The Role of Stage at Diagnosis in Colorectal Cancer Black–White Survival Disparities: A Counterfactual Causal Inference Approach

Linda Valeri1,2, Jarvis T. Chen3, Xabier Garcia-Albeniz4, Nancy Krieger3, Tyler J. VanderWeele4,5, and Brent A. Coull5

Abstract

**Background:** To date, a counterfactual framework has not been used to study determinants of social inequalities in cancer. Considering the case of colorectal cancer, for which racial/ethnic differences in stage at diagnosis and survival are well documented, we quantify the extent to which black versus white survival disparities would be reduced if disparities in stage at diagnosis had been eliminated in a large patient population.

**Methods:** We obtained data on colorectal cancer patients (diagnosed between 1992 and 2005 and followed until 2010) from US-SEER (Surveillance, Epidemiology, and End Results) cancer registries. We employed a counterfactual approach to estimate the mean survival time up to the 60th month since diagnosis for black colorectal cancer patients had black–white disparities in stage at diagnosis been eliminated.

**Results:** Black patients survive approximately 4.0 [confidence interval (CI), 4.6–3.2] months less than white patients within five years since diagnosis. Had disparities in stage at diagnosis been eliminated, survival disparities decrease to 2.6 (CI, 3.4–1.7) months, an approximately 35% reduction. For patients diagnosed after the age of 65 years, disparities would be halved, while reduction of approximately 30% is estimated for younger patients. Survival disparities would be reduced by approximately 44% for women and approximately 26% for men.

**Conclusions:** Employing a counterfactual approach and allowing for heterogeneities in black–white disparities across patients’ characteristics, we give robust evidence that elimination of disparities in stage at diagnosis contributes to a substantial reduction in survival disparities in colorectal cancer.

**Impact:** We provide the first evidence in the SEER population that elimination of inequities in stage at diagnosis might lead to larger reductions in survival disparities among elderly and women.

Introduction

Use of a counterfactual causal inference framework is recognized as a valuable contribution to quantifying the causal effects of potential interventions (1). To our knowledge, however, this framework has not been applied to analysis of the contribution of stage at diagnosis to social inequalities in cancer outcomes. Here, we adopt a counterfactual causal inference framework to investigate determinants of black–white disparities as proposed by VanderWeele and Robinson (2), framed by an understanding that inequitable race relations, not “race” per se, are the cause of racial/ethnic health inequities, that is, unjust, unfair, and preventable social inequalities in health (3–8).

The case we have chosen is colorectal cancer, as black–white disparities in both stage at diagnosis and survival are well documented. Differences in colorectal cancer mortality rates between black and white patients started manifesting in 1980 (9) and the black–white mortality ratio peaked for women in 2005 at 1.5 and for men in 2006 at 1.6 (10). As colorectal cancer is the third leading cause of cancer-related death in both men and women (11), to reduce the uneven burden of cancer in the United States it is crucial to investigate rigorously why these disparities exist.

The determinants of colorectal cancer mortality disparities are multifactorial, with black–white disparities often attributed to differences in socioeconomic and behavioral factors, as well as in quality and access to prevention practices and to care. Since the 1970s, blacks have lagged behind whites in access to both screening and improved treatment options (12). Therefore, many have suggested that the large differences in stage at diagnosis and poorer prognosis for distant-stage disease may be driving mortality disparities (12).

Mixed evidence is found on black–white disparities in tumor characteristics (13, 14) and differences in tumor location are not considered to be main drivers of mortality disparities (14).

Previous studies reported that differences in stage at diagnosis accounted for up to 60% of the observed mortality disparities (15–17). These studies, however, were conducted in smaller patient populations and using regression approaches valid only under strong assumptions, which are violated in this context. In particular, a popular approach to estimate the extent to which
racial/ethnic disparities in survival is explained by intermediate factors is to compare estimates of the effect of interest from a regression model with and without adjustment for the hypothesized factor. As is now well documented in the epidemiologic and biostatistics literature, this approach is biased if there are interactions between race/ethnicity and stage at diagnosis (18).

In this article, we apply a counterfactual framework, using data from the National Cancer Institute’s (NCI) Surveillance, Epidemiology, and End Results (SEER) Program, to estimate the extent to which black–white survival disparities among colorectal cancer patients would be reduced had disparities in the intermediate factor, stage at diagnosis, been eliminated. We estimate a clinically meaningful, model-free measure of survival to quantify black–white disparities and the impact of this hypothetical intervention across baseline patients’ characteristics, providing important insights to patients, clinicians, and policy makers. In particular, we allow for interactions in race and stage in their impact on survival and, recognizing that people embody diverse forms of inequality simultaneously (3–4), we investigate differences in survival disparities according to age and gender.

Materials and Methods

Study population

We obtained data from NCI’s SEER Program, which provides information on tumor site, stage, and individual demographic information for cancers occurring in approximately 28% of the U.S. population. Patients from the Atlanta, Connecticut, Detroit, Iowa, New Mexico, San Francisco-Oakland, Seattle-Puget Sound, Utah, San Jose-Monterey, Los Angeles, and Rural Georgia registries were included in the study. The California registries were combined and the Rural Georgia and Atlanta registries were combined. The study population consisted of non-Hispanic white and non-Hispanic black cancer patients diagnosed between 1992 and 2005 and followed until 2010. We selected patients diagnosed above the age of 18, with microscopically confirmed adenocarcinoma histology and colorectal cancer as first diagnosis. We included patients diagnosed at stage I–IV according to the American Joint Committee on Cancer (AJCC) staging criteria (19). Patients from the Hawaii registry were excluded from the analysis due to the low percentage of non-Hispanic blacks. Finally, we excluded patients for whom the reporting source was nursing home/hospice, autopsy, or death certificate. Supplementary Fig. S1 and Supplementary Table S1 display the flowchart of eligibility criteria for the present study and the codes we used to extract information about tumor characteristics from SEER data. Census and American Community Survey (ACS) data were merged to SEER data to obtain area-based measures of socioeconomic position (20–21).

Stage at diagnosis was categorized as stage I–IV, according to the AJCC staging criteria. For the purpose of the present analyses, patients with in situ cancers and unstaged cancers were excluded from the primary analyses. Approximately 6% of the sample had unstaged cancer at diagnosis (n = 9,192).

Survival time in months from date of diagnosis was calculated under the assumption that, if last known vital status was alive, then the patients were presumed alive and the database study cut-off date was used as date of last contact. Survival time was censored for 27% of black and 30% of white population. Individuals who died within 5 years since diagnosis had survival time information reliably recorded, as registries conduct active patients’ follow-up for at least 5 years.

Age at diagnosis, categorized as age<50, 50–65, and >65, year at diagnosis (1992–1995, 1996–2000, 2001–2005), gender (men versus women), grade of tumor differentiation (I–IV), and registry were considered as potential confounders of the stage–survival association. Because individual-level SEP data are not available in the SEER registry, median household income at county level, an aggregate measure of SEP, was linearly interpolated from 1990 and 2000 Census, and 2009 American Community Survey data and categorized in quintiles.

Statistical analysis

The distribution of baseline characteristics between the two groups was assessed for differences using χ2 statistics. For non-Hispanic whites and non-Hispanic blacks within each confounder stratum (denoted by x), we estimated nonparametrically the restricted mean survival time (RMST) up to 60 months of follow up. This quantity is estimated as the area under the Kaplan–Meier curve from diagnosis to 60 months (22). We denote the difference of RMST for the blacks versus whites as the disparity in survival.

\[
\text{Disparity}(x) = \text{RMST (black, } x) – \text{RMST (white, } x).
\]  

( Eq.1)

The overall disparity measure was then obtained via random effect meta-analysis (23–24) across all confounder strata (x) to provide more precise inferences by borrowing strength while allowing for heterogeneous underlying disparity measures in each stratum.

We estimated the residual disparity between black and white individuals that would be observed had we intervened on stage at diagnosis. The causal contrast can be formally defined as the difference of the RMST in black versus white patients had we intervened to shift the stage at diagnosis distribution among blacks to match that of the whites, the most advantaged patients (2). Effectively, the residual disparity is calculated as the difference in the RMST for blacks versus whites once the stage-specific RMSTs for black patients are multiplied by the probability of being diagnosed at that stage in the reference population (white patients), adjusting for confounders of the stage–survival relationship. The quantity is estimated using the following standardization formula:

\[
\text{Residual Disparity}(x) = \sum_{\text{stage}} \{\text{RMST (black, stage, } x) \} – \text{RMST (white, stage, } x)\} \cdot p(\text{stage | white, } x).
\]  

( Eq.2)

To compute the residual disparity after the hypothetical intervention on stage, we estimated nonparametrically the area under the Kaplan–Meier curve, truncated at month 60, for white and black individuals in each stage and confounder (x) strata; the probability of being diagnosed at a certain stage in the white population was also empirically estimated. Equation 2 correctly estimates the effect of a shift in the distribution of stage at diagnosis in the black population towards that of the white population under the assumption that all confounders of the stage–survival relationship are adjusted for in the analysis.

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(2). The overall residual disparity measure was again obtained via random effect meta-analysis across all covariate strata.

Finally, percent disparity reduction was estimated as:

\[
\text{Disparity Reduction} = \frac{\text{disparity} - \text{residual disparity}}{\text{disparity}}
\]  
(Eq. 3)

Data were analyzed using the R software package (25).

Sensitivity analyses

We explored the robustness of our results to selection bias introduced by missing data on stage at diagnosis. Information on stage at diagnosis was missing for approximately 6% of the sample. Although the number of unstaged patients was not large, we observed an important difference in the 5-year survival probability between individuals with complete information and individuals with missing this information (50% vs. 35%). The survival probability and Kaplan–Meier curves for unstaged patients in the present study lied between those calculated for stage III and stage IV patients. Therefore, we assessed the sensitivity of our results to the extreme assumption that all unstaged individuals had stage IV at diagnosis.

Results

Our analytic sample consists of 166,722 colorectal cancer patients (148,010 non-Hispanic whites and 18,712 non-Hispanic blacks) with no missing age at diagnosis. As shown in Table 1, black and white individuals display significant differences in many of the baseline characteristics. White individuals display a survival advantage (log-rank test \(P < 0.0001\)) and are more likely than black patients to survive past 5 years from diagnosis (observed 5-year survival from death due to all causes: 49% vs. 40%, \(P < 0.001\)). Blacks are more likely to be diagnosed at stage IV than whites (24% vs. 17%; \(P < 0.001\)).

In the SEER population, we find that RMST by stage at diagnosis for whites differs substantially from that estimated for black patients. For white patients diagnosed at stage I, the area under the Kaplan–Meier curve is 51.7 months: that is, one expects a white colorectal cancer patient diagnosed at stage I to be alive for 51.7 months of the 60 months of follow-up. In contrast, the RMST for blacks is 48.5 months. For patients diagnosed at stage II, we observe RMST of 47.9 for whites versus 45.5 months for blacks; for stage III patients, 42.6 months for whites vs. 41.8 months for blacks; and for stage IV patients, 18.3 months for whites vs. 15.5 months for blacks. In this population we find the survival differences to be larger among patients diagnosed at stage I (Table 2). Race–stage interaction was significant in an accelerated failure time model, indicating that disparities were significantly larger for patients diagnosed at stage I (LRT \(P < 0.001\), Supplementary Table S2). Figure 1 summarizes the estimates of the observed RMST for black and white patients, and the counterfactual RMST for black patients under the scenario of a shift of stage at diagnosis.

Table 1. Baseline characteristics of colorectal cancer patients \(n (\%)\): US SEER registries, 1992–2005

<table>
<thead>
<tr>
<th>Stage</th>
<th>Black (n = 18,712)</th>
<th>White (n = 148,010)</th>
<th>(P)</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>3,131 (17%)</td>
<td>31,998 (22%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>II</td>
<td>5,079 (27%)</td>
<td>46,195 (31%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>III</td>
<td>4,724 (25%)</td>
<td>36,631 (25%)</td>
<td>0.1242</td>
</tr>
<tr>
<td>IV</td>
<td>4,442 (24%)</td>
<td>25,298 (17%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Unstaged</td>
<td>1,296 (7%)</td>
<td>7,888 (5%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>5-year survival</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>65</td>
<td>7,659 (40%)</td>
<td>72,076 (49%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Grade</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>I</td>
<td>1,393 (7.4%)</td>
<td>11,555 (8%)</td>
<td>0.0834</td>
</tr>
<tr>
<td>II</td>
<td>12,403 (66%)</td>
<td>91,578 (63%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>III</td>
<td>2,852 (15.2%)</td>
<td>29,163 (20%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>IV</td>
<td>84 (0.4%)</td>
<td>1,078 (0.1%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Unknown</td>
<td>1,006 (10.4%)</td>
<td>12,365 (8.3%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Age at diagnosis</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;50</td>
<td>2,396 (13%)</td>
<td>10,623 (7%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>50–65</td>
<td>5,921 (32%)</td>
<td>33,246 (23%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>&gt;65</td>
<td>10,395 (55%)</td>
<td>104,141 (70%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Male</td>
<td>8,962 (48%)</td>
<td>7,333 (39%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Date at diagnosis</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1992–1995</td>
<td>4,838 (26%)</td>
<td>42,783 (29%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>1996–2000</td>
<td>6,541 (35%)</td>
<td>54,606 (37%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>2001–2005</td>
<td>7,333 (39%)</td>
<td>50,621 (34%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Registry</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>San Francisco-Oakland, Los Angeles, San Jose-Monterey</td>
<td>7,300 (39%)</td>
<td>45,574 (31%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Connecticut</td>
<td>1,371 (7.3%)</td>
<td>22,009 (15%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Atlanta and Rural Georgia</td>
<td>3,588 (9%)</td>
<td>7,663 (5.2%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Iowa</td>
<td>2,20 (1.2%)</td>
<td>23,444 (16%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Detroit</td>
<td>5,485 (29%)</td>
<td>19,047 (13%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>New Mexico</td>
<td>115 (0.6%)</td>
<td>5,278 (3.3%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Utah</td>
<td>36 (0.2%)</td>
<td>6,456 (4.3%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Seattle-Puget Sound</td>
<td>597 (3.2%)</td>
<td>18,539 (12%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Median county income (in 2010, $)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$21,116–$50,254</td>
<td>3,303 (21%)</td>
<td>25,298 (20%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>$50,255–$55,092</td>
<td>4,571 (29%)</td>
<td>24,622 (19%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>$55,093–$62,512</td>
<td>3,695 (23%)</td>
<td>25,356 (20%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>$62,513–$71,724</td>
<td>2,313 (15%)</td>
<td>26,943 (20%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>$71,725–$110,970</td>
<td>1,975 (12%)</td>
<td>27,008 (21%)</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>
distribution. The estimates are reported by age at diagnosis. Individuals diagnosed before the age of 65 survive longer than older patients. Comparing the histograms on the first row (RMSTs for blacks) with the histograms on the second row (RMSTs for whites) we see that whites experience longer survival. Comparing the histograms on the second row (RMSTs for whites) with the histograms on the third row (RMSTs for blacks after the hypothetical intervention) we observe a shift of the RMSTs for the black population towards that of the white; however, the disparity is not eliminated.

Figure 2 displays the density of the estimated disparity (black line) and residual disparity (gray line) across covariate strata, separating the three age groups. The distribution of the disparity measures displays high variability. This variability reflects both the heterogeneity in the underlying disparities across covariate strata as well as the uncertainty in the estimates themselves. Greater disparities are noted for patients diagnosed before the age of 65 years. Had we intervened eliminating disparities in stage at diagnosis, the residual disparity gets closer to zero but still important differences are observed.

By combining the estimates employing random effect meta-analysis, a significant disparity in overall survival is observed for which the difference in RMST truncated at year 5 for black versus white is $-4.0$ months [confidence interval (CI), $-4.6$ to $-3.3$; Table 3].

The residual disparity in colorectal cancer survival had we intervened on the stage at diagnosis distribution is $-2.6$ months (CI, $-3.4$ to $-1.7$, Table 3), indicating that elimination of disparities in stage at diagnosis may lead to a reduction of overall survival disparities in the first 5 years after diagnosis of 35% (CI, 11%–55%; Table 3). Our nonparametric analyses account for heterogeneities across the several factors. Black–white disparities are slightly larger among men and the impact of the hypothetical elimination of stage at diagnosis disparities appears stronger among women (44% vs. 26%). Estimates of the disparity by age groups reveal that patients diagnosed before the age of 65 experience significant greater disparities in survival with respect to the elderly population (disparity$_{\text{age}<50} = -5.9$, disparity$_{\text{age}50-65} = -5.1$

<table>
<thead>
<tr>
<th>Stage</th>
<th>Estimate (CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>-3.2 (-4.4, -1.7)</td>
</tr>
<tr>
<td>II</td>
<td>-2.4 (-3.6, -1.0)</td>
</tr>
<tr>
<td>III</td>
<td>-0.8 (-2.1, 0.6)</td>
</tr>
<tr>
<td>IV</td>
<td>-2.6 (-5.9, -1.5)</td>
</tr>
</tbody>
</table>

Table 2. Disparity in restricted mean survival time truncated at month 60 since diagnosis by stage at diagnosis

Figure 1.
Histograms of restricted mean survival time truncated at month 60 by age at diagnosis for white, black, and black after the hypothetical intervention. The intervention consists on the shift of the stage distribution of the black population to match that of the white population, resulting in the elimination of the disparities in stage at diagnosis. The restricted mean survival time is estimated in strata of confounding factors (age at diagnosis, grade of tumor differentiation, county median income, date at diagnosis, gender, and state). The y-axis represents the percent of individuals for which a certain RMST is estimated and the x-axis represents the RMST in months.
Stage at Diagnosis and Black-White Colorectal Cancer Survival Disparities

Figure 2.
Density of difference in restricted mean survival time by age at diagnosis for white versus black (disparity, black) and white versus black under the hypothetical intervention (residual disparity, gray) across confounders’ strata. The intervention consists on the shift of the stage distribution of the black population to match that of the white population. Dashed lines represent the estimated disparity and residual disparity using random effect meta-analysis for each age group.

Discussion
This is the first study, to our knowledge, to employ a counterfactual causal inference approach to quantifying the role of stage in diagnosis in black–white cancer survival disparities. The method relies on the assumption that we adjust for factors that affect survival and stage at diagnosis and that differences in stage at diagnosis are mostly the result of differential access to preventive care and may be mitigated by fostering equal access to screening and follow-up practices.

Using this approach, we assessed in the SEER population of colorectal cancer patients how much black–white survival disparities would be reduced if disparities in stage at diagnosis were eliminated. We find that black patients within five years since diagnosis survive approximately 4.0 months less than whites. The elimination of disparities in stage at diagnosis would contribute to a reduction of about 35% of this difference among colorectal cancer patients.

Our patient population, diagnosed between 1992 and 2005, experienced important disparities in access to screening (26,27). However, in the 2010 CDC Behavioral Risk Factor Surveillance System survey, blacks and whites reported more similar screening rates (66.3% for whites and 65% for blacks; ref. 28). Universal insurance coverage through Medicare and the recent elimination of out-of-pocket costs for screening likely contributed to these changes. It remains to be seen whether reductions in disparities in screening will result in equalization of the distribution of stage at diagnosis and reductions in black–white disparities as anticipated by our analysis. Our data suggest that, to the extent that similar screening rates lead to the elimination of disparities in stage at diagnosis, survival disparities could substantially reduce in the near future. Importantly, we find that such improvements might be particularly apparent for subpopulations (e.g., women and the elderly). The reasons for these heterogeneities should be further researched. We hypothesize that differentials in comorbidities might explain the differences in the impact of stage at diagnosis.
actions between race and stage at diagnosis (18). In our study, we
found that stage differences accounted for 60% of the excess mortality among blacks. That study
from Pennsylvania State Tumor Registry (16), found that differ-
ences in colorectal cancer survival (30) and their interplay
with socioeconomic characteristics are important drivers of
cancer outcomes as well. Furthermore, we show that there are
disparities in colorectal cancer survival disparities. This might be relevant for
the study of determinants of black–white disparities in other
cancer outcomes as well. Therefore, we show that there are still determinants of the black–white disparities that are unex-
plained. Therefore, other factors may play a larger role in explain-
ing such disparities among younger groups and in males. In parti-
cular, socioeconomic characteristics are important drivers of
disparities in colorectal cancer survival (29) and their interplay
need to be further investigated.

A popular approach to estimate the extent to which racial/
ethnic disparities are explained by intermediate factors is to
estimate the role of screening and stage-specific relative colorectal cancer
survival in a population aged 50 years and older, estimated that screening and stage-specific survival explained a little more
than 50% of the disparity in colorectal cancer mortality between blacks and whites (30). Although a powerful tool that
can reflect the natural history of a complex disease, microsi-
mulation comes with some limitations. Large-scale microsimu-
lation models require many parameters that may not be avail-
able from existing data sources. In addition, the degree of
model detail does not go hand in hand with overall prediction
power and potential for bias in complex models is often
overlooked.

Our approach has several advantages. First, we formalized
the quantities of interest accounting for the causal nature of the
question. Adopting the counterfactual framework was key for
clarifying the objects of inference and the identification
conditions. Moreover, informed by an ecosocial understand-
ing of how people jointly embody diverse types of inequality
(3–4), we considered different colorectal cancer patients’ sub-
groups according to gender and age. We estimated differences
in the white and black patients using restricted mean survival
times, a clinically meaningful measure of disparity, employing
a nonparametric approach adjusting for potential confoun-
ders. Moreover, we report disparities on the difference scale,
which has clearer public health interpretation. Finally, we
evaluated the sensitivity of our analyses to selection bias due
to missing stage at diagnosis.

There are limitations to this analysis. Our estimates could be
biased due to unmeasured residual confounding. Comorbid-
ities were not measured in the SEER cancer registry and these
factors affect stage at diagnosis as well as cancer survival.
We expect this bias to lead to an overestimation of the impact
of the hypothetical intervention. Furthermore, as survival is
our outcome, lead-time bias might affect our estimates. In the
study we have mitigated the impact of this bias by excluding
patients diagnosed with in situ cancers. The direction of such
bias in our study would again be an over estimation. Microsi-
mulation would be valuable complement to our analyses to
assess the sensitivity of the results to this bias. Finally, stage at
diagnosis measure is a crude representation of actual cancer
natural history. Microsimulation comes with some limitations. Large-scale microsimulation
models require many parameters that may not be avail-
able from existing data sources. In addition, the degree of
model detail does not go hand in hand with overall prediction
power and potential for bias in complex models is often
overlooked.

Our study carries important implications for future work. We
show the importance of accounting for heterogeneities across
gender and age to assess the role of stage at diagnosis in explaining
colorectal cancer survival disparities. This might be relevant for
the study of determinants of black–white disparities in other
cancer outcomes as well. Furthermore, we show that there are still determinants of the black–white disparities that are unexplained. Therefore, other factors may play a larger role in explaining such disparities among younger groups and in males. In particular, socioeconomic characteristics are important drivers of disparities in colorectal cancer survival (29) and their interplay need to be further investigated.

A popular approach to estimate the extent to which racial/ethnic disparities are explained by intermediate factors is to estimate the role of screening and stage-specific relative colorectal cancer survival in a population aged 50 years and older, estimated that screening and stage-specific survival explained a little more than 50% of the disparity in colorectal cancer mortality between blacks and whites (30). Although a powerful tool that can reflect the natural history of a complex disease, microsimulation comes with some limitations. Large-scale microsimulation models require many parameters that may not be available from existing data sources. In addition, the degree of model detail does not go hand in hand with overall prediction power and potential for bias in complex models is often overlooked.

Our approach has several advantages. First, we formalized the quantities of interest accounting for the causal nature of the question. Adopting the counterfactual framework was key for clarifying the objects of inference and the identification conditions. Moreover, informed by an ecosocial understanding of how people jointly embody diverse types of inequality (3–4), we considered different colorectal cancer patients’ subgroups according to gender and age. We estimated differences in the white and black patients using restricted mean survival times, a clinically meaningful measure of disparity, employing a nonparametric approach adjusting for potential confounders. Moreover, we report disparities on the difference scale, which has clearer public health interpretation. Finally, we evaluated the sensitivity of our analyses to selection bias due to missing stage at diagnosis.

There are limitations to this analysis. Our estimates could be biased due to unmeasured residual confounding. Comorbidities were not measured in the SEER cancer registry and these factors affect stage at diagnosis as well as cancer survival. We expect this bias to lead to an overestimation of the impact of the hypothetical intervention. Furthermore, as survival is our outcome, lead-time bias might affect our estimates. In the study we have mitigated the impact of this bias by excluding patients diagnosed with in situ cancers. The direction of such bias in our study would again be an overestimation. Microsimulations would be valuable complement to our analyses to assess the sensitivity of the results to this bias. Finally, stage at diagnosis measure is a crude representation of actual cancer stage; this might lead to an underestimation of the role of stage at diagnosis.

Other factors, such as socioeconomic position and quality of treatment have also been reported to contribute to explain both differences in colorectal cancer incidence and survival among racial/ethnic groups (31–32), but their doing so does not mean that interventions to modify stage at diagnosis are irrelevant. We note that a recent comprehensive screening program deployed in the state of Delaware resulted in the elimination of stage at diagnosis disparities and in the reduction of survival disparities (33). Finally, in our study we evaluate the role of stage at diagnosis, a more direct intervention concerns screening uptake and follow-up rates. Linked SEER-Medicare data would allow evaluating potential screening regimes in the elderly population. A recent report from the CDC (28) found that colorectal cancer screening rates among whites were substantially higher than those among Hispanics, Asian/Pacific Islanders, and American Indian/Alaska Natives (AI/AN). Important disparities in cancer survival have been observed comparing white patients with Hispanic and Asian patients. Therefore, future research will consider other racial/ethnic groups as well.
Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

Authors’ Contributions

Conception and design: L. Valeri, J.T. Chen, N. Krieger, T.J. VanderWeele, B.A. Coull

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Acquisition of data (provided animals, acquired and managed patients, provided facilities, etc.): L. Valeri, J.T. Chen

Analysis and interpretation of data (e.g., statistical analysis, biostatistics, computational analysis): L. Valeri, J.T. Chen, N. Krieger, T.J. VanderWeele, B.A. Coull

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