Black Caregivers’ Perspectives on Racism in ASD Services: Toward Culturally Responsive ABA Practice

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Abstract
Significant racial and ethnic disparities in health care and service access exist. In the present article, we reviewed qualitative studies investigating the racism-related experiences of Black caregivers of children with autism spectrum disorder (ASD) in the U.S. health care system. Specifically, we examined institutional racism (i.e., systemic racism) and individual racism directed toward Black families when they seek diagnoses and services for their children with ASD. Additionally, we summarized culturally responsive and context-specific practice guidelines to work collaboratively with Black caregivers of children with ASD for applied behavior analysis practitioners.

Keywords: Autism • Applied behavior analysis • Black caregivers • Racial discrimination • Systemic racism

Research has shown that there are disparities in health care and access to services between Black and white families in several different domains, such as diagnosis, access to care, and the quality of care in the United States (Liptak et al., 2008; Magaña et al., 2012; Magaña et al., 2015; Mandell et al., 2002; Montes & Halterman, 2011). Many studies have pointed out that Black children with autism spectrum disorder (ASD) are diagnosed later, more often misdiagnosed with a conduct-related or adjustment disorder, and more likely to be not diagnosed at all compared to their white counterparts (Mandell et al., 2002; Mandell et al., 2009; Williams et al., 2020; but see Emerson et al., 2016, for different results). Furthermore,
Black children were less likely to receive early intervention programs (Feinberg et al., 2011) and overall service access than white children (Liptak et al., 2008).

Researchers have attributed racial disparities to many factors, such as provider–caregiver interactions, accessibility to services, and geographical differences (Mandell et al., 2002; Williams et al., 2020), which are rooted in the United States’ long history of racial segregation and discrimination. Disparities in educational opportunities in communities of color have also contributed to there being few Black behavioral and mental health professionals (Williams et al., 2020), which often means Black caregivers are interacting with non-Black providers. Racism has a clear impact on caregivers, as previous research has shown that racial discrimination among Black mothers is a contributing factor to depressive symptoms (McNeil et al., 2014; Odom et al., 2010), a concerning finding that calls for a family-centered, culturally responsive practice approach targeting Black children with ASD.

**Toward Culturally Responsive and Context-Specific Applied Behavior Analysis (ABA) Practice**

In the present article, we examined studies in the social science and education literature on Black families’ experiences with racism in the U.S. health care system (Burkett et al., 2015; Dababnah et al., 2018; Gourdine et al., 2011; Lovelace et al., 2018; Pearson & Meadan, 2018; Solomon et al., 2015; Solomon & Lawlor, 2013). The main method for data collection used in these studies is qualitative caregiver interviews. Qualitative interviews are not commonly used by behavior analysts; however, these methods can be valuable sources of information when one investigates the conditions in which socially significant behavior is observed in a given verbal community. For these studies, Black caregivers’ reports of racist experiences in their interactions with the U.S. health care system are critical concerns that the behavior-analytic community needs to consider. By examining these studies, we wish to contribute to behavior analysts’ ability to widen their approaches to the investigation of the social validity of their practice when working with Black families (see Robertson, 2016).

There has been much discussion on culturally responsive practice in ABA (see the special issues on diversity and inclusion issue in *Behavior Analysis in Practice*). As Brodhead (2019) advocated, culture always matters when implementing ABA services. Such discussion often results in a broad and general set of themes and training content that are not context specific, given the difficulty narrowing down the characteristics of a given culture (see Fong et al., 2016, for discussion). This difficulty partly stems from a behavior analyst’s practice to operationalize each concept; and indeed, it is a daunting task to operationalize each cultural practice. Our approach provides an alternative method to develop context-specific practices. That is, we start by looking at the occurrences of verbal behavior of a group member related to racial discrimination in a particular context and analyze what environmental arrangements cause such verbal instances. Then, we can develop guidelines to prevent these issues in a specific context.

Our current article focuses on Black caregivers of children with ASD in the context of the U.S. health care system. Given racism may not be observable by simply looking at health care policies and regulations, we, the scholars, must turn to the lived experiences of people of color interacting with these institutions to examine their reports of racial discrimination. In the following sections, we briefly define our working definitions of racism. Then, we outline findings from several qualitative studies on how different types of racism can manifest themselves in the lives of families raising Black children with ASD (Burkett et al., 2015; Dababnah et al., 2018; Gourdine et al., 2011; Lovelace et al., 2018; Pearson & Meadan, 2018; Solomon et al., 2015; Solomon & Lawlor, 2013). Finally, after we explore the service system from Black caregivers’ perspectives and point to situations where context-specific issues might arise between Black families and ABA practitioners, we provide recommendations for ABA practitioners to prevent and address instances of racism. We believe that our article continues the important conversations of highlighting the need for culturally responsive and context-specific practice in ABA.

**Why Is Understanding Race and Racism Important?**

Prior to incorporating recommendations for practice, it is critically important to understand the definition of race and racism. Borrowing from the social sciences, we are using conceptual definitions, a set of abstract characteristics associated with a given concept/term (Timasheff, 1947), which is important when investigating race. Race is a “socially constructed belief that the human race can be divided into biologically discrete and exclusive groups” that are based on physical, cultural, or social classifications (Golash-Boza, 2016, p. 130; see also Matsuda et al., 2020, for their attempt to operationally define racism). As a related construct, racism represents ideologies and their resultant practices that introduce racial inferiority and superiority (Golash-Boza, 2016). In a social system, these practices can be enacted through individuals and institutions. Individual racism can be both intentional and unintentional and could manifest as disrespect, bigotry, devaluation, and dehumanization (Golash-Boza, 2016; Jones, 2000). Institutional racism (i.e., systemic racism) is manifested as unequal access to both materials and societal resources (e.g., education, employment, advocacy, infrastructure) for people of color, which includes access to quality medical services (see Golash-Boza, 2016, for a review; Jones, 2000). Thus, it is important to investigate how racism not only impacts the
individuals whom we as practitioners support but also manifests itself within all social institutions, such as education and health care.

Racism Manifested Through Institutions

The White community may have access to more information than the Black community. (Burkett et al., 2015, p. 3249).

Institutional racism manifested itself in several ways in our review of the literature of caregivers raising children with ASD. Specifically, we noted instances of racialized differences in access to resources (e.g., information, services), provider interactions (e.g., service rejections, few Black therapists), and clinical research, all rooted in systemic inequities. These findings situate the difficulties that Black caregivers of children with ASD face in a much larger historical and societal context, rather than attribute them to a lack of personal effort.

In the previous quotation, a Black caregiver of children with ASD reported racial differences in available information. This theme of differential access to information based on race arose often in the studies we reviewed. For example, studies reported some parents shared that, due to the lack of educational information about ASD, they had to learn about ASD by themselves, very often through the internet (Burkett et al., 2015; Lovelace et al., 2018). This type of challenge points to the dearth of educational programs for Black parents in their community. Without systemic and provider supports, Black caregivers needed to educate themselves on ASD.

Systemic challenges persisted for Black families in the ASD service system. Black caregivers reported multiple service rejections for reasons unknown to them (Lovelace et al., 2018). Pearson and Meadan (2018) noted that 9 out of 11 Black mothers in their study had to advocate for their children with ASD to access services. Similarly, Solomon and Lawlor (2013), while interviewing Black mothers of children with ASD in Los Angeles, noted that racial discrimination emerged as a frequent theme. That is, mothers reported that they and their children were mistreated by “institutions authorizing autism-related services” (p. 107). Furthermore, mothers in this study shared that their children were denied services and parents were “looked down” upon, and mothers noted that these negative experiences are mostly because of racial discrimination (p. 107). Other studies found that Black families experienced a lack of or a delay in services because of ineffective community institutions. For example, “nice programs” were far from their homes and inaccessible without a car (Lovelace et al., 2018, p. 12). Likewise, Pearson and Meadan (2018) reported that Black caregivers in rural areas shared they could not access ABA services as the centers were too far away and there were no in-home sessions because of the lack of nearby therapists and clinicians. Additionally, some Black caregivers noted a differential quality of treatment based on insurance type, and that children with public insurance did not receive equally good-quality health care service compared to private insurance (Burkett et al., 2015). Williams et al. (2020) found that when Black families sought services, they would face multiple difficulties in the health care system, and as a result, many of them would not ultimately receive services.

In addition to accessibility issues because of structural inequities, it is important to reflect on the lack of clinical research that has included the Black population (see Brodhead et al., 2014, for review of participants’ cultural background). Davenport et al. (2018), for example, conducted a systematic review aiming at examining the sample diversity in research on social skills interventions for children with ASD and whether the studies culturally adapted their interventions based on the sample characteristics. Their analysis found that in 79 U.S. studies, only 6.2% included Black children in the sample (compared to 72.1% white children), and no studies adapted their interventions for Black participants. Thus, when Black families seek support through the health care system, there is only a small chance that they will find a treatment that is tailored specifically to Black children and their families, despite the clear value of cultural adaptations (see Robertson, 2016, for the social validity of parent-implemented behavior intervention for Black children with ASD).

Racism Manifested Through Interactions With Health Care Providers

In addition to fighting against and eliminating systemic racism, ABA practitioners need to be cognizant of the ways in which individuals in the health care system can display discriminatory activities. The assumptions, stereotypes, and beliefs about clients, conscious or not, can impact a person’s ability to deliver relevant and effective services (Garb, 1997). For example, a service provider might have an assumption of a Black client based on race (prejudice) and act on that assumption (discrimination). One study found a Black mother of a child with ASD experienced racial discrimination in her interactions with her child’s doctor, saying,

You assume because I’m Black . . . that I might not be as smart. . . . There are assumptions like . . . I’m just this person that has multiple kids. This is a Black woman. If you see my three kids, you assume I don’t have a husband. (Dababnah et al., 2018, p. 329)

As this quote demonstrates, discrimination that caregivers report does not necessarily have to be explicit to be harmful. Rather, service providers may unconsciously make...
assumptions and related comments due to implicit bias (Hagiwara et al., 2020).

Disrespectful interactions with health care professionals led some Black parents to believe that unfair treatment was due at least partly to their race (Burkett et al., 2015; Dababnah et al., 2018; Lovelace et al., 2018; Pearson & Meadan, 2018; Solomon & Lawlor, 2013). For example, one mother said,

Like I’ve said I’ve had lots of bad experiences. I can’t think of anything else to put it on other than my color . . . and you know we’re in [primarily White neighborhood]. . . . I hate to say it’s because of your color. . . . I try to look at everything else and then when everything else doesn’t all line up then you have to say maybe it was my color. Maybe it was because I was Black. Maybe it was because I didn’t come in . . . talking in a certain way . . . maybe they had a different expectation of me and when they saw I had expectations for my child, maybe they didn’t want to fulfill that. You don’t want to think that way but you have to look at that. It’s too blatant for you not to. (Lovelace et al., 2018, p. 7)

In the next section, we discuss Black caregivers’ experiences facing racism in two health care situations: diagnosis and services.

Experiencing Racism During the Diagnostic Process

Lovelace et al. (2018) reported that race plays a significant role for Black mothers of children with ASD who navigate the health care system. For example, one Black mother described the challenges many Black caregivers encounter getting an ASD diagnosis for their children: “I felt like a lot of the psychiatrists and stuff that, like, do you think I’m lying? . . . I was ignored or what I was saying was trivial to them, and it was very demeaning” (Lovelace et al., 2018, p. 8). Although a timely diagnosis is crucial in the treatment of ASD symptoms (Harris & Handleman, 2000), Black parents of children with ASD reported multiple experiences of racism in the diagnostic process, which very often contributed to delayed diagnoses. A delay in diagnosis not only is counterproductive to skill acquisition but also can contribute to an increased risk of challenging behaviors. For instance, Solomon and Lawlor (2013) found a parent of a Black girl with ASD reported that a delayed diagnosis compromised her daughter’s safety, as she had severe elopement behavior.

The literature we reviewed provided numerous examples of health care providers’ disrespectful behavior toward caregivers, including neglecting parents’ worries regarding their child’s growth, minimizing caregivers’ knowledge regarding their children, making negative assumptions about Black caregivers’ socioeconomic status, and accusing parents of lying about their child’s symptoms (Burkett et al., 2015; Dababnah et al., 2018; Gourdine et al., 2011; Lovelace et al., 2018; Pearson & Meadan, 2018). For example, one mother of a Black child said,

I remember telling [pediatrician] that [child] is not saying any words. The pediatrician said, “Well, some kids, it takes some time . . .” . . . I kept going to the pediatrician. “Something’s not right. Why is he not speaking?” and [pediatrician] kept coming up with all these excuses. . . . (Dababnah et al., 2018, p. 328)

This mother tried to obtain help from a doctor to get her child evaluated but faced multiple obstacles in the health care system to do so. In another example, some health care professionals suggested caregivers institutionalize their child, even before asking the parents’ opinions or assessing their caregiving skills and resources (Gourdine et al., 2011).

Mandell et al. (2002) found that Black parents on average made three times more visits to doctors to obtain an ASD diagnosis than their white counterparts. Similarly, Black parents needed to make multiple visits to medical offices in order to receive a referral for a diagnostic evaluation (Dababnah et al., 2018; Pearson & Meadan, 2018). Due to the prolonged negotiation with health care providers, in some cases, these challenges led some parents to seek support through advocacy (Pearson & Meadan, 2018) or request intervention directly “without their provider’s referral” (Dababnah et al., 2018, p. 328). The protracted struggles for referrals, as well as health care providers’ attitudes toward Black families, were characterized by Black caregivers as frustrating, discriminatory, insensitive, and demeaning (Dababnah et al., 2018; Lovelace et al., 2018; Pearson & Meadan, 2018; Solomon & Lawlor, 2013).

Experiencing Racism While Seeking and Accessing Services

Even after Black families obtained their child’s ASD diagnosis, the interactions with health care providers were still challenging. Families experienced different discriminatory treatments, such as a wraparound service specialist’s avoidance of integrated community outings in a Black client’s neighborhood (Lovelace et al., 2018). The mother attributed the specialist’s repeated denials of the community outings to the specialist’s belief that the family’s neighborhood was not safe for community programs, although the family lived in a mainly white neighborhood. The mother stated, “I didn’t live right in the shoot ’em up central” (p. 7).

Another example was a Black mother’s report that her doctor wrongly judged her by saying, “‘Well, you can’t do this with Medicaid.’ I said, ‘I don’t even have Medicaid, I have private insurance’” (Dababnah et al., 2018, p. 329).
Even when their children received services, some caregivers reported that health care providers did not take into consideration parental knowledge regarding their children and their opinions regarding treatment options (Burkett et al., 2015; Lovelace et al., 2018). Although parents desired a good relationship with health care professionals, perceptions about how professionals evaluated their ability to implement effective treatment at home overshadowed how caregivers interacted with them (Burkett et al., 2015).

Caregivers’ previous experiences of explicit and implicit racism often contributed to parental behavior change, in that they became vigilant in interactions with health care providers. For example, a Black mother withdrew her child with ASD from a service because she felt her child was not welcomed, and the mother did not want to expose her child to negative experiences (Lovelace et al., 2018). When they did bring their children to professional care, caregivers reported being watchful of their children’s interactions with the professionals because they did not fully trust health care professionals (Burkett et al., 2015). Solomon et al. (2015) reported that the mother of a Black child with ASD was vigilant in health care settings because of the potential “intersection of race and disability (i.e., the direct and implicit ways in which race can exacerbate potential impediments to her son’s care)” (p. 550). Therefore, when the mother was asked to check off whether her son was Black while filling out the form during the doctor visit, she decided to write “human” instead. Furthermore, Lovelace et al. (2018) said, when seeking services, a mother sometimes modified her accent over the phone, as she feared rejection from professional services if they realized she was Black. The authors wrote, “Often parents have to resort to code-switching or speaking in a manner that can mask another’s perceptions as to their race or ethnicity. While on a phone call, Latisha said that on the phone she can speak “prim and proper” and doesn’t always “sound” African American, but the person figured it out and didn’t let her son in.” (Lovelace et al., 2018, p. 8)

**Recommendations for Culturally Responsive and Context-Specific ABA Practice**

Black families of children with ASD face multiple difficulties navigating the health care system. It is important that ABA practitioners recognize the potential barriers Black families may have overcome to arrive at the point where they request ABA service. Thus, ABA practitioners should embrace the perspectives and values shared by Black families to not only facilitate the effectiveness of their interventions but also fight against racial discrimination and promote equity2 in health care services (Burkett et al., 2015; Dababnah et al., 2018; Lovelace et al., 2018; Pearson & Meadan, 2018).

Although the barriers that Black families of children with ASD face may not always be tied to their race (e.g., white and other racial/ethnic groups also reported certain challenges; see Sansosti et al., 2012), the frequency with which racial discrimination is experienced by Black individuals and their caregivers is of great concern (e.g., Burkett et al., 2015; Dababnah et al., 2018; Lovelace et al., 2018). Black families have reported that having trusting relationships with health care professionals is very important to them. Their reports provide insights into the values and competencies that health care professionals need to develop to build a good relationship with Black families of children with ASD (Burkett et al., 2015; Pearson & Meadan, 2018).

This issue is timely for behavior analysts. The demand for ABA services in the United States has been increasing constantly over the years, “with a 1,942% increase from 2010 to 2018 and a 127% increase from 2017 to 2018” (Behavior Analyst Certification Board [BACB], 2019, p. 2). With the increasing demands, one would expect that more ABA practitioners will provide services to culturally diverse populations. Therefore, training on cultural diversity and cultural humility is needed for successful ABA service provision (see Dennison et al., 2019; Fong et al., 2016; Fong & Tanaka, 2013; and Wright, 2019, for discussion on cultural diversity and cultural humility). However, recent surveys with BACB-certified practitioners show that graduate training, fieldwork, and supervision do not provide adequate education on multiculturalism and diversity (Beaulieu et al., 2019; Conners et al., 2019) and that there are very few opportunities for continued education on these topics (Beaulieu et al., 2019).

In the following sections, we discuss implications and recommendations for ABA practitioners to fight against racism in the U.S. health care system.

**Combating Institutional Racism**

Although the BACB recently began providing public information regarding the racial and ethnic backgrounds of its certified ABA practitioners, the lack of racially diverse representation is apparent from the shortage of multilingual and multicultural personnel and training programs (Dennison et al., 2019). The behavior-analytic field can increase the diversity of ABA professionals through targeted opportunities for Black professionals to choose future careers in ABA and the support of individuals through scholarships, mentorships, and

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2 In the context of racial justice within the health care system, equality refers to providing standardized service to people of all ethnic backgrounds, whereas equity refers to ensuring culturally responsive service.
advocacy (Fong et al., 2016). Additionally, there has been an increasing effort to empower the Black community within the ABA field through the creation of networks, such as Black Applied Behavior Analysts, Inc. This, combined with a more inclusive and responsive behavior-analytic field, will more likely help mitigate institutional racism.

Furthermore, our field can help increase Black communities’ access to culturally relevant information on ASD; culturally responsive ABA approaches, assessments, and interventions for ASD; and parent training on policy and advocacy concerns that are important for racially diverse communities. In addition, the field can be more forward thinking, combating the lack of practitioners and clinical options in certain rural communities. Although the COVID-19 pandemic has increased the availability of telehealth options for caregivers and their children (see Unholz-Bowden et al., 2020, for a review of telehealth effectiveness), continued efforts to advocate for underserved populations are needed to alleviate some of the impacts of institutional racism.

Combating Provider Racial Bias

In some cases, ABA practitioners might not be aware of their own biases toward Black communities and may unintentionally engage in racial discrimination while providing services. Based on the summary of the aforementioned qualitative studies, we identified four main areas where ABA practitioners should be especially sensitive in order to diminish racism manifested via interactions: (a) build knowledge about Black cultural values, (b) show care and strengthen partnerships, (c) create culturally relevant interventions, and (d) be aware of one’s own prejudice and cultural values.

Build Knowledge About Black Cultural Values

ABA practitioners should make meaningful efforts to understand the cultural values of each family they serve, although one should not have an a priori assumption that each family adopts the same cultural values. Practitioners learn about the values that a particular family endorses during initial and ongoing assessments and service provision. Although there is considerable variation in cultural values within Black communities, we highlight two major cultural themes from the literature: (a) respect and (b) awareness of Black families’ attitudes toward ASD.

Respect Respect is an important value in the Black community, and it has been passed down in families over generations (Burkett et al., 2017). A child who “behaved” showed respect for others and reflected the discipline practices of their parents. As one Black father described, “You did not disrespect your mom or dad . . . even if your father wasn’t in your life” (Burkett et al., 2017, p. 498). They view parenting as an important way to teach respect to their children and would employ traditional and nontraditional discipline practices such as directives or mimicking the disruptive behavior of their children. Black parents tend to expect their children with ASD to live up to the same standards as those of their typically developing siblings (Burkett et al., 2015; Burkett et al., 2017). Understanding these values may explain some Black families’ preferences for certain treatment options and increase their adherence to treatment plans. Pearson et al. (2018) proposed that speech-language pathologists (SLPs) should value Black families’ beliefs and the role of respect in their lives without judgment. This can also inform how ABA practitioners should approach discussions of treatment procedures with Black caregivers before they are implemented.

Awareness of Black Families’ Attitudes Toward ASD The impacts of institutional and individualized racism have affected how Black families view ASD. Previous studies have highlighted some Black community members did not believe in an ASD diagnosis, associated shame with having a child with ASD, or did not immediately agree that a child needs an intervention (Dababnah et al., 2018; Lovelace et al., 2018; Pearson & Meadan, 2018). ABA practitioners have to be aware of the impact of social and cultural systems and their influence on Black families’ engagement with ASD. For instance, one Black mother suggested that professionals should carefully discuss ASD within the family’s broader sociocultural context and build a relationship with the parents before providing services or referring them to other professionals (Pearson & Meadan, 2018). Furthermore, one Black mother said,

I think that putting more knowledge out there specifically geared towards African Americans [is important]. Examples of African Americans that have children on the spectrum that are getting services, that are actually benefiting in some way, and encouraging them to be more involved in their child’s services, and their child’s education as well, it would really, really, really, really, really, help. (Lovelace et al., 2018, p. 12)

Show Care and Strengthen Partnerships

Respect within the family–professional relationship is very important (Burkett et al., 2015; Lovelace et al., 2018). To better understand how to show care and build a successful parent–professional partnership, we identify two areas: (a) listen to what Black caregivers have to say and (b) explain service provision, timelines, and objectives transparently.
Listen to What Black Caregivers Have to Say

Black caregivers shared that they need professionals who listen to what they have to say regarding their child and their concerns (Burkett et al., 2015; Lovelace et al., 2018; Pearson & Meadan, 2018). As presented earlier, professionals have disregarded Black parents’ concerns in various health care settings (e.g., diagnostics, service planning). Caregivers also reported their children’s diagnoses were delayed because their primary health care providers did not acknowledge early parental concerns (Dababnah et al., 2018). Thus, it is crucial to listen to Black parents, not only to establish a respectful relationship, but also to facilitate effective screening and intervention processes.

A part of developing a respectful therapeutic relationship with Black families is to include them when developing a service plan for their children. Dennison et al. (2019) discussed the importance of social validity in ABA when working with families from diverse backgrounds. Listening to and acknowledging families’ values about treatment outcomes should be important for quality ABA services. As one Black mother of a child with ASD reflected, “I don’t really go too much by the blueprint as they say. I try to adapt the blueprint to what I have” (Burkett et al., 2015, p. 3249). Therefore, without parental opinions and feedback, ABA practitioners will be uncertain as to whether a given intervention is relevant for a family and whether they will adhere to given recommendations.

Another way to listen to Black families is to examine parents’ beliefs about the child’s diagnosis, their expectations about treatment, and their knowledge regarding different treatment options systematically (see Mandell & Novak, 2005, for useful questions to ask). Similarly, La Roche et al. (2018) suggested the Cultural Formulation Interview could also be a beneficial clinical tool to understand cultural factors from parental perspectives related to ASD diagnosis and treatment. Moreover, from a functional contextualism approach, Taylor et al. (2019) developed training curricula that could equip ABA practitioners with techniques for compassionate care. Their training curricula focus on “active listening, collaborating with caregivers, understanding a family’s culture, being kind, asking open-ended questions, avoiding technical jargon, and caring for the entire family” (p. 660).

Explain Service Provision, Timelines, and Objectives Transparently

ABA practitioners should fully explain service provision, service timelines, and service objectives as clearly as possible. Transparency in service provision is an important factor in any family’s participation in services. A lack of transparency can impact adherence and continued participation in behavioral health services, and for Black families who experience an increased level of discrimination, service provision must be described in clear, relevant ways (Carpenter-Song et al., 2010, as cited in Williams et al., 2020). Williams et al. (2020) further suggested that transparency between service providers and Black clients will decrease mistrust and facilitate collaboration, which is particularly important in order to increase treatment effectiveness and generalization.

Create Culturally Relevant Interventions

ABA practitioners should adapt and modify their treatment goals to increase cultural relevance. One of the biggest mistakes that providers can make is thinking that there is a singular path to supporting Black families. Working with each family to co-create target goals that reflect the family’s value system improves the cultural relevance of treatment recommendations (Kauffman et al., 2008). By including families in treatment design, ABA practitioners will understand not only their preferences for specific goals but also how families’ experiences shape why certain decisions are made for their children (Burkett et al., 2015).

Creating culturally relevant assessment and intervention materials is also important. For example, Blake et al. (2017) adapted parent-mediated social communication interventions by modifying a picture of a father and a child to that of a mother and a child and changing the women’s dresses in the pictures to reflect the clients’ cultural norms. In another example, Pearson et al. (2018) discussed how SLPs modified their communication devices, such as the Picture Exchange Communication System (PECS) and a speech-generating device, to adapt to the racial background of families. Specifically, a therapist can alter a picture of a person in PECS to have the same skin tone as that of a Black student or make the generated voice of the speech-generating device match a Black student’s family member’s voice (dialect, tone, inflection, semantics, and other factors). Pearson et al. (2018) also suggested using social stories to teach culturally different expectations that a Black student might encounter between home and school, such as the loudness of a voice or an acceptable level of body movements.

Pearson et al. (2018) also provided valuable suggestions for skill assessments. They argued that SLPs should exercise caution when evaluating Black students’ language, and this recommendation is also critical for behavior analysts. Incorporating a Black linguistic consciousness (Baker-Bell, 2020) will help analysts in the evaluation of Black students’ speech patterns and allow them to understand the characteristic verb markings that are incorporated in Black language. As ABA professionals become more mindful of these characteristics when assessing their clients’ behavioral repertoire, they will improve the effectiveness and social appropriateness of interventions and assessments.

Be Aware of One’s Own Prejudice and Cultural Values

For all professionals, it is important to recognize that no one is free of prejudice. Many researchers (Ford & Airhihenbuwa,
2010; La Roche et al., 2018) thus have emphasized that clinicians should understand their own cultures, biases, and values when they provide services to Black clients in order to improve the quality of interactions with them. Similarly, ABA practitioners should be aware of their own culture and biases when working with clients (Fong et al., 2016; Fong & Tanaka, 2013; Wright, 2019). Fong et al. (2016) illustrated how preconceived notions can impact a behavior analyst’s effectiveness with families of different backgrounds. The authors offered examples of how the analyst should be aware of their own reactions to information that families share, especially when it differs from their own cultural norms such as attitudes toward family and gender roles. In line with the recommendations of Fong et al. to assess cultural bias, we think it is also important that ABA practitioners seek out professional development on implicit bias and how it can be triggered during the course of service (e.g., see more information on implicit bias training; Hagiwara et al., 2020). Other practice-oriented frameworks offer providers methods to address institutional racism and other forms of oppression within their work (e.g., see Shaia, 2019).

Conclusion

Despite some hints of improvements in the area of early diagnosis and referral to early intervention programs for Black children with ASD (Dababnah et al., 2018), general disparities in the health care system between Black and white families still persist in the United States. Black caregivers continue to face more obstacles navigating the U.S. health care system compared to their white counterparts, which leads to irreparable disparities in Black and white individuals with ASD. Through our review of qualitative studies that reflect the experiences of Black caregivers of children with ASD with health care service providers, we have found that Black caregivers face racism at almost all steps in the health care system. ABA practitioners and the behavior-analytic field must respond to this persistent social justice issue and act to eliminate the barriers Black families face when accessing the health care system, so that their children with ASD can receive proper and effective interventions to improve their quality of life without facing racism. By identifying the contexts in which such racial discrimination might occur, behavior analysts can develop culturally responsive and context-specific practice to prevent and eliminate racism in the health care system.

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