, 2020.
- 3. Rothan HA, Byrareddy SN. The epidemiology and pathogenesis of coronavirus disease (COVID-19) outbreak. *J Autoimmun*. 2020;109(February):102433. doi:10.1016/j.jaut.2020.102433
- 4. Chen N, Zhou M, Dong X, et al. Epidemiological and clinical characteristics of 99

- cases of 2019 novel coronavirus pneumonia in Wuhan, China: a descriptive study. *Lancet*. 2020;395(10223):507-513. doi:10.1016/S0140-6736(20)30211-7
- 5. Kamboj M, Sepkowitz KA. Nosocomial infections in patients with cancer. *Lancet Oncol.* 2009;10(6):589-597. doi:10.1016/S1470-2045(09)70069-5
- 6. Koch L, Jansen L, Brenner H, Arndt V. Fear of recurrence and disease progression in long-term (≥5 years) cancer survivors A systematic review of quantitative studies.

 *Psychooncology. 2013;22(1):1-11. doi:10.1002/pon.3022
- 7. Kosir U, Wiedemann M, Wild J, Bowes L. Cognitive Mechanisms in Adolescent and Young Adult Cancer Patients and Survivors: Feasibility and Preliminary Insights from the Cognitions and Affect in Cancer Resiliency Study. *J Adolesc Young Adult Oncol*. 2019;00(00):1-6. doi:10.1089/jayao.2019.0127
- Marjerrison S, Barr RD. Unmet Survivorship Care Needs of Adolescent and Young Adult Cancer Survivors. *JAMA Netw Open*. 2018;1(2):e180350.
 doi:10.1001/jamanetworkopen.2018.0350
- 9. Barr RD, Ferrari A, Ries L, Whelan J, Bleyer WA. Cancer in adolescents and young adults: A narrative review of the current status and a view of the future. *JAMA Pediatr*. 2016;170(5):495-501. doi:10.1001/jamapediatrics.2015.4689
- Dinmohamed AG, Visser O, Verhoeven RHA, et al. Fewer cancer diagnoses during the COVID-19 epidemic in the Netherlands. *Lancet Oncol.* 2020. doi:10.1016/S1470-2045(20)30265-5
- 11. van de Haar J, Hoes LR, Coles CE, et al. Caring for patients with cancer in the COVID-19 era. *Nat Med.* 2020. doi:10.1038/s41591-020-0874-8
- 12. Schrag D, Hershman DL, Basch E. Oncology Practice During the COVID-19
 Pandemic. *JAMA*. 2020;published. doi:doi:10.1001/jama.2020.6236
- 13. Nekhlyudov L, Duijts S, Hudson S V, Jones JM, Keogh J, Love B. Addressing the

- needs of cancer survivors during the COVID-19 pandemic. *J Cancer Surviv*. 2020. doi:https://doi.org/10.1007/s11764-020-00884-w
- 14. Holmes EA, O'Connor RC, Perry VH, et al. Multidisciplinary research priorities for the COVID-19 pandemic: a call for action for mental health science. *The Lancet Psychiatry*. 2020;0366(20):1-14. doi:10.1016/S2215-0366(20)30168-1
- Galea S, Merchant RM, Lurie N. The mental health consequences of COVID-19 and physical distancing. *JAMA Intern Med.* 2020:E1-E2.
 doi:10.1001/jamainternmed.2020.1562
- 16. Barnett M, McDonnell G, DeRosa A, et al. Psychosocial outcomes and interventions among cancer survivors diagnosed during adolescence and young adulthood (AYA): a systematic review. *J Cancer Surviv*. 2016;10(5):814-831. doi:10.1007/s11764-016-0527-6
- 17. Kosir U, Wiedemann M, Wild J, Bowes L. Psychiatric disorders in adolescent cancer survivors: A systematic review of prevalence and predictors. *Cancer Rep*.
 2019;(December 2018):e1168. doi:10.1002/cnr2.1168
- 18. Brooks SK, Webster RK, Smith LE, et al. The psychological impact of quarantine and how to reduce it: rapid review of the evidence. *Lancet*. 2020;395:912-920. doi:10.1016/S0140-6736(20)30460-8
- 19. Löwe B, Wahl I, Rose M, et al. A 4-item measure of depression and anxiety:

 Validation and standardization of the Patient Health Questionnaire-4 (PHQ-4) in the general population. *J Affect Disord*. 2010;122(1-2):86-95.

 doi:10.1016/j.jad.2009.06.019
- Kroenke K, Spitzer RL, Williams JBW, Löwe B. An Ultra-Brief Screening Scale for Anxiety and Depression: The PHQ-4. *Psychosomatics*. 2009;50(6):613-621. doi:10.1016/s0033-3182(09)70864-3

- 21. Team RC. R: A language and environment for statistical computing. 2020. https://www.r-project.org/.
- 22. Wickham H. ggplot2: Elegant Graphics for Data Analysis. 2016.
- 23. Wickham H, Francois R, Henry L, Muller K. dplyr: A Grammar of Data Manipulation. 2020. http://cran.r-project.org/package=dplyr.
- 24. Revelle W. psych: Procedures for Psychological, Psychometric, and Personality Research. 2019. https://cran.r-project.org/package=psych.
- 25. Aust F, Barth M. papaja: Create APA Manuscripts with R Markdown. 2018.
- 26. Krippendorf K. Content Analysis: An Introduction to Its Methodology. Fourth Edi. Los Angeles: SAGE Publications Ltd; 2018.
- 27. Elo S, Kääriäinen M, Kanste O, Pölkki T, Utriainen K, Kyngäs H. Qualitative Content Analysis. *SAGE Open.* 2014;4(1):215824401452263. doi:10.1177/2158244014522633
- 28. Boulad F, Kamboj M, Bouvier N, Mauguen A, Kung AL. COVID-19 in children with cancer in New York City. *JAMA Oncol.* 2020;Published:3-4. doi:doi:10.1001/jamaoncol.2020.2028 Author
- 29. Desai A, Sachdeva S, Parekh T, Desai R. Covid-19 and cancer: Lessons from a pooled meta-analysis. *JCO Glob Oncol*. 2020;6:557-559. doi:10.1200/GO.20.00097
- 30. Qiu J, Shen B, Zhao M, Wang Z, Xie B, Xu Y. A nationwide survey of psychological distress among Chinese people in the COVID-19 epidemic: Implications and policy recommendations. *Gen Psychiatry*. 2020;33(2):19-21. doi:10.1136/gpsych-2020-100213
- 31. Casanova M, Bagliacca EP, Silva M, et al. How young patients with cancer perceive the Covid-19 (coronavirus) epidemic in Milan, Italy: is there room for other fears?

 *Pediatr Blood Cancer. 2020;(March):e28318. doi:10.1002/pbc.28318
- 32. Wilkins KL, D'Agostino N, Penney AM, Barr RD, Nathan PC. Supporting

- Adolescents and Young Adults with cancer through transitions: Position statement from the Canadian Task Force on Adolescents and Young Adults with cancer. *J Pediatr Hematol Oncol.* 2014;36(7):545-551. doi:10.1097/MPH.0000000000000103
- 33. Matthews T, Danese A, Caspi A, et al. Lonely young adults in modern Britain: Findings from an epidemiological cohort study. *Psychol Med.* 2019;49(2):268-277. doi:10.1017/S0033291718000788
- 34. Greenzang KA, Fasciano KM, Block SD, Mack JW. Early Information Needs of Adolescents and Young Adults About Late Effects of Cancer Treatment. *Cancer*. 2020;0:1-8. doi:10.1002/cncr.32932
- 35. Drew D, Kable A, van der Riet P. The adolescent's experience of cancer: An integrative literature review. *Collegian*. 2019;26(4):492-501. doi:10.1016/j.colegn.2019.01.002
- 36. Ramsey WA, Heidelberg RE, Gilbert AM, Heneghan MB, Badawy SM, Alberts NM. eHealth and mHealth interventions in pediatric cancer: A systematic review of interventions across the cancer continuum. *Psychooncology*. 2019;29(1):17-37. doi:10.1002/pon.5280
- 37. Chalmers JA, Sansom-Daly UM, Patterson P, McCowage G, Anazodo A. Psychosocial assessment using telehealth in adolescents and young adults with cancer: A partially randomized patient preference pilot study. *J Med Internet Res.* 2018;7(8):e168. doi:10.2196/resprot.8886
- 38. Sansom-Daly UM, Wakefield CE, Bryant RA, et al. Feasibility, acceptability, and safety of the Recapture Life videoconferencing intervention for adolescent and young adult cancer survivors. *Psychooncology*. 2019;28(2):284-292. doi:10.1002/pon.4938

Figure captions

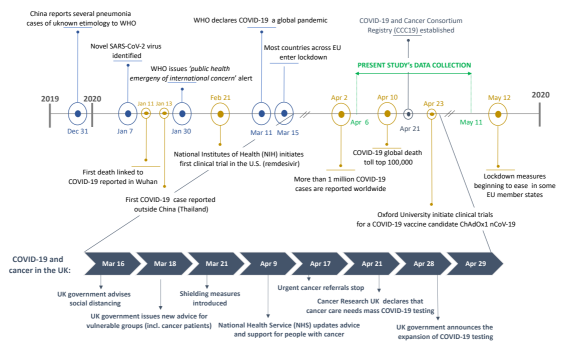


Figure 1: Global and UK COVID-19 context at the time of this study.

This figure aimed at providing future readership with COVID-19 context at the time of writing (May 12, 2020). By mid-March, most countries across Europe began nation-wide lockdowns. At the beginning of April, most cancer centers began adapting their treatment protocols. By late April COVID-19 a Cancer Consortium Registry was established with currently over 100 participating institutions globally reporting results about COVID-19 outcomes in cancer patients. There are currently several vaccine trials which have already began testing in healthy human participants.

Table 1
Demographic and Medical information of participants

Variable	n	%	M(SD)	Range
Age	177	100	29.33 (6.17)	[18 - 39]
Gender				
Female	154	87		
Male	20	11		
Prefer not to answer	3	2		
Region				
North America	93	53		
Central and South America	2	1		
Europe	78	44		
Middle East	1	1		

Africa	0	0		
Asia	1	1		
Oceania	3	2		
Living arrangement				
Alone	24	14		
Shared house	13	7		
Partner only	45	25		
Parents	20	11		
Family	72	41		
Other	4	2		
Treatment status				
Undergoing treatment	57	32		
Completed within 6 months	24	14		
Completed more than 6 months ago	96	54		
Immunosuppressant medications				
Yes	41	23		
No	122	69		
Prefer not to answer	14	8		
Table 1 continued				
Variable	n	%	M (SD)	Range
Mental health diagnosis				
Yes	66	37		
No	108	61		
Prefer not to answer	3	2		
Prefer not to answer	3	2		

Table 2
Psychological distress among AYAs

	Anxiety	Depression	Total	_
n	M (SD)	M (SD)	M (SD)	

Total sample	177	2.61 (1.94)	1.92 (1.71)	4.53 (3.29)
Undergoing treatment	57	2.84 (1.89)	2.07 (1.69)	4.91 (3.22)
Completed within 6 months	24	3.54 (2.02)	2.29 (1.90)	5.83 (3.60)
Completed more than 6 months ago	96	2.24 (1.87)	1.73 (1.66)	3.97 (3.15)
		Anxiety	Depression	Total
	n	n (%)	n (%)	n (%)
Total sample	n 177	n (%)	n (%) 34 (19%)	n (%) 51 (29%)
Total sample Undergoing treatment		. ,		
·	177	56 (32%)	34 (19%)	51 (29%)

Note. PHQ-4 items are rated on a 4-point Likert scale ranging from 0 to 3. The upper half of the table contains means and *SD*, while the bottom half contains number and proportion of individuals scoring in the clinical range.

de wo or Health anxiety Br	road statements and escriptions of anxiety and corries that are not specified redescribed in detail road statements about corries of own health,	"My anxiety levels tend to be quite high. I had felt like I had my anxiety under control, but this situation has thrown me off." "I worry that my current health problems will not be
wo	orries of own health,	
co	ypochondria, includes oncerns about individuals' tental health and illness iggers	taken as seriously in the healthcare system as they were before."
fee	Ientions of lacking support, beling low and sad due to olation, feelings of oneliness	"Whereas I was isolated prior to the global lockdown, I was still able to have the physical support of my family given that they were healthy. I have now lost that, making navigating cancer treatment all the more lonely."
	roader descriptions of low lood or morale, depression	"Increased feelings of depression and overeating."

Death/dying	Refers to the impact on mental health and wellbeing due to disruptions in daily living and activities Explicit mentions about dying and mortality	"My biggest source of happiness was my job and not being able to see my students is heartbreaking." "I am constantly worried that if I fall sick, I will end up dying considering my cancer history and weak immune system."
Worry about COVID-19 – self	Worries about how one's body would react to COVID-19, worries about severe complications due to COVID-19, including worries about receiving adequate care if they contracted COVID-19	"I am worried that if I caught it, I would be one of the ones that do not do well as I have had radiation to my spleen and still have low lymphocytes."
Worry about COVID-19 – others	Worries about close individuals or family members contracting the virus	"Mostly worried about family/elder members of the family wanting to get outside more."
Cancer care	Changes and disruptions in cancer care, which are unfavorable; worry about continuing with care and going to the hospital where many people are infected	"I worry about what stage my cancer will be when they are finally able to do my surgery/treatment."
Return to normal	Concerns about how to return to normal, what normal will look like; worries about returning back to work	"Going back to work eventually stresses me."
Finance	Direct financial impact from job loss or disruption; loss of insurance	"I have lost my job, risk losing my insurance, and will be in active treatment until late summer."

Table 3: Categories of content on COVID-19 impact on psychological wellbeing

SUPPLEMENTARY MATERIAL – Breakdown by treatment phase

	Better than before	Same as before	More anxious than
	N (%)	N (%)	before
			N (%)
Undergoing	7 (12%)	14 (25%)	36 (63%)
treatment $(n = 57)$			
Completed within 6	2 (8%)	9 (38%)	13 (54%)
months $(n = 24)$			

Completed more	7 (7%)	29 (30%)	60 (63%)
than 6 months ago			
(n = 96)			

Table 1: Table showing the proportion of individuals feeling more anxious or distressed now, compared to the time prior to the pandemic, stratified by treatment phase

	Received info about	Did not receive info about
	COVID	COVID
	N (%)	N (%)
Undergoing treatment (n =	27 (47%)	30 (53%)
57)		
Completed within 6	4 (17%)	19 (83%)
months $(n = 23)$		
Completed more than 6	11 (10%)	82 (90%)
months ago (n = 91)		

Table 2: Table showing the proportion of individuals who received direct communication about COVID-19 from the health care team by treatment phase

	Want more	Don't want more	Not sure
	information	information	N (%)
	N (%)	N (%)	
Undergoing	33 (59%)	13 (23%)	10 (18%)
treatment $(n = 56)$			
Completed within 6	13 (57%)	4 (17%)	6 (26%)
months $(n = 23)$			
Completed more	50 (55%)	16 (18%)	25 (27%)
than 6 months ago			
(n = 91)			

Table 3: Table showing the proportion of individuals who would have wanted more information about COVID-19, stratified by treatment phase

	COVID-19 Information sources	Satisfaction (mean)	Trustworthiness (mean)
Undergoing treatment (n = 57)	General news (n = 44) Medical Professionals (n = 43) Social Media (n = 37)	6/10	5.8/10
Completed within 6 months (n = 23)	General news (n = 22)	6.3/10	6.5/10

	Medical		
	Professionals (n =		
	14)		
	Social Media (n =		
	12)		
Completed more	General news (n =	6.3/10	6.2/10
than 6 months ago	69)		
(n = 91)	Social Media (n =		
	60)		
	Medical		
	Professionals (n =		
	44)		

Table 4: Table showing the most commonly used information resources for COVID-19, and reported satisfaction and perceived trustworthiness of these resources, stratified by treatment phase

NOTE: Satisfaction and trustworthiness are rated on a scale from 0 (least satisfied/trustworthy) to 10 (most satisfied/trustworthy).