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COVID-19 vaccination and breakthrough infections in patients with cancer

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Background: Vaccination is an important preventive health measure to protect against symptomatic and severe COVID-19. Impaired immunity secondary to an underlying malignancy or recent receipt of antineoplastic systemic therapies can result in less robust antibody titers following vaccination and possible risk of breakthrough infection. As clinical trials evaluating COVID-19 vaccines largely excluded patients with a history of cancer and those on active immunosuppression (including chemotherapy), limited evidence is available to inform the clinical efficacy of COVID-19 vaccination across the spectrum of patients with cancer.

Patients and methods: We describe the clinical features of patients with cancer who developed symptomatic COVID-19 following vaccination and compare weighted outcomes with those of contemporary unvaccinated patients, after adjustment for confounders, using data from the multi-institutional COVID-19 and Cancer Consortium (CCC19).

Results: Patients with cancer who develop COVID-19 following vaccination have substantial comorbidities and can present with severe and even lethal infection. Patients harboring hematologic malignancies are over-represented among vaccinated patients with cancer who develop symptomatic COVID-19.

Conclusions: Vaccination against COVID-19 remains an essential strategy in protecting vulnerable populations, including patients with cancer. Patients with cancer who develop breakthrough infection despite full vaccination, however, remain at risk of severe outcomes. A multilayered public health mitigation approach that includes vaccination of close contacts, boosters, social distancing, and mask-wearing should be continued for the foreseeable future.

Key words: COVID-19, vaccination, SARS-CoV-2, neoplasm, cancer

INTRODUCTION

The development of effective vaccines against COVID-19, the disease caused by SARS-CoV-2, has allowed widespread vaccination programs aimed at reducing symptomatic and severe COVID-19.1,2 The presence of underlying immunosuppression and receipt of recent systemic therapy for cancer have been associated with prolonged or severe infection and may reduce the efficacy of vaccination.3-8

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VACCINE AND BREAKTHROUGH INFECTIONS IN PATIENTS WITH CANCER

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INTRODUCTION

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Lower seroconversion rates following the receipt of COVID-19 vaccines have been observed in patients with underlying malignancy compared with non-cancer controls, with more concerning findings seen in patients with hematologic malignancies compared with those with solid cancers.\textsuperscript{7,9}

The clinical impact of these serological alterations is yet to be evaluated and the characteristics of breakthrough COVID-19 in vaccinated patients with cancer have not been reported, potentially due to low prevalence of SARS-CoV-2 during the time period of the published reports. Clinical trials evaluating COVID-19 vaccines report a high efficacy in the general population, with few-to-no severe breakthrough infections.\textsuperscript{1,2} As patients with a history of cancer and those on active immunosuppression (including chemotherapy) were largely excluded from these landmark studies, scant evidence informs the clinical efficacy of COVID-19 vaccination across the spectrum of patients with cancer. Given the predisposition of patients with cancer to suffer from severe outcomes in SARS-CoV-2 infection\textsuperscript{10} and considering their potential susceptibility to mount a less effective immune response following vaccination,\textsuperscript{11} better characterization of breakthrough COVID-19 following vaccination in this vulnerable population is needed.

Here, we sought to report the clinical attributes of patients with cancer who develop breakthrough SARS-CoV-2 infections and compare their weighted outcomes with those seen in a contemporary unvaccinated population, using data from the multi-institutional COVID-19 and Cancer Consortium (CCC19) registry.

**PATIENTS AND METHODS**

The large international CCC19 registry captures data on patients with a current or prior history of cancer who develop COVID-19 through a REDCap survey with methodology outlined previously.\textsuperscript{10,12} Deidentified data are collected using a comprehensive set of variables related to demographics, cancer status, anticancer therapies, SARS-CoV-2 infection, and COVID-19 vaccination. Data on COVID-19 vaccination were routinely collected on every newly entered case beginning with the first global approval in November 2020. Eligible cases included adult patients (>18 years of age) accrued from 1 November 2020 to 31 May 2021 with current or prior history of invasive cancer and laboratory-confirmed SARS-CoV-2 infection. Patients were excluded if vaccination status or timing was unknown or if the vaccine was administered after SARS-CoV-2 infection. We also excluded cases with poor data quality (score \textgreater;5 using the previously defined metric).\textsuperscript{13}

The primary endpoint was 30-day all-cause mortality among fully vaccinated patients compared with the unvaccinated population after Inverse Probability of Treatment Weighting (IPTW) to adjust for baseline clinical variables. Secondary endpoints included rates of intensive care unit (ICU) admission and/or mechanical ventilation (MV), and hospitalization rates in fully vaccinated, compared with unvaccinated patients after IPTW to adjust for baseline clinical variables.

Patients were categorized as fully vaccinated at the time of COVID-19 when two doses of vaccine had been administered and diagnosis of COVID-19 was recorded >4 weeks from first dose (BNT162b2),\textsuperscript{2} or when two doses of vaccine had been administered and a diagnosis of COVID-19 was recorded >6 weeks from the first dose (mRNA-1273),\textsuperscript{3} or in the event of a single dose of vaccination and a positive diagnosis >28 days post-vaccine (Ad.26.COV2.S).\textsuperscript{14} Patients who received at least one dose of vaccine and developed COVID-19 but did not meet the previous criteria were considered partially vaccinated. Unvaccinated patients were defined as having no known prior exposure to COVID-19 vaccination before COVID-19 diagnosis.

IPTW was used to adjust for differences in baseline clinical variables between fully vaccinated and unvaccinated patients [Supplementary Methods, available at https://doi.org/10.1016/j.annonc.2021.12.006] using age (as a continuous variable truncated at 90 years due to Health Insurance Portability and Accountability Act requirements), biologic sex (female; male), race (non-Hispanic White; Hispanic; non-Hispanic Black; other), smoking status (current or former smoker; never smoker), Eastern Cooperative Oncology Group performance status (ECOG PS 0; 1; \textgeq;2), baseline corticosteroid use [none; \textleq;10 mg/day prednisone dose equivalent (PDE); >10 mg/day PDE],\textsuperscript{15} lymphopenia [absolute lymphocyte count (ALC), \textleq;1000 versus >1000 per \textmu l], modified Charlson comorbidity index (mCCI, 0; 1; \textgeq;2; Supplementary Table S1, available at https://doi.org/10.1016/j.annonc.2021.12.006), cancer status (active and progressing versus not active and progressing), cancer type (solid organ tumor; hematologic neoplasm; both), and recent systemic anticancer therapy (any of cytotoxic chemotherapy, immunotherapy, targeted therapy, or endocrine therapy in the 3 months before COVID-19 diagnosis, or not).

The data dictionary used in the analysis, along with instructions on how to access full CCC19 data dictionary and code to generate derived variables is found in Supplementary Table S2, available at https://doi.org/10.1016/j.annonc.2021.12.006. All data analyses were carried out using R 4.0.3 and the R packages Hmisc 4.4-2, Matchit 4.2.0, ipw 1.0-11, survey 4.0, sandwich 3.0-1, and glmnet 4.1-1.

This study was exempt from Institutional Review Board (IRB) review (VUMC IRB#200467), was approved by IRBs at participating sites per respective institutional policy, and is registered on ClinicalTrials.gov (NCT04354701).

**RESULTS**

Overall, we identified 1787 patients with cancer and COVID-19 who met inclusion criteria, of whom 1656 (97%) were unvaccinated, 77 (4%) partially vaccinated, and 54 (3%) fully vaccinated [Supplementary Table S3, available at https://doi.org/10.1016/j.annonc.2021.12.006]. Baseline clinical factors by vaccination status are shown in Table 1.

Median age of fully vaccinated patients was 65.5 years [interquartile range (IQR) 57.0-72.8 years], 35 (65%) were
female and 38 (70%) were non-Hispanic White. A total of 19 (35%) had hematologic malignancies and 17 (31%) had an mCCI of ≥2. Before IPTW was carried out, more patients in the fully vaccinated cohort relative to the unvaccinated cohort had an underlying hematologic malignancy (35% versus 20%), were female (65% versus 55%), non-Hispanic White (70% versus 60%), received baseline prednisone or equivalent >10 mg/day (17% versus 7%), had ALC <1000/µl (46% versus 28%), or received systemic therapy within the prior 3 months (56% versus 43%).

Among the 54 fully vaccinated patients who developed COVID-19, 35 (65%) were hospitalized, 10 (19%) were admitted to ICU or required MV, and 7 (13%) died within 30 days. Comparable rates were observed in the unvaccinated group (Table 1).

| Table 1. Baseline clinical factors by baseline vaccination status |
|---------------------------------------------------------------|
|                  | Fully vaccinated | Partially vaccinated | Unvaccinated |
|                  | N (%)           | N (%)               | N (%)        |
| Total patients (N = 1787) | 54 (3)          | 77 (4)              | 1656 (93)    |
| Median age, years (IQR) | 65.5 (57.0-72.8) | 68.0 (58.0-78.0)    | 64.0 (54.0-74.0) |
| Female sex | 35 (65)        | 38 (49)             | 903 (55)     |
| Non-Hispanic White | 38 (70)        | 58 (75)             | 999 (60)     |
| ECOG performance status ≥2 | 9 (17)        | 14 (18)             | 224 (14)     |
| Modified Charlson comorbidity index ≥2 | 17 (31)      | 26 (34)             | 413 (25)     |
| Current or former smoker | 26 (48)      | 33 (43)             | 732 (44)     |
| Hematologic malignancy | 19 (35)       | 18 (23)             | 339 (20)     |
| Cancer status active and progressing | 10 (19)      | 17 (22)             | 237 (14)     |
| Systemic treatment within 3 months | 30 (56)      | 31 (40)             | 720 (43)     |
| Baseline prednisone dose-equivalent >10 mg/day | 9 (17)      | <5 (<6)             | 117 (7)      |
| ALC <1000/µl (not drawn/not available N = 902) | 25 (46)     | 26 (34)             | 469 (28)     |

COVID-19 infection severity

- 30-day mortality | 7 (13) | <5 (<6) | 160 (10)
- Admitted to intensive care | 10 (19) | 11 (14) | 215 (13)
- Hospitalized | 35 (65) | 45 (58) | 834 (50)

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We found no significant differences in ICU/MV or hospitalization rates between the vaccinated patients compared with the unvaccinated cohort after adjustment (AOR 1.13, 95% CI: 0.54-2.37 and AOR 1.25, 95% CI: 0.68-2.30, respectively). In both vaccinated and unvaccinated patients, higher ICU/MV and hospitalization rates were identified in patients with lymphopenia (AOR 1.43, 95% CI: 1.03-1.97 and AOR 1.62, 95% CI: 1.20-2.20, respectively), the presence of comorbid conditions (mCCI of ≥2 versus 0: AOR 2.67, 95% CI: 1.83-3.90, and AOR 2.42, 95% CI: 1.71-3.43, respectively; mCCI of 1 versus 0: AOR 1.83, 95% CI: 1.24-2.71, and AOR 1.65, 95% CI: 1.27-2.15, respectively), poor ECOG PS (ECOG PS ≥2 versus 0: AOR 2.19, 95% CI: 1.32-3.64, and AOR 3.68, 95% CI: 2.46-5.53, respectively; ECOG PS 1 versus 0: AOR 1.62, 95% CI: 1.05-2.48, and AOR 1.35, 95% CI: 1.02-1.78, respectively) and hematologic as opposed to solid cancers (AOR 2.00, 95% CI: 1.41-2.83, and AOR 2.42, 95% CI: 1.82-3.22, respectively).

Secondary outcome regressions and sensitivity analyses are described in Supplementary Methods and Supplementary Tables S4-S8, available at https://doi.org/10.1016/j.annonc.2021.12.006.

DISCUSSION

To our knowledge, this is the first study to evaluate the clinical characteristics and outcomes of patients with cancer who experience breakthrough infection following COVID-19 vaccination. Vaccination has been widely effective at reducing the severity of SARS-CoV-2 infection, but the protection afforded by this preventive modality can be wane over time. Based on the accumulation of knowledge
to date, it appears that patients with hematologic malignancy are less likely to mount an effective immune response.\textsuperscript{16} The enrichment of hematologic malignancies (35\% versus 20\%) in this cohort is consistent with evidence that these patients may have a blunted serologic response to vaccination secondary to disease or therapy,\textsuperscript{17,18} compounding their probability to develop more severe COVID-19 outcomes relative to patients with other types of tumors.\textsuperscript{19}

The described vaccinated cohort may be subject to ascertainment bias as a result of selective reporting, if reporting sites did not attain complete coverage of all eligible cases. Furthermore, it is possible that the vaccination exposure was not captured in some of the ostensibly unvaccinated patients. Neither post-vaccination titers nor T-cell-mediated immunity metrics are routinely checked in clinical care and were not captured in this study, leaving open the possibility that at least some of the patients manifesting COVID-19 did not have adequate immunity. Nevertheless, any infection in a fully vaccinated person meets the Centers for Disease Control and Prevention (CDC) definition of breakthrough infection (\url{https://www.cdc.gov/coronavirus/2019-ncov/vaccines/effectiveness/why-measure-effectiveness/breakthrough-cases.html}). Notably, boosters were not yet available during the timeframe of this study, and are not part of the definitions of fully vaccinated used in the original trials, which we adopted.\textsuperscript{1,2,14} The mortality and ICU/MV rates of 13\% and 19\%, respectively, recorded in the fully vaccinated group, with no significant differences compared with the unvaccinated group, represent a considerable residual risk. Additionally, this contrasts with a substantially lower rate of severe outcomes previously

### Table 2. Results of regression analysis following truncated inverse probability of treatment weighting

|                     | 30-Day mortality AOR (95% CI) | Intensive care unit/mechanical ventilation AOR (95% CI) | Hospitalization AOR (95% CI) |
|---------------------|-------------------------------|------------------------------------------------------|-----------------------------|
| Vaccination status  |                               |                                                      |                             |
| (ref = unvaccinated)|                               |                                                      |                             |
| Fully vaccinated     | 1.08 (0.41-2.82)              | 1.13 (0.54-2.37)                                     | 1.25 (0.68-2.30)            |
| Age (per 10 years increase) | 1.25 (1.08-1.44)          | 1.07 (0.96-1.20)                                     | 1.28 (1.18-1.39)            |
| Sex (ref = female)  |                               |                                                      |                             |
| Male                | 1.32 (0.92-1.90)              | 1.44 (1.06-1.95)                                     | 1.22 (0.98-1.52)            |
| Cancer status active and progressing (ref = not active and progressing) | 6.07 (4.00-9.19)          | 1.96 (1.30-2.96)                                     | 2.42 (1.71-3.43)            |
| Modified Charlson comorbidity index (ref = 0) |                               |                                                      |                             |
| 1                   | 1.66 (1.07-2.59)              | 1.83 (1.24-2.71)                                     | 1.65 (1.27-2.15)            |
| 2                   | 2.10 (1.36-3.24)              | 2.67 (1.83-3.90)                                     | 2.42 (1.71-3.43)            |
| ECOG performance status (ref = 0) |                               |                                                      |                             |
| 1                   | 2.26 (1.25-4.06)              | 1.62 (1.05-2.48)                                     | 1.35 (1.02-1.78)            |
| 2                   | 4.34 (2.35-8.02)              | 2.19 (1.32-3.64)                                     | 3.68 (2.46-5.53)            |
| On baseline corticosteroids >10 mg PDE/day (ref = none) |                               |                                                      |                             |
| Yes                 | 1.28 (0.69-2.39)              | 1.37 (0.81-2.31)                                     | 1.81 (1.13-2.88)            |
| Lymphopenia <1000/\mu l (ref = ≥1000/\mu l) |                               |                                                      |                             |
| Yes                 | 1.68 (1.11-2.55)              | 1.43 (1.03-1.97)                                     | 1.62 (1.20-2.20)            |
| Cancer type (ref = solid) |                               |                                                      |                             |
| Hematologic         | 1.20 (0.75-1.91)              | 2.00 (1.41-2.83)                                     | 2.42 (1.82-3.22)            |

AOR, multivariable adjusted odds ratio; CI, confidence interval; PDE, prednisone dose-equivalent; ref, reference.

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Figure 1. Forest plot showing results of regression analysis following truncated Inverse Probability of Treatment Weighting by clinical outcomes.

CI, confidence interval; ECOG, Eastern Cooperative Oncology Group; OR, odds ratio; PDE, prednisone dose-equivalent.
identified in vaccinated healthy individuals as opposed to matched unvaccinated controls, outlining the potential vulnerability of patients with cancer.

Lymphopenia, which has a strong association with severe SARS-CoV-2 infection, was present in 46% of fully vaccinated patients and 28% in the unvaccinated patients. This finding supports the notion that lymphopenic patients with cancer are at high risk for severe disease, even after vaccination. It has been previously shown that lymphocyte-depleted patients such as those receiving anti-CD20 monoclonal antibodies or chimeric antigen receptor T-cell (CAR-T-cell) treatment have a much weaker serological response to COVID-19 vaccines. Our results appear to confirm the clinical relevance of such previous observations.

Alternatively, more severe infection with resultant lymphopenia may have been present at diagnosis in vaccinated patients, potentially due to a lower suspicion of SARS-CoV-2 infection following full vaccination, resulting in a delayed presentation of vaccinated patients with cancer after COVID-19 symptom onset.

Some differences are apparent in the baseline characteristics of the fully vaccinated cohort relative to the unvaccinated population and can partially explain differences in clinical outcomes, even after adjustment. A higher prevalence of active and progressive cancer (19% versus 14%) and of recent receipt of systemic anticancer therapy (56% versus 43%) was seen in the fully vaccinated population relative to unvaccinated patients, with both being previously identified risk factors for adverse COVID-19 outcomes. The number of affected patients in this study is too small to make any definitive conclusions about specific types of anticancer therapy that might be associated with breakthrough infection.

Chronic corticosteroid use (>10 mg/day PDE), which is associated with higher odds of hospitalization from COVID-19 viral respiratory illness in a cohort of patients with autoimmune conditions, also appeared to be more prevalent at baseline in the fully vaccinated cohort compared with the unvaccinated patients (24% versus 14%).

Following regression analysis, the association between advancing age, active and progressing cancer, ECOG PS ≥2, mCCI ≥2, and lymphopenia with mortality suggests that these established prognostic factors for COVID-19 outcomes remain relevant in defining those patients who may still be at risk of severe outcomes following completion of vaccination.

Limitations of this study include the dependence on clinically annotated data and the reliance on time intervals (instead of exact dates) to capture vaccine administration relative to COVID-19 diagnosis. Given the timeframe of this analysis, it is unlikely that any patients had infection >6 months out from vaccination, such that waning immunity is not likely a major factor; future studies will need to consider this as well as receipt of boosters. The true population prevalence of vaccinated individuals during the time period is unknown and likely changed substantially, as at-risk populations were variably prioritized before vaccines became more generally available in spring 2021. As such, the rate of serious illness or mortality from COVID-19 among the total population of vaccinated patients with cancer is unknown, as vaccination was widely administered and likely protected the majority from serious illness. Strengths include the high-quality data with a robust quality assurance process, along with a comprehensive list of clinical, demographic, and laboratory variables.

With the emergence of the B.1.617.2 (delta) variant, which displays a higher transmissibility than previous strains of the virus, and for which available vaccines appear to show decreased neutralization and efficacy with a higher rate of breakthrough cases, the current findings underscore the need to maintain the implementation of public health measures required to control infection spread and protect vulnerable populations. Delta and subsequent variants will continue to raise the possibility of immune escape from the first generation of vaccines. Additional research is needed to further categorize the patients who remain at risk of symptomatic COVID-19 following vaccination, and test strategies that may reduce this risk. Based on experience in patients with prior organ transplantation, the strategy of administering a third primary series vaccination dose to increase antibody titers is one option recently suggested by the CDC to be considered for immunosuppressed patients, including patients with cancer on systemic anticancer therapies (https://www.cdc.gov/vaccines/covid-19/clinical-considerations/covid-19-vaccines-us.html). Further correlation of antibody responses to clinical benefit will be needed to validate the use of these strategies, as well as boosters and emerging strategies such as ‘mix and match’, where a different booster vaccine is administered from the primary vaccination course.

Overall, vaccination remains an invaluable strategy in protecting vulnerable populations, including patients with cancer, against COVID-19. Patients with cancer who develop breakthrough infection despite full vaccination, however, remain at risk of severe outcomes and should not be neglected. A mitigation approach that includes vaccination of close contacts, boosters, social distancing, and mask-wearing in public should be continued for the foreseeable future.

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DISCLOSURE

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