



Healthcare Disparities in Otolaryngology

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Abstract

Purpose of Review The purpose of this review is to summarize some of the recent research studies on healthcare disparities across various subspecialties within otolaryngology. This review also highlights the impact of COVID-19 pandemic on disparities and proposes potential interventions to mitigate disparities.

Recent Findings Significant healthcare disparities in care and treatment outcomes have been reported across all areas of otolaryngology. Notable differences in survival, disease recurrence, and overall mortality have been noted based on race, ethnicity, socioeconomic status (SES), insurance status, etc. This is most well-researched in head and neck cancer (HNC) within otolaryngology.

Summary Healthcare disparities have been identified by numerous research studies within otolaryngology for many vulnerable groups that include racial and ethnic minority groups, low-income populations, and individuals from rural areas among many others. These populations continue to experience suboptimal access to timely, quality otolaryngologic care that exacerbate disparities in health outcomes.

Keywords Healthcare · Disparities · Access · Otolaryngology · Workforce · COVID-19

Introduction

Health and healthcare disparities are pervasive in the USA and globally, leading to preventable morbidity and mortality. Importantly, these disparities are modifiable and therefore demand our urgent attention. *Healthcare disparities* are differences in access to healthcare for members of different groups that

contribute to inequities in health outcomes [1]. This is distinct from *health disparities* that refer to differences in the incidence, prevalence, mortality, and overall disease burden among specific groups [2]. Innumerable studies have documented healthcare disparities in the USA based on patient race/ethnicity, socioeconomic status (SES), insurance status, education level, language, gender, sexual orientation, and geographic location among many others [3–9]. “Social Determinants of Health (SDH)” are defined as the conditions in which people are born, grow, work, live, learn, and age that shape their daily life. SDH can be grouped into five domains: access to education, access to healthcare, financial stability, neighborhood, and community [10]. These factors are interconnected and collectively form the socioecological framework as shown in Fig. 1.

Research on healthcare disparities focuses on detecting, understanding, and ameliorating disparities [11, 12]. However, most of the research to date in otolaryngology has been on detecting disparities (showing that they exist) with a growing literature based on understanding mechanisms [11, 12]. Implementation research, trials, and health policy interventions are greatly needed within our field. This review article focuses on healthcare disparities in otolaryngology, including only recent and high impact articles despite the abundance of epidemiological literature in the setting of increased national focus on health disparities since 2020.

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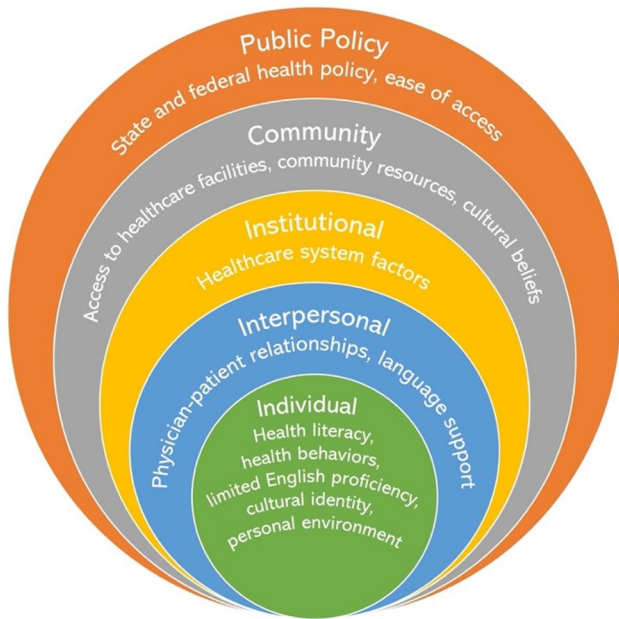


Fig. 1 Socioecological framework applied to healthcare with examples within each category

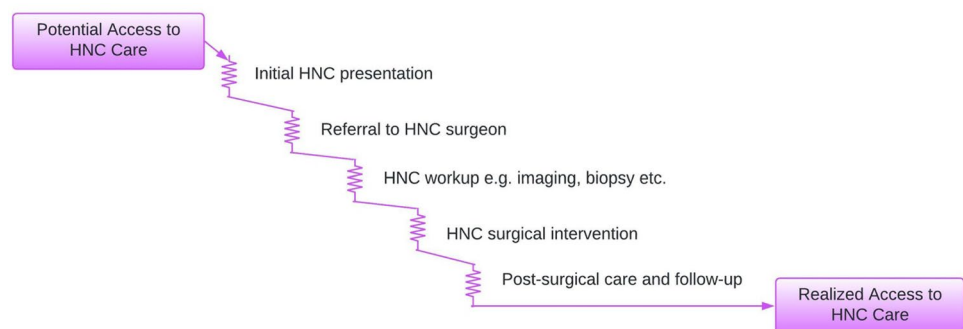
Disparities in Access to Healthcare

The Institute for Healthcare Improvement defines “access” as timely patient care in the right location with the right clinical team [13]. Disparities in access to care often underlie health inequities.

The “Voltage Drop” Framework of Access to Care

A “voltage drop” framework [14] can be used to show how patients may face multiple “resistance points” along the continuum of surgical care, as shown in Fig. 2 [15]. Patients may face barriers to initial presentation, referral to an otolaryngologist, and treatment; each may affect the patient’s outcome. Resistance points vary by disease, patient population, healthcare system, and public policy. Insurance, healthcare facilities in geographic proximity, affordability, and referral pathway support can help overcome resistance points [15].

Fig. 2 Voltage drop framework applied to head and neck cancer (HNC) care



Global Metrics for Disparities in Otolaryngology Care

Healthcare disparities are local, national, and global in scale. Globally, five billion people lack timely access to surgical and anesthesia care; the majority of these individuals are from low/middle-income countries (LMICs) [16]. The Lancet Commission on Global Surgery (LCoGS) recommended six metrics to measure access to timely, high quality, and affordable surgical care (Table 1) [16]. These measures have informed national surgical action plans and can be applied to otolaryngology [17–22].

Healthcare Disparities Across Otolaryngology Subspecialties

In this section, we discuss the evidence-based healthcare disparities across otolaryngology subspecialties.

Disparities in Head and Neck Cancer

HNC is the seventh most common cancer in the world, accounting for 3% of cancer deaths globally [23]. There are severe disparities in HNC prevalence and outcomes compared to other cancers [24].

Race and Ethnicity

A systematic review on health disparities in HNC found that 77% of the 148 reviewed articles in HNC examined race/ethnicity [25••]. Risk of HNC differs by race/ethnicity because of modifiable risk factors such as systemic racism, occupational exposures such as asbestos [26, 27], and lack of preventative care such as HPV vaccination. HPV- oropharyngeal cancer was more common among Black individuals (55.3%) compared to White individuals (30.3%) [28]. The incidence of HPV + oropharyngeal cancer is declining among Black individuals [29], but oral HPV infection is higher in Black individuals (9.8%) versus White (7.5%), Hispanic (7.2%), and Asian American (3.0%) individuals [30]. This is likely due to disparities in HPV awareness [31] and vaccination rates,

Table 1 Proposed indicators by the Lancet Commission on Global Surgery for measuring access to surgical care [16]

No.	LCoGS indicator
1	Access to timely surgical care
2	Surgical workforce density
3	Surgical volume
4	Perioperative mortality rate
5	Protection against impoverishing expenditure
6	Protection against catastrophic expenditure

concerning for future cancer risk [32]. The incidence of laryngeal cancer among Black males is higher compared to White males [29]. Nationally, Black patients disproportionately present at a younger age with advanced-stage laryngeal cancer and have nodal metastases compared to other races [33]. For oral cavity cancer, there is a higher incidence in Native American and Asian American/Pacific Islander populations [29]. Lao-tian individuals have 18.04 times higher risk of developing this nasopharyngeal carcinoma than non-Hispanic White individuals [34]. Compared to White patients, American Indian patients had a greater risk of alcohol abuse (68% vs 42%), current smoking (67% vs 49%), living more than 1 h away from a cancer center (81% vs 30%), and delayed presentation with stages III and IV HNC (74% vs 55%) [35].

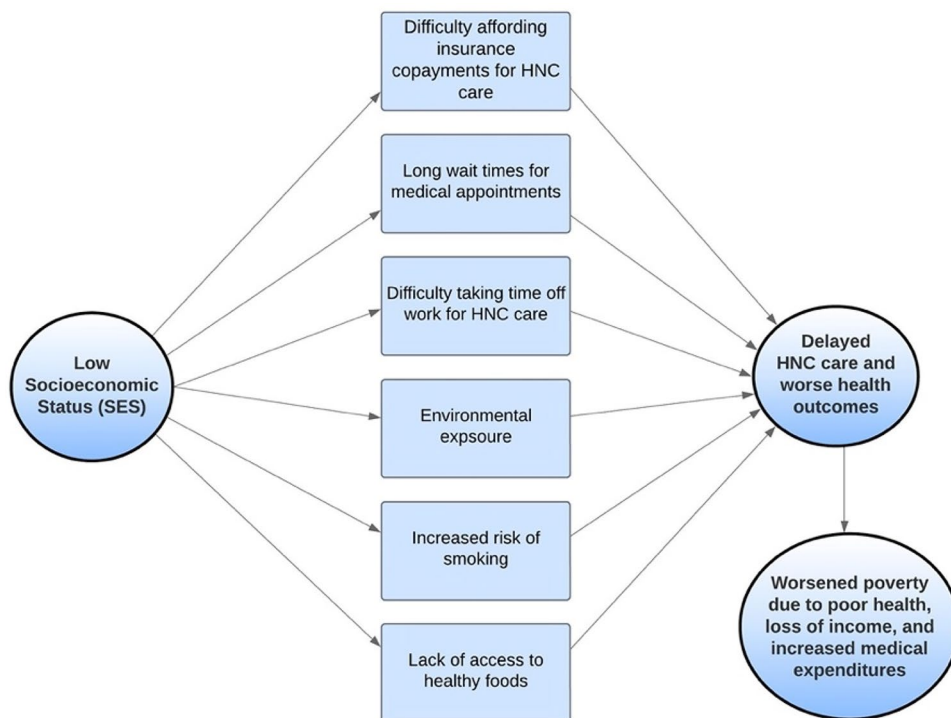
Significant racial and ethnic disparities also exist for HNC survival. A systematic review and meta-analysis reported that Black patients have inferior survival than White patients for HPV- oropharyngeal squamous cell

carcinoma [36]. Another meta-analysis found that Black HNC patients have poorer survival versus White individuals with a hazard ratio (HR) of 1.27 after controlling for socioeconomic factors [37]. In oropharyngeal squamous cell carcinoma, Black patients had worse survival compared to White patients (HR 1.22), after controlling for age, stage, and treatment [28]. The 5-year overall survival for laryngeal cancer was also lower for Black patients (52.7%) compared to White (60.6%), Hispanic (59.5%), and Asian (65.7%) patients [33]. American Indian patients with HNC and nasopharyngeal cancer experience worse survival compared with White patients [35, 38]. These racial/ethnic disparities call for interventions to eliminate the underlying factors.

Insurance and Socioeconomic Status

Low SES increases risks of worse health outcomes for HNC through multiple mechanisms (Fig. 3). A systematic review found that the risk of oral cancer is over 200% higher in low-income countries compared to high-income countries [39]. In the USA, low income, low education level, minority status, and lack of health insurance were each associated with advanced stage at presentation for HNC, with poverty most predictive [40]. The incidence rate of HPV-negative oropharyngeal cancer was higher among individuals with lower median household income [28]; lower SES was associated with worse survival [28]. Low SES is associated with advanced T-stage presentation and p16-negative status [41].

Fig. 3 Effect of low socioeconomic status for head and neck cancer care



Robust insurance and financial resources increase patients' ability to prevent and present early with HNC. The Affordable Care Act decreased the number of uninsured Americans from 50 to 27 million, with improved access to surgical care due to Medicaid expansion [42]. Insurance coverage does not fully protect against out-of-pocket costs. Relative out-of-pocket expenses for HNC patients was found to be 3.93% of patients' income compared to 3.07% of income for patients with other types of cancers [24]. Medicaid insurance is often not as robust as private coverage. HNC patients with Medicaid present at a later stage with larger tumor size, more distant metastases, and more lymph node involvement at diagnosis versus privately insured patients [43]. Medicaid patients are less likely to receive indicated surgical treatment [43]. Sittig et al. reported that 5-year overall survival for HNC patients with Medicaid, Medicare, or other government insurance was 50.6% compared to 65.5% for those with private insurance [44]. Within the USA, uninsured status was associated with higher mortality for HPV + oropharyngeal cancer (HR 3.12) [45]. Patients presenting with advanced-stage laryngeal cancer were more likely to be uninsured or have Medicaid [46]. Laryngectomy patients with Medicaid had increased length of stay and associated hospital charges compared to privately insured patients [47]. Insurance status also affects risk of mortality for laryngeal cancer. Among patients with advanced-stage laryngeal cancer, Medicaid patients have a 57% increased risk of mortality compared to those with private insurance [48].

Rurality

"Timely access to care" is challenging for rural areas in the USA and globally [16]. US otolaryngologists are disproportionately in high-income urban areas [49–51]. Rural counties therefore have fewer otolaryngologists per population [51]. The incidence and death rates from smoking-related cancers including laryngeal cancer are higher for rural residents [52]. Residence in a county with two or fewer otolaryngologists was found to be associated with poorer survival for HNC patients [40]. Rural individuals may have to travel much farther to see an otolaryngologist, making it harder to access specialty HNC care. Among HNC survivors, those in rural areas report lower health-related quality of life than those living in urban areas [53]. Moreover, the suicide rates for HNC patients living in rural areas are higher than the general population and those living in urban areas [54].

Sex and Gender

Disparities have been shown based on patient biological sex and gender identity. Males have a higher risk of HNC [55]. One study in Korea found that the incidence rate of

HNC was 0.19 in males compared to 0.06 in females, even when controlled for smoking and alcohol use [55]. Female patients present with more advanced laryngeal cancer than male patients [46], but have lower mortality rates [56]. Women with non-oropharyngeal HNC had better 5-year overall survival than men (56.3% versus 54.4%, respectively) [57]. Gender also plays a role in clinical trial participation. Although women make up 26.2% of the HNC patients in the USA, they comprise only 17% of patients in HNC clinical trials [58].

Patient Language and Health Literacy

Patients with limited English language proficiency (LEP) and lower health literacy have reduced access to care and poorer outcomes. Patients may not understand risk factors, treatment recommendations, or instructions for recovery. Low education level was associated with an increased risk of HNC (OR 2.50) [39]. Approximately 13.8% of the HNC survivors have inadequate health literacy [59]. Patients with LEP were underrepresented in most outpatient specialty clinics including otolaryngology [60]. Morris et al. found that the incidence rate of papillary thyroid cancer were lower in the counties with non-English speaking individuals without a high school degree, showing reduced healthcare access for these individuals [61]. Language support resources can reduce these barriers.

Disparities in Otology

A recent systematic review on disparities in otology included 52 research articles. Publications most commonly focused on racial/ethnic disparities [62•].

Race and Ethnicity

Disparities in race/ethnicity in otologic disease severity at presentation are well described. Black, Hispanic, and Asian populations have larger vestibular schwannomas at diagnosis versus White patients [63]. In the all-African American Jackson Heart Study prospective cohort, 29.5% of participants had tinnitus, highlighting the high prevalence [64]. There are racial/ethnic disparities in undergoing otologic surgical treatment. Non-Hispanic White patients were more likely than non-White patients to undergo surgery for recurrent acute otitis media or chronic otitis media with effusion that provided significant improvement in quality of life [65]. Compared to White patients, patients who identified as a racial minority were half as likely to pursue cochlear implantation despite qualifying for the procedure [66]. This may be due to racism, patient-provider interactions, social stigma, financial cost, insurance status/type, or lack of educational outreach [67]. Postoperative complication rates are

inequitable. African American patients had more skin-site complications after bone-anchored hearing aid implantation compared to other racial/ethnic groups [68].

Insurance and Socioeconomic Status

Better insurance coverage and higher SES both improve access to otologic treatments. Patients with Medicare who were also eligible for Medicaid were 41% less likely to use hearing care services such as visit to audiologists and hearing therapists and purchase of hearing aids, and twice as likely to report hearing issues with their hearing aids compared to high-income Medicare patients [69]. Nieman et al. conducted a successful community-delivered hearing care intervention for low-income and African American older adults with hearing loss in Baltimore. The intervention included training sessions to use a listening device, educational sessions on age-related hearing loss, counseling sessions on optimizing communication, and a brief aural rehabilitation session. After 3 months, they found improved outcomes [70].

Children with higher family income and parental education are more likely to receive cochlear implants at a younger age [71]. Similarly, adults with military insurance were 13 times more likely to receive cochlear implants than adults with Medicaid [66]. Medicaid-insured children receiving cochlear implants have fivefold greater odds of postoperative complications than those with private insurance [72]. This may be partly due to more missed follow-up appointments for children with Medicaid, which could serve as an intervention target. Education level and occupation have been found to influence timing of care for hearing loss [73]. Tran et al. found that patients with inadequate health literacy were more likely to present with more severe hearing loss [74]. Another study found that there is a higher incidence of chronic otitis media among patients that belong to a region with lower education level [75].

Rurality

Hearing loss and hearing impairment are more prevalent in residents of rural counties compared to urban counties, likely due to occupational and recreational noise exposure in rural areas where farming is a common source of income [76]. Rural residents described lack of geographic proximity to hearing care as a major barrier [76]. Rural residents have longer commutes to cochlear implantation centers versus urban residents [77, 78]. The time interval between onset of hearing loss and cochlear implantation was also greater for rural adult patients [77]. Children from rural regions with congenital hearing loss obtained hearing aids at the median age of 47.7 weeks after birth compared to only 26 weeks

for urban children [79]. Similarly, rural children received cochlear implants at the median age of 182 weeks after birth compared to 104 weeks for urban children, highlighting the delayed care for the children in rural counties [79]. Rurality is also associated with lower income and Medicaid coverage which exacerbates geographic barriers [77, 78].

Disparities in Rhinology

Healthcare disparities in rhinology are well established in certain domains, such as with allergic fungal rhinosinusitis (AFRS) and exposures to allergens, fungus, and pollution that contribute to allergic rhinitis, upper airway disease, and asthma.

Race and Ethnicity

White patients are significantly more likely to have seen a physician for their sinonasal symptoms compared to patients who identify as Hispanic and Native American [80]. From a language perspective, English-speaking patients were more likely to see a physician for their sinonasal symptoms compared to Spanish-speaking patients [80]. A study in South Florida found that Hispanic individuals tend to have a higher risk of severe chronic rhinosinusitis (CRS) and longer sinonasal symptom duration compared to non-Hispanic individuals [81]. Another study found that disproportionately fewer Black patients seek tertiary rhinology care than Black residents in southeast Wisconsin (9.3% vs. 13.8%) [82]. A retrospective study on AFRS consisted of significantly more African American patients with Medicaid or no insurance than expected [83]. Patients with AFRS were more likely to be residing in counties with higher poverty level, lower income level, and higher number of African American individuals than CRS [83]. Patients who identified as Black or Hispanic and were hospitalized for epistaxis had increased incidence of complications, longer length of stay, and consequently higher hospital charges compared to White patients [84].

Patients from minority racial and ethnic groups are underrepresented in endoscopic sinus surgery (ESS) outcome studies and CRS clinical trials. A cohort study investigating ESS outcomes reported that only 18% of the CRS patients electing ESS belonged to minority groups compared to the national average of 35% [85]. A systematic review found that among patients enrolled in prospective CRS clinical trials between 2010 and 2020 in the USA, 81.67% identified as White, 5.35% as Black, 1.27% as Asian, and 0.12% as American Indian [86]. Additionally, patient reported outcome metrics are often not implemented in non-academic centers or with patients with limited English proficiency [87]. Patients with increased healthcare access are more likely to be diagnosed with rhinologic conditions, therefore leading to inadequate data regarding true disease burden.

Insurance and Socioeconomic Status

Patients with CRS had longer wait times for ESS at public hospitals compared to private hospitals (68 days versus 45 days) and were more likely to be lost to follow-up [88]. Most (86%) public hospital patients with CRS in one local New York City study had Medicaid or no insurance compared to only 4% of CRS patients at private hospitals with better quality [88]. Another study reported that 60% of patients receiving tertiary rhinology care for CRS at an academic medical center in southeast Wisconsin had private insurance compared to only 49.8% of patients generally seen at the medical center [82]. Medicaid and self-pay insurance status were associated with ED presentation for acute rhinosinusitis among adult and pediatric patient population [89], likely reflecting poor access to primary care. Among patients with CRS electing ESS, patients with higher household income were more than twice as likely to experience a clinically meaningful quality of life improvement after ESS compared to patients with lower income level [90].

Climate Change

Climate change is exacerbating rhinologic disparities and is a growing focus in rhinology. Weather extremes induced by global warming have changed pollination patterns and may exacerbate allergic rhinitis and asthma. Worsening air pollution affects allergen-mediated respiratory diseases such as CRS [91]. This will disproportionately affect racial/ethnic minority groups and individuals with low SES, exacerbating disparities.

Disparities in Laryngology

A systematic review in adult laryngology found that disparities based on race/ethnicity were most frequently studied while the most commonly examined condition was laryngeal cancer [92•]. We have included laryngeal cancer in the head and neck cancer section and summarize other disparities below.

Race and Ethnicity

Individuals who identify as African American, Hispanic, or other minority race or ethnicity are less likely to report voice problems compared to White individuals [93], highlighting barriers to presentation. Additionally, Hispanic patients with voice disorders are more likely to delay care because they could not reach a medical office by telephone and due to long wait times at the doctor's office compared to White patients [93]. These disparities also exist for complication rates. Black patients are more likely to have iatrogenic laryngotracheal stenosis compared to White patients [94]. Moreover, among patients with laryngotracheal stenosis, Black patients are more likely to have a temporary tracheostomy

as well as be dependent on tracheostomy compared to White patients [94]. After stroke, Asian patients are more likely to experience poststroke dysphagia (OR 1.73) compared to Caucasian individuals [95].

Biologic Sex, Gender, and Gender Affirming Voice Treatment

Men are less likely to report voice disorders compared to women. 63.9% of women report voice disorders compared to only 36.1% men [96]. However, women are less likely to be referred to an otolaryngologist for their voice disorder compared to men [97]. On the other hand, women are more likely to receive treatment for their voice disorders compared to men [98].

Transgender voice care is a rapidly growing area of focus in laryngology. Transgender and nonbinary (TGNB) individuals have historically been marginalized in society and are vulnerable to healthcare access limitations [99]. About 27% of TGNB people living in rural areas travel more than 75 miles to access gender affirming medical care [100]. Travel is a barrier to gender-affirming healthcare needs such as voice therapy that requires frequent sessions. A 2015 US transgender survey revealed that 46% of transfeminine respondents who want to pursue behavioral voice therapy have not received it yet [100]. Roblee et al. suggested that the use of telehealth may be beneficial in mitigating barriers to voice therapy care, particularly for TGNB people living in rural counties.

Disparities in Pediatric Otolaryngology

Several systematic reviews describe the incidence, prevalence, management, and outcomes of various pediatric otolaryngologic diseases [101–103]. A systematic review on disparities reported that pediatric otologic diseases were most frequently studied and the most cited disparities include race and ethnicity, insurance status, and SES [101].

Race and Ethnicity

Numerous studies have reported differences in the prevalence and management of pediatric otolaryngologic diseases based on race and ethnicity [103]. A 2011 systematic review reported that Black children may have greater risk and higher prevalence for sleep-disordered breathing (SDB) compared to non-Black children [104, 105]. Black and Hispanic children are less likely to be diagnosed with frequent ear infections, hay fever, streptococcal pharyngitis, and sinusitis compared to White children, highlighting lack of access to care and underdiagnosis in children from racial and ethnic minority groups [106]. Hispanic children with thyroid cancer present with 27-mm tumor compared to 20 mm for White children [107]. Racial disparities in surgical interventions are well described. White children are more likely to undergo adenotonsillectomy compared

to Black or Hispanic children [104, 108]. According to the 2014 National Health Interview Survey, 12.6% White children received tympanostomy tubes for ear infections compared to only 4.8% of Black children and 4.4% of Hispanic children [109]. Black and Hispanic children are less likely to receive cochlear implantation early compared to White children [110]. Children from minority groups suffer more complications. Hispanic children have been found to have a higher mortality rate (28%) following tracheostomy than non-Hispanic White (7.9%) and non-Hispanic Black (8.7%) children [111]. Black children are more likely to have laryngeal stenosis, bronchopulmonary dysplasia, and an extended length of hospital stay following tracheostomies compared to White children [112].

Insurance and Socioeconomic Status

Socioeconomic disadvantage is the most studied factor contributing to healthcare disparities within pediatric otolaryngology. Spilsbury et al. found that children living in neighborhoods with severe socioeconomic disadvantage are more likely to be diagnosed with pediatric obstructive sleep apnea (OSA), even after controlling for premature birth, obesity, and race [113]. Children with public insurance have lower access to specialty care than those with private insurance. One study measured children's access to specialty care including otolaryngology care for common health conditions such as OSA and chronic bilateral otitis media [114]. They found that 66% of children with Medicaid were denied an appointment at specialty clinics, compared to 11% of children with private insurance [114]. Among 89 specialty clinics that accepted both insurance types, the average wait time for children with Medicaid was 22 days longer than that for privately insured children [114]. Children with Medicaid waited longer to be referred to an otolaryngologist for acute otitis media versus those with private insurance [115]. As mentioned in the otology section, children with public insurance are diagnosed with sensorineural hearing loss (SNHL) later than those with private insurance (20 versus 4 months age) and wait longer after diagnosis to receive cochlear implantation (25.5 versus 11 months) [116]. Children with Medicaid and self-pay insurance status are more likely to present to the ED for acute rhinosinusitis, highlighting problems accessing primary care [89, 117]. Among pediatric patients with acute bacterial sinusitis, those with Medicaid or no insurance had higher rates of complications compared to those with private insurance [100].

Other

Some studies have highlighted family factors contributing to disparities among children within otolaryngology. Children

with SNHL referred for cochlear implantation were more likely to have married parents (91% versus 70%) [118]. Another study found that children were less likely to attend their appointment at otologic medical clinics if their parents were younger with lower education level [119].

Disparities in Comprehensive Otolaryngology and Sleep Otolaryngology

Healthcare disparities also exist for conditions seen mostly by general otolaryngologists such as adult obstructive sleep apnea (OSA) and adult tonsillar diseases. A case-control study found that African American patients with sleep disordered breathing (SDB) are younger than Caucasian patients, suggesting that young African American individuals may have an increased risk of sleep apnea compared to young White individuals [120]. Black female patients with OSA are significantly younger at the time of diagnosis than White female patients and Black male patients with OSA have significantly lower oxygen saturation levels than White male patients [121]. Black male patients also present with more severe OSA at diagnosis with mean apnea hypopnea index of 52.4 ± 39.4 events/h compared with 39.0 ± 28.9 in White male patients, suggesting delayed diagnosis [122]. This study also found that Black male patients were 1.61 times more likely to have witnessed apneas and 1.56 times more likely to have drowsy driving at the time of OSA diagnosis compared to White male patients [122]. Black patients have decreased rates of surgery for OSA [123]. Lower household income, female sex, and Black and Hispanic race were associated with increased revisits for acute pain after adult tonsillectomy [124].

Disparities in Facial Plastic and Reconstructive Surgery

Healthcare disparities in aesthetic and reconstructive surgery are less well studied. One of the most serious complications of reconstructive flap surgery is postoperative sepsis. Sparenberg et al. found that African American race and male gender were significantly associated with sepsis following a reconstructive flap surgery [125]. Among HNC patients receiving free flap reconstruction, patients with Medicaid or no insurance were more likely to experience wound-related complications such as flap failure, fistula formation, donor site or primary site breakdown, and infection [126]. Another study reported that age over 55 years is associated with major medical complications following free flap reconstruction of the head and neck [127]. Other areas with significant attention include cultural and racial/ethnic differences in desired cosmetic surgery outcomes such as facial rejuvenation and appropriate dermatologic care for different skin tones. Keloid formation has been another topic of study with differences based on race/ethnicity [128, 129].

Effects of COVID-19 Pandemic on Healthcare Disparities in Otolaryngology

The USA has significant healthcare disparities despite having the world's most expensive healthcare system [130]. This is partly due to underinvestment in a social safety net and public health. The COVID-19 pandemic focused efforts on SDH.

The COVID-19 pandemic further compromised access to care and exacerbated disparities. Otolaryngology had among the most severe reductions in surgical volume and outpatient visits in the USA [131]. Massachusetts noted a 63% decrease in otolaryngology office visits and a greater decrease in operative procedures [132]. Consequently, there was an increase in the number of advanced presentations such as

Table 2 Proposed strategies and interventions to reduce healthcare disparities in otolaryngology

Category	Recommended interventions	Examples
Prevention	Implement health outreach programs	Proactive health outreach programs such as tobacco cessation programs geared towards patients with low-SES or from racial/ethnic minority groups can help reduce healthcare disparities [144].
Early detection	Extend health insurance coverage	Expand Medicaid and other government health insurance programs to improve access to healthcare. For instance, the Delaware Cancer Treatment Program provided insurance coverage for colorectal cancer screenings to the uninsured and low SES state residents that decreased the percentage of African American individuals with malignant diseases from 79 to 40% [145].
	Increase access to otolaryngologic care services	Improve geographic distribution of otolaryngologists by adapting the “hub-and-spoke model” of care and expanding residency training programs to areas with low trainee to population ratio [146].
Patient and community support	Prioritize patient education and community support	Interventions by community-based health workers have appeared to be effective for health education and social support [147]. Routine health literacy assessments in clinical practice may help identify patients who might require additional help in navigating the healthcare system [142].
	Community-delivered, convenient care for patients with barriers to care	Community-delivered hearing care intervention for low-income and African American older adults with hearing loss may improve outcomes similar to hearing aids and the usual standard of care [70].
	Enhance the diversity of clinical trials and intervention studies	Classification systems such as PROGRESS and CONSORT have been helpful in improving the reporting of health equity considerations in randomized clinical trials, systematic reviews and meta-analyses [142].
Support for patients with limited English proficiency (LEP)	Increase access to interpreter services	Use of professional interpreters for LEP patients is associated with decreased communication errors, increased patient comprehension, improved clinical outcomes, and increased satisfaction with clinical services [148].
	Improve institutional cultural competency	Hospital-wide initiatives such as cultural competency programs that include educational sessions to clinical and non-clinical staff focusing on healthcare disparities and culturally competent patient care may enhance health equity [149].
Workforce	Reduce implicit bias among otolaryngologists	Encouraging clinicians to engage with others' perspectives and implement intentional strategies to overcome biases can help reduce implicit bias [150].
	Enhance the diversity of otolaryngology workforce	Develop early pathways for UIM to have increased exposure to otolaryngology supplemented by mentorship and research opportunities; expand institutional and national efforts in residency recruitment [151].

orbital emergencies due to acute sinusitis [133]. Black and Hispanic children and those with Medicaid were disproportionately less likely to attend otolaryngology appointments [134]. During the pandemic, patients with Medicaid and lower income presented with higher disease severity for CRS and SNHL compared to those with private insurance and higher income [135]. An equity approach is needed to transform how we recover from the pandemic and improve access to healthcare.

Otolaryngology Workforce

The diversity of the otolaryngology workforce significantly influences access to affordable, high value otolaryngologic care. Racial and ethnic concordance between providers and patients leads to improved communication, increased patient satisfaction, and greater healthcare utilization which minimizes delays in care [136, 137]. Moreover, physicians from minority racial and ethnic groups are more likely to provide care to patients in medically underserved areas who may be low-income, uninsured, or insured with Medicaid [138, 139]. However, the demographics of the otolaryngology workforce in the USA do not match those of the general population. In 2016, African American individuals comprised 12.6% of the US population but made up only 2.1% of otolaryngology residents and 2.4% of otolaryngology faculty [140]. Similarly, Hispanic individuals are underrepresented at all levels in otolaryngology compared to the general US population. Only 5.5% of otolaryngology residents self-identified as Hispanic compared to 17% of the US population in 2016 [140]. Women are also a minority in the field of otolaryngology, accounting for only 17.1% of practicing otolaryngologists in the USA in 2020 [141]. It is pertinent to implement strategies nationwide to create a diverse otolaryngologic workforce to help mitigate healthcare disparities.

Achieving Health Equity in Otolaryngology

The field of otolaryngology has focused on detecting disparities and identifying potential factors involved, but progress can be made in identifying actionable targets for intervention to eliminate these disparities [12, 142]. The Accreditation Council for Graduate Medical Education reported that less than 5% of participating institutions had a systematic approach in place to address healthcare disparities within their organizations between 2017 and 2020 [143]. Therefore, there must be an urgent call to action to focus on identifying solutions and implementing strategies to reduce and eventually eliminate healthcare disparities (Table 2).

Conclusions

The need to eliminate healthcare disparities in otolaryngology has been recognized through the abundance of research studies in the field. These studies have identified disparities across all areas in otolaryngology for many vulnerable groups that include racial and ethnic minority groups, low-income populations, and individuals from rural areas among many others. However, there have been little effort in developing and implementing interventions to reduce these disparities.

Declarations

Conflict of Interest Sana Batool has nothing to disclose. Ciersten A. Burks reports United Against Racism Subspecialty Grant, Mass General Brigham. Regan W. Bergmark reports Nesson Fellow, Brigham and Women's Hospital (Institutional faculty grant funding and salary support for research on disparities in timely access to high quality surgical care); United Against Racism Subspecialty Grant, Mass General Brigham; Grant funding, I-Mab Biopharma.

Human and Animal Rights and Informed This article does not contain any studies with human or animal subjects performed by any of the authors.

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Papers of particular interest, published recently, have been highlighted as:

- Of importance
- Of major importance

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