Role of Implicit Bias in Pediatric Cancer Clinical Trials and Enrollment Recommendations Among Pediatric Oncology Providers

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BACKGROUND: Provider implicit bias can negatively affect clinician-patient communication. In the current study, the authors measured implicit bias training among pediatric oncology providers and exposure to implicit association tests (IATs). They then assessed associations between IATs for race and socioeconomic status (SES) and recommendations for clinical trial enrollment. METHODS: A prospective multisite study was performed to measure implicit bias among oncology providers at St. Jude Children’s Research Hospital and affiliate clinics. An IAT was used to assess bias in the domains of race and SES. Case vignettes were used to determine an association between bias and provider recommendation for trial enrollment. Data were analyzed using Student t tests or Wilcoxon tests for comparisons and Jonckheere-Terpstra tests were used for association. RESULTS: Of the 105 total participants, 95 (90%) had not taken an IAT and 97 (92%) had no prior implicit bias training. A large effect was found for (bias toward) high SES (Cohen d, 1.93) and European American race (Cohen d, 0.96). The majority of participants (90%) had a vignette score of 3 or 4, indicating recommendation for trial enrollment for most or all vignettes. IAT and vignette scores did not significantly differ between providers at St. Jude Children’s Research Hospital or affiliate clinics. No association was found between IAT and vignette scores for race (P = .58) or SES (P = .82). CONCLUSIONS: The authors noted a paucity of prior exposure to implicit bias self-assessments and training. Although these providers demonstrated preferences for high SES and European American race, this did not appear to affect recommendations for clinical trial enrollment as assessed by vignettes. Cancer 2021;127:284-290. © 2020 The Authors. Cancer published by Wiley Periodicals LLC on behalf of American Cancer Society. This is an open access article under the terms of the Creative Commons Attribution-NonCommercial License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited and is not used for commercial purposes.

KEYWORDS: clinical trials, enrollment, implicit association tests, implicit bias, pediatric oncology.

INTRODUCTION

Advancements in cancer treatments, due in part to well-designed clinical trials, have led to an overall 5-year survival rate of >85% for pediatric patients with cancer.1 Clinical trial participation is the standard approach for improving outcomes for childhood cancer in academic centers.2 Because of the structured nature of pediatric clinical trials, participants receive uniformly adjudicated treatment and have lower mortality and complication rates.3 Despite these benefits, enrollment rates are variable and range from <20% to 86%.2,4 Known barriers to enrollment include structural challenges such as trial availability, clinical barriers such as eligibility restrictions, patient-level obstacles including travel and financial concerns, and provider-level concerns including time constraints and implicit bias.3,5,6

Among the St. Jude Children’s Research Hospital (SJCRH) affiliate network, we noted a wide range of clinical trial enrollment rates. Similar to national data, average enrollment approximates 50% to 60%; however, enrollment from some affiliate sites is as low as 20%. In an effort to understand this variation, we previously investigated physician-perceived barriers to clinical trial enrollment across the SJCRH affiliate network.7 We identified 3 major barriers: language discordance, transportation issues, and complex trial design. We also observed that socioeconomic status (SES) was a factor pediatric oncology providers occasionally considered when deciding whether to recommend a clinical trial (unpublished data).
Studies exploring the role of implicit bias in clinician decision making have demonstrated variable associations between implicit bias and patient care. Maina et al reviewed a decade of research regarding implicit bias in health care and found that implicit bias was associated with disparities in treatment in 6 of the 14 studies that examined outcomes. Implicit race bias has been associated with poorer communication in race-discordant clinical interactions, and research has demonstrated that children of a specific race may receive suboptimal care. For example, Sabin et al demonstrated that pediatrician implicit pro-White biases were associated with a reduced likelihood of prescribing appropriate narcotic pain medication for an African American patient. Implicit bias is negatively associated with supportive communication and length of clinical interactions for adult patients with cancer. Pediatricians implicitly associate African American patients with noncompliance, which may affect enrollment into clinical trials. However, to the best of our knowledge, the role of implicit bias in enrolling pediatric oncology patients into clinical trials has not been systematically or prospectively studied.

Maximizing clinical trial enrollment is a cornerstone of the mission of SJCRH. We hypothesized that implicit bias in the domains of race and SES influences recommendations by pediatric oncology providers for patient enrollment into clinical trials. To test this hypothesis, we measured implicit biases in race and SES via the implicit association test (IAT) and determined whether implicit bias scores were associated with provider recommendations for enrolling patients described in 4 case vignettes (CVs) into a clinical trial.

MATERIALS AND METHODS
The current study was a prospective, nontherapeutic, noninterventional study that included pediatric oncology faculty and advanced practice providers (APPs) involved in direct patient care at SJCRH and its affiliate clinics. The study was approved by the institutional review board at SJCRH. The 8 SJCRH affiliate clinics are located in Shreveport and Baton Rouge, Louisiana; Charlotte, North Carolina; Huntsville, Alabama; Johnson City, Tennessee; Peoria, Illinois; Springfield; Missouri; and Tulsa, Oklahoma. SJCRH is a tertiary specialized pediatric cancer center located in Memphis, Tennessee, whereas the affiliate clinics are in community-based hospitals. The racial demographics of the catchment areas for SJCRH and the affiliate clinics vary. Three areas have a slight majority African American population. One has a heterogeneous population with no majority racial group. Four catchment areas have a majority White population, and one has mixed racial demographics that include 10% American Indian. Patients in the affiliate network have access to pediatric cancer trials sponsored by SJCRH, and resources for clinical trial participation, including translated informed consent documents and patient advocacy training, are available at all sites.

Pediatric oncology providers at SJCRH and the affiliate clinics were recruited for participation via email. A reminder email was sent at 3 weeks and again at 1 week before the close of the study period of 6 weeks. Before starting IATs and CVs, we asked each participant their practice location, provider type (physician or APP), years in practice, number of IATs previously taken, and history of implicit bias training. No additional demographic or identifying information was collected from participants to preserve confidentiality and anonymity. APPs (ie, nurse practitioners and physician assistants) were included in the study because they actively participate in discussions regarding clinical care and research and are partners to physicians in the decision-making process for clinical trial enrollment in pediatric oncology.

We conducted the current study in collaboration with Project Implicit, a nonprofit organization that studies and measures implicit social cognition. The order of IATs and CVs was randomly assigned independently by Project Implicit staff. Testing was performed using a secure website.

Study Measures
We used race and SES IATs to measure how pediatric oncology providers associate specific traits with distinct social categories. IATs are timed cognitive tests, measuring the relative association strength between 2 pairs of concepts, including a target concept such as race (eg, European American vs African American) and an evaluation concept (eg, good vs bad). For the SES IAT, positive scores indicated an implicit preference for upper SES over lower SES. Positive scores on the race IAT indicated implicit preference for European American over African American individuals. Negative scores indicated the opposite preferences. Both the race and SES IATs required approximately 10 minutes each to complete.

The CVs were written by senior physicians on the study team and had not been used previously (see Supporting Table 1). Each participant was given 4 CVs in 2 domains (high/low SES and White/African American race). CVs are a validated strategy of data collection to demonstrate associations of race and social bias. All 4 CVs
described a new pediatric cancer diagnosis with an option to enroll the patient into a phase 2 or phase 3 clinical trial. We chose 4 diagnoses for the CVs that were representative of 4 frontline SJCRH-initiated clinical trials (Hodgkin lymphoma, acute lymphoid leukemia, acute myeloid leukemia, and infant leukemia). A degree of uncertainty was incorporated into the CVs because uncertainty may add to bias in real-life clinical decision making. Socioeconomic descriptions embedded in CVs included employment type, educational level, housing, and insurance status. CV responses were timed (90 seconds). Responses were either yes or no to recommend enrolling the patient into a clinical trial, with elaboration embedded in the answer choices or as an optional written response (see Supporting Table 1).

**Statistical Analysis**

IAT scores were calculated using a standard scoring algorithm. Scores ranged from −2 to +2, with 0 indicating no relative preference between conditions. CVs were scored as 0 to 4 (0 indicated enrolling no CV patients and 4 indicated enrolling all 4 CV patients).

Random assignment resulted in 6 possible orders of IAT and CV tests. We randomized study elements to remove the potential effects of participants viewing the IAT prior to the CV or vice versa. To assess IAT effect sizes, we used the Cohen d statistic, which is a standardized effect size measure. Cohen d values are interpreted as follows: a Cohen d of 0.2 indicates a small effect, a Cohen d of 0.5 indicates a medium effect, and a Cohen d of 0.80 indicates a large effect. We calculated Cohen d scores to measure the standardized distance from the mean IAT scores to an ideal score of 0. We also calculated Cohen d values to measure the mean difference in IAT scores between providers from SJCRH and the affiliate clinics using pooled within-group standard deviations.

Summary statistics included the mean, standard deviation, median, and range for continuous variables and the count and frequencies for categorical variables. Two-sample Student t tests or Wilcoxon rank sum tests were used for continuous variables and chi-square or Fisher exact tests were used for categorical variables. One-sample Student t tests or Wilcoxon signed-rank tests were used to determine whether the sample means or medians were different from 0. The nonparametric Spearman rank correlation (Spearman correlation coefficient [ρ]) and Jonckheere–Terpstra tests were used to ascertain correlations between race and SES IAT and CV scores. All P values were 2-sided, and P < .05 was considered statistically significant. Analyses were performed using SAS (version 9.4) or R (version 3.6.1) statistical software.

**RESULTS**

Of the 251 providers invited, 105 (42%) completed all 3 components of the study (race IAT, SES IAT, and CVs). Four additional participants were excluded because they did not complete all 3 components. The participants included 81 providers from SJCRH (77%) and 24 from the affiliate clinics (23%). The overall distribution by provider type was similar noted between SJCRH and the affiliate clinics (P = .7). The majority of providers (90%) had not taken any IATs.

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**TABLE 1. Participant Characteristics by Site**

| Characteristics                        | SJCRH N = 81 No. (%) | Affiliate Clinic N = 24 No. (%) | Total N = 105 P a |
|----------------------------------------|-----------------------|--------------------------------|------------------|
| Type of provider                       |                       |                                |                  |
| Physician                              | 25 (31)               | 15 (62.5)                      | 40               |
| APP                                    | 56 (69)               | 9 (37.5)                       | 65               |
| Years of practice                      |                       |                                |                  |
| <10                                    | 39 (48)               | 10 (42)                        | 49               |
| 10-20                                  | 23 (28)               | 9 (37)                         | 32               |
| >20                                    | 19 (24)               | 5 (21)                         | 24               |
| Taken an IAT before                    |                       |                                |                  |
| No                                     | 73 (90)               | 22 (92)                        | 95               |
| Yes                                    | 8 (10)                | 2 (8)                          | 10               |
| Previous formal implicit bias training |                       |                                |                  |
| No                                     | 76 (94)               | 21 (88)                        | 97               |
| Yes                                    | 5 (6)                 | 3 (12)                         | 8                |

Abbreviations: APP, advanced practice provider; IAT, implicit association test; SJCRH, St. Jude Children’s Research Hospital.

aThe Fisher exact or Pearson chi-square tests were used to compare groups.
before the study nor had they received previous implicit bias training (92%). The percentages of APPs and physicians who had taken previous IATs (P = .18) or received previous training (P = .48) did not differ significantly. The percentages of participants from SJCRH or the affiliate clinics also did not differ significantly with regard to previous IATs (P = 1.00) or training (P = .38).

We reported the ranges and medians, in addition to Cohen d values. Among all participants, we found a very strong preference for high SES over low SES.

The overall, SJCRH, and affiliate clinic mean SES IAT scores were 0.71 (Cohen d = 1.93), 0.69 (Cohen d = 1.88), and 0.76 (Cohen d = 2.09), respectively, which all were statistically significantly different from zero (P < .001) (Table 2). The median SES IAT scores were 0.75 for overall, 0.74 for SJCRH, and 0.82 for the affiliate clinics, indicating a strong preference for high SES (Fig. 1). Among all the participants for the race

### TABLE 2. IAT Scores by Site

| Bias Test   | SJCRH N = 81 | Affiliate Clinic N = 24 | Total N = 105 | SJCRH Versus Affiliate |
|-------------|--------------|-------------------------|---------------|------------------------|
|             | Mean (SD)    | Cohen d<sup>a</sup>    | P<sup>b</sup> | Mean (SD)    | Cohen d<sup>a</sup>  | P<sup>b</sup> | Mean (SD)    | Cohen d<sup>a</sup>  | P<sup>b</sup> |
| SES IAT     | 0.69 (0.37)  | 1.88 <.001             |               | 0.76 (0.36)  | 2.09 <.001             |               | 0.71 (0.37)  | 1.93 <.001             | 0.17 .56  |
| Race IAT    | 0.39 (0.42)  | 0.92 <.001             |               | 0.41 (0.35)  | 1.15 <.001             |               | 0.39 (0.41)  | 0.96 <.001             | 0.05 .84  |

Abbreviations: IAT, implicit association test; SD, standard deviation; SES, socioeconomic status; SJCRH, St. Jude Children’s Research Hospital.

<sup>a</sup>Cohen d value was calculated for SJCRH or affiliate clinic providers as an effect size to measure the standardized distance from the mean IAT scores to an ideal score of 0.

<sup>b</sup>P values were calculated using 1-sample Student t tests.

<sup>c</sup>Cohen d value was calculated as an effect size to measure the mean difference between SJCRH and affiliate clinic providers using the pooled within-group SD.

<sup>d</sup>P values were calculated using 2-sample Student t tests.

**Figure 1.** Box and whisker plots of implicit association test (IAT) score distributions by site. The zero point indicates no preference in either direction. The median is represented by dark horizontal bars. For both race and socioeconomic status (SES), the median values did not differ between providers at St. Jude Children’s Research Hospital (P = .57) and the affiliate clinics (P = .84).
TABLE 3. Continuous IAT Scores for SES and Race Based on Randomization Ordera

| Variable          | Overall               | CVs First N = 29 | CVs Not First N = 76 | p
|-------------------|-----------------------|------------------|----------------------|----------
| SES IAT score     |                       |                  |                      |          |
| Mean (SD)         | 0.71 (0.37)           | 0.52 (0.45)      | 0.78 (0.30)          | .007     |
| Median (range)    | 0.75 (−0.53 to 1.42)  | 0.53 (−0.53 to 1.34) | 0.81 (−0.01 to 1.42) |
| Race IAT score    |                       |                  |                      |          |
| Mean (SD)         | 0.39 (0.41)           | 0.37 (0.39)      | 0.40 (0.42)          | .78      |
| Median (range)    | 0.42 (−0.51 to 1.49)  | 0.38 (−0.33 to 1.49) | 0.47 (−0.51 to 1.25) |

Abbreviations: CV, case vignette; IAT, implicit association test; SD, standard deviation; SES, socioeconomic status.
aRandomization assignment order was CVs before IATs versus IATs before CVs.

TABLE 3. Continued...
The majority of providers offered clinical trials in all 4 CVs, and neither SES nor race bias were found to significantly affect CV scores. This finding is unsurprising because pediatric oncology is a clinical trial–centric field, and most children with cancer in the United States are offered trial enrollment at some point during cancer care or survivorship. SJCRH is a research hospital, and providers at SJCRH and its affiliate clinics are encouraged to enroll patients into clinical trials. The limited qualitative data in the current study suggested that providers believe trial enrollment is beneficial. It is interesting to note that some providers directly indicated their consideration of social factors when deciding to offer clinical trial enrollment. The offer for trial enrollment in the CVs suggests that the standard approach at large pediatric oncology centers to recommend clinical trial enrollment may supersede the potential effect of other factors, such as implicit bias.

Although the participant IAT scores did not appear to affect CV scores, the randomization order was significant for SES. The participants who completed the CVs first demonstrated less bias on the SES IAT. Therefore, the CVs may have functioned as a limited educational tool, personalizing patients and prompting awareness of the participants’ potential biases.

We did not collect the participants’ demographic information (eg, race, ethnicity, or sex) to protect anonymity and thus could not use this information to analyze potential differences in implicit bias. Although to our knowledge IATs are the most widely used instrument with which to identify bias, these tools have their own limitations and should be interpreted in context. IATs are sensitive to the context in which they are taken, because scores can change from one test to another. Despite these limitations, IATs capture attitudes that are distinct from those of self-reports. Although CVs are constrained in their ability to represent or convey nuances of real-life clinical care, they are a tool used to study medical decision making and have been used in similar studies. Our vignettes were designed specifically for the current study and thus were not previously used or pretested, which is a limitation of this study. Furthermore, each participant in the current study was exposed to all 4 CVs, which may have impacted their responses. Although the participation rate for this study was similar to that of other surveys of health care providers, selection bias may have affected the results. However, the similar distribution of participant types across sites assuages that concern.

The findings of the current study have highlighted a need for increased awareness of and formal education regarding implicit bias, as well as interventions that support diversity and inclusion. Educational curricula may include awareness tools, such as IATs, didactics regarding social determinants of health, and interactive workshops exploring provider bias. Expert facilitation is essential to diffuse defensiveness in implicit bias education. Although strong implicit bias was not found to be associated with recommendations for enrollment into clinical trials as defined by theoretical CVs, recent evidence has demonstrated that implicit bias education can have a positive impact on providers. Diversity and inclusion education have been emphasized for trainees by formal organizations such as the Accreditation Council for Graduate Medical Education. The data from the current study have suggested that this education should be offered to providers at all phases of their career continuum. In addition to formal training, diversity and inclusion efforts may be supported by offices dedicated to equity and supported by leadership, diversity officers, and human resource efforts focused on diversifying the health care workforce.

The lack of an association between the IATs and the CV scores suggested that implicit bias does not influence the conscious decisions of providers to recommend pediatric cancer clinical trial participation. However, CVs are an insufficient substitute for real-time clinical decision making, and further studies are needed to assess the effect of implicit bias on pediatric oncology patient-provider communication at the time of trial enrollment. Null results in implicit bias vignette studies may be the result of the vignette’s inability to capture the high time pressures and cognitive demands of busy real-world clinical settings. Previous studies in fields such as surgery and adult medicine demonstrated that interventions, including consensus treatment guidelines and protocols, can mitigate the effects of implicit bias on clinical care. For example, in a real-world study of the association between provider implicit bias and hypertension treatment that demonstrated null results, the authors speculated that emphasis of guideline adherence in the organizations studied may mitigate the impact of implicit bias on treatment decisions. Specifically, checklists and protocols that discourage provider discretion are potentially protective against bias. The protocolized nature of pediatric oncology practice may be similarly protective. Additional interventions to address and mitigate implicit bias in medical settings and other industries include recruiting a diverse health care workforce, care checklists and algorithms, and patient navigators. Continued investigation and adaptation of these tools and training methods for pediatric oncology is warranted to further understand
implicit bias and its potential effect on patient care, including clinical trial enrollment.

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**AUTHOR CONTRIBUTIONS**
Dylan E. Graetz: Writing of the initial draft, analysis of the data, critical revision of the article for important intellectual content, and primary author of the article. Arshia Madni: Writing of the initial draft and critical revision of the article for important intellectual content. Jeffrey Gossett: Data collection and analysis, writing of the initial draft, and critical revision of the article for important intellectual content. Guolian Kang: Conceptualization of the study, data collection and analysis, writing of the initial draft, and critical revision of the article for important intellectual content. Janice A. Sabin: Study conceptualization and critical revision of the article for important intellectual content. Victor M. Santana: Study conceptualization, data collection and analysis, writing of the initial draft, and critical revision of the article for important intellectual content. Carolyn L. Russo: Study conceptualization, data collection and analysis, writing of the initial draft, critical revision of the article for important intellectual content, and corresponding author. All authors approved the final article as submitted and agreed to be accountable for all aspects of the work.

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