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# The Prevalence and Characteristics of Children With Profound Autism, 15 Sites, United States, 2000–2016

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### **Abstract**

**Objectives:** Autism spectrum disorder (autism) is a heterogeneous condition that poses challenges in describing the needs of individuals with autism and making prognoses about future outcomes. We applied a newly proposed definition of profound autism to surveillance data to estimate the percentage of children with autism who have profound autism and describe their sociodemographic and clinical characteristics.

**Methods:** We analyzed population-based surveillance data from the Autism and Developmental Disabilities Monitoring Network for 20 135 children aged 8 years with autism during 2000–2016. Children were classified as having profound autism if they were nonverbal, were minimally verbal, or had an intelligence quotient <50.

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Disclaimer

The findings and conclusions in this article are those of the authors and do not necessarily represent the official position of the Centers for Disease Control and Prevention.

**Declaration of Conflicting Interests** 

The authors declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article. Supplemental Material

Supplemental material for this article is available online. The authors have provided these supplemental materials to give readers additional information about their work. These materials have not been edited or formatted by *Public Health Reports*'s scientific editors and, thus, may not conform to the guidelines of the *AMA Manual of Style*, 11th Edition.

**Results:** The percentage of 8-year-old children with profound autism among those with autism was 26.7%. Compared with children with non–profound autism, children with profound autism were more likely to be female, from racial and ethnic minority groups, of low socioeconomic status, born preterm or with low birth weight; have self-injurious behaviors; have seizure disorders; and have lower adaptive scores. In 2016, the prevalence of profound autism was 4.6 per 1000 8-year-olds. The prevalence ratio (PR) of profound autism was higher among non-Hispanic Asian/Native Hawaiian/Other Pacific Islander (PR = 1.55; 95 CI, 1.38–1.73), non-Hispanic Black (PR = 1.76; 95% CI, 1.67–1.86), and Hispanic (PR = 1.50; 95% CI, 0.88–1.26) children than among non-Hispanic White children.

**Conclusions:** As the population of children with autism continues to change, describing and quantifying the population with profound autism is important for planning. Policies and programs could consider the needs of people with profound autism across the life span to ensure their needs are met.

### Keywords

autism; surveillance; public health

Autism spectrum disorder (ASD; autism) is a developmental disability that manifests during childhood, affects multiple areas of development, and usually lasts throughout a person's life. Autism varies in presentation and can be associated with a wide range of social, communication, and behavioral problems and outcomes. This heterogeneity poses challenges in describing the needs of people with autism and making prognoses about future outcomes.

The *Diagnostic and Statistical Manual of Mental Disorders, Fifth Edition* (DSM-5) introduced severity levels for people with autism in the domains of social communication and restricted and repetitive behaviors to indicate the degree of support needed,<sup>2</sup> but they lack validation and their application might not be consistent.<sup>3,4</sup> Increasing efforts have been made in scientific and family advocacy communities to define people with autism who will likely have the greatest need for supports and services.<sup>5</sup> A recent Lancet commission defined "profound autism" as having autism with an overall or subdomain intelligence quotient (IQ) score <50 or being nonverbal or minimally verbal.<sup>6</sup> The commission indicated that children with profound autism were also more likely to have self-injurious behavior or epilepsy and require around-the-clock supervision than children without profound autism.<sup>6</sup>

As a result of this new and highly debated definition, there is a need for population-based information about the prevalence and characteristics of people who meet the criteria for profound autism. This information is critical for developing effective public health policies and programs to provide support across the life span. For example, one autism advocacy group is primarily focused on recognition and policy solutions for forms of autism that "by virtue of any combination of cognitive and functional impairments, necessitate continuous or near-continuous supervision, services and supports over the lifespan. Individuals in this category are often nonverbal or have limited use of language, are intellectually disabled, and, in a subset, exhibit challenging behaviors that interfere with safety and well-being." This new definition of profound autism may help to quantify the population with high levels of

need in these areas. In this article, we applied the newly recognized definition of profound autism to population-based surveillance data in selected US communities to (1) estimate the percentage of children with profound autism and compare their sociodemographic and clinical characteristics with those of children who have non–profound autism and (2) estimate the population prevalence for both groups across time.

### **Methods**

### **Study Design and Population**

The Autism and Developmental Disabilities Monitoring (ADDM) Network has conducted population-based surveillance of 8-year-old children with autism at selected US sites in even-numbered years since 2000 to estimate the prevalence of autism. The ADDM Network uses records-based surveillance methodology in which staff review records from medical, education, and service providers, as previously described. Children were classified as having autism after record review by clinicians who applied a standard case definition of ASD based on behavioral criteria for ASD described in the *DSM-IV* (2000–2014) and *DSM-5* (2016). Sites linked children's records to birth certificate information from their states to identify additional birth, maternal, and demographic characteristics. We obtained population denominators from the National Center for Health Statistics vintage 2018 bridged-race postcensal population estimates. For study areas comprising subcounty school districts, we implemented a standardization process using public school enrollment counts to adjust the population estimates. The Centers for Disease Control and Prevention determined this was a public health surveillance activity, deemed not to be research, and, therefore, exempt from institutional review board review.

We excluded 2 sites (Missouri and Pennsylvania) from this analysis because of a lack of access to educational records in which cognitive functioning data are typically included. We analyzed data from 15 sites (Alabama, Arkansas, Arizona, Colorado, Florida, Georgia, Maryland, Minnesota, North Carolina, New Jersey, South Carolina, Tennessee, Utah, Wisconsin, and West Virginia). We also excluded 2 ADDM surveillance years (2012 and 2014) because data on verbal status were not captured during these years.

Sociodemographic information collected included state of residence, sex (male, female), race and ethnicity (non-Hispanic Black [hereinafter, Black], non-Hispanic White [hereinafter, White], non-Hispanic Asian/Native Hawaiian/Other Pacific Islander [hereinafter, API], non-Hispanic American Indian/Alaska Native [hereinafter, AI/AN], or Hispanic), and highest maternal educational attainment (high school diploma, bachelor's degree, or >bachelor's degree). We also included census tract—level socioeconomic indicator rank values: (1) education: percentage of the population aged 25 years with high school diploma or General Educational Development (GED), (2) poverty: percentage of families with children aged <19 years living below the federal poverty level, and (3) median annual household income: percentile rank for census tract. Birth characteristics collected included birthweight <2500 g (low birthweight; yes or no) and gestational age <37 weeks (preterm; yes or no). Clinical and educational characteristics collected included cognitive functioning scores, adaptive functioning composite scores, verbal status, self-injurious

behavior, aggressive behavior, staring spells or seizure-like activity, and special education eligibility category.

### **Profound Autism Criteria**

We categorized children as having profound autism if they were either nonverbal or minimally verbal or had an IQ <50.6 We considered children to be nonverbal or minimally verbal if any of the following were identified in the records: (1) most recent evaluation at 48 months of age describing a child as nonverbal (median [IQR], 79 [65–93] months) or child determined to be nonverbal (no spontaneous words or word approximations) by clinician record review, (2) language classified primarily as echolalia or jargon by clinician review, or (3) being administered an Autism Diagnostic Observation Scale Module 1 (a gold standard observational measure appropriate for nonverbal or minimally verbal children) at age 48 months (median [IQR], 60 [53–70] months). <sup>11</sup> A sensitivity analysis increasing the age cutoff to 60 months minimally lowered (by <1 percentage point) the overall percentage of children with profound autism; thus, we kept the 48 months cutoff to capture children who may not have had a more recent language assessment recorded. If none of these indicators were identified, we categorized the child as being verbal.

We categorized children as having an IQ <50 if the most recent standard score results of an IQ test for either the overall IQ score or IQ subdomain score was <50. IQ data were not available for 19% (n = 3894) of children and were imputed.

### Statistical Analyses

To impute missing data, we used the multivariate imputation by chained equations approach with 20 imputed datasets. <sup>12</sup> We used a separate imputation model for each variable: logistic regression for binary data (self-injurious behavior, aggressive behavior, IQ <50, preterm birth, and low birth-weight) and polytomous logistic regression for categorical data (census-tract socioeconomic level, maternal education level, and adaptive score level). We created a descriptive summary of the percentage of the population with profound autism and prevalence ratios (PRs) among autism cases and 95% CIs comparing characteristics of children with profound autism with children with non–profound autism on the complete case (ie, nonimputed) and multiply imputed (imputed) datasets. We also calculated the median and IQR of age of first autism diagnosis among children with an age of diagnosis (age of diagnosis was not included in the imputation models because some children who met the ADDM autism case definition never had a documented clinical diagnosis) for the profound autism and non–profound autism groups. We further described characteristics of the profound autism and non–profound autism populations in the multiply imputed dataset.

For the 2016 surveillance year, we calculated the overall, sex-specific, race and ethnicity—specific, and site-specific population prevalence estimates of children with profound and non–profound autism per 1000 children aged 8 years and associated PRs to compare prevalence estimates between groups within the profound and non–profound autism populations, respectively. For study years 2000–2010, we calculated the overall population prevalence estimates by year to examine changes in the prevalence of profound and non–

profound autism over time. We used Stata release 17 (StataCorp LLC) and R version 4.1.2 (R Foundation for Statistical Computing) for data management and analyses.

## Results

The final sample included 20 135 children aged 8 years identified as having autism across 7 surveillance years and 15 sites (eTable 1 in Supplemental Material). Of these children, 26.7% (Table 1) had profound autism in the imputed data compared with 29.4% of children in the nonimputed data (eTable 2 in Supplemental Material); we imputed the IQ level of approximately 8% of all cases of profound autism.

The prevalence of profound autism in 2016 was 4.59 (95% CI, 4.31–4.85) cases per 1000 8-year-old children (Figure 1 and Table 2) with a ratio of 3.1 non–profound autism cases for every 1 profound autism case. Prevalence estimates of profound autism per 1000 8-year-old children were highest among API (6.05; 95% CI, 4.75–7.34) and Black (6.00; 95% CI, 5.32–6.69) children and lowest among White children (3.65; 95% CI, 3.31–3.98). In contrast, White children had the highest prevalence of non–profound autism (15.34; 95% CI, 14.67–16.01) compared with all other racial and ethnic groups. The prevalence of profound autism varied by site, ranging from 2.76 (95% CI, 2.23–3.28) in Colorado to 6.37 (95% CI, 5.23–7.51) in North Carolina. New Jersey had the highest prevalence of non–profound autism (25.99; 95% CI, 24.26–27.72).

During 2000–2016, the prevalence of non–profound autism increased from 3.94 to 14.26 cases per 1000 children aged 8 years; the prevalence of profound autism increased from 2.68 to 4.59 cases per 1000 children aged 8 years (Figure 2).

Profound autism was more common among girls than among boys (31.9% vs 25.6%; PR = 1.25; 95% CI, 1.18–1.32) (Table 1). Profound autism was also more common among Black (PR = 1.76; 95% CI, 1.67–1.86), API (PR = 1.55; 95% CI, 1.38–1.73), and Hispanic (PR = 1.50; 95% CI, 1.40–1.61) children than among White children. Apart from study year 2000, when the highest percentage of children with profound autism was observed (40.5%), the percentage of children with autism who had profound autism was similar across study years (2002–2016 range, 24.3%–27.9%). The lowest percentage of children with profound autism was observed in the most recently included study year, 2016 (24.3%). Sites varied in the percentage of their autism population who had profound autism, ranging from 21.0% in Colorado to 38.3% in South Carolina. Profound autism was more common among children who had documented self-injurious behaviors (PR = 1.48; 95% CI, 1.41–1.55) or seizure-like behaviors (PR = 1.33; 95% CI, 1.26–1.40) than among children without these behaviors. A higher percentage of children with an adaptive score 70 had profound autism (38.8%) than children with scores of 71–85 (14.2%) and >85 (8.0%). Lower communitylevel socioeconomic status and maternal education levels were associated with a higher percentage of profound autism. Children with autism who were born preterm or had a low birthweight were more likely to have profound autism than children with autism who were not born preterm (PR = 1.12; 95% CI, 1.02-1.22) or with a low birthweight (PR = 1.30; 95% CI, 1.20–1.41). The percentage of children with profound autism being served in an autism special education eligibility category was 30.2%. Among children with a known age

of diagnosis (74.2% for profound autism and 65.5% for non–profound autism), the median (IQR) age of diagnosis was 46 (34–62) months among children with profound autism and 57 (38–76) months among children with non–profound autism.

Among children with profound autism, most met the case definition of profound autism through only 1 of the criteria: 42.7% had met only the IQ score <50 criterion and 38.1% met only the nonverbal/minimally verbal criterion; 19.3% met both criteria (Table 3). Most children with profound autism had aggressive behavior (52.7%) and an adaptive functioning score 70 (79.4%); 36.5% had self-injurious behavior, and 31.0% had seizures or seizure-like behaviors.

### **Discussion**

Among the 20 135 children aged 8 years with autism in the 2000–2016 surveillance years, approximately 27% overall met the case definition of profound autism. In 2016, the most recent surveillance year, the prevalence of profound autism was 1 in 218 children aged 8 years. To our knowledge, this study is the first US population-based study quantifying the prevalence and describing the characteristics of the population with profound autism. This analysis used data from a long-running surveillance system for autism in the United States, allowing profound autism to be quantified in a well-described cohort of schoolaged children. It is important to identify these children because they are underrepresented in autism research and intervention studies and generally have the greatest need for services and supports. <sup>13,14</sup> Children with profound autism may have considerable medical complexity and likely may not be able to live independently or perform tasks of daily living as they age. <sup>15,16</sup> It is essential that policies, programs, and resources are tailored to the profound autism population across the life span to ensure their needs are met. <sup>17,18</sup>

While the prevalence of both profound and non–profound autism increased over time (2000–2016), the increase was greater for non–profound autism (from 1 in 254 to 1 in 70 children aged 8 years) than for profound autism (from 1 in 373 to 1 in 218 children aged 8 years); while attenuated, this trend remained when we used the 2002 study year as the starting reference point. This finding suggests that the composition of the identified autism population changed as the overall identified autism prevalence increased. Because the profound autism population has lower proportional representation among all autism cases compared with earlier years, it is critical to support policy and programming that are focused on the unique support needs of people with profound autism. 19–21

Variation across sites in the prevalence of autism per 1000 children aged 8 years in 2016 was greater for non–profound autism (site range, 10.4–26.0) than for profound autism (site range, 2.8–6.4). The difference in autism prevalence estimates by site within a study year could reflect geographic differences in identification and access to services, especially given the wide range of prevalence estimates of non–profound autism.

The overall Black to White PR of 1.0 in 2016 from a previously published report masks differences in prevalence by profound autism status. The prevalence of profound autism among Black children was higher than among White children, while the prevalence of non—

profound autism was higher among White children than among Black children. We found similar prevalence differences for the other racial and ethnic minority groups compared with White children. One possibility is that children of racial and ethnic minority groups with non–profound autism are less likely than White children with non–profound autism to be identified in the community.<sup>22</sup> The higher prevalence of profound autism in these racial and ethnic minority populations may also be due, in part, to a higher overall prevalence of intellectual disability compared with White children.<sup>23</sup> Additional research is needed to better understand the relationship of race and ethnicity and associated structural discrimination with profound autism status.

While boys were significantly more likely than girls to have autism, among autism cases, girls had 1.25 times the prevalence of profound autism as boys. This finding is consistent with preliminary characterization of the profound autism population. One explanation for this finding is that girls with non–profound autism may be underidentified because they compensate more than boys for social deficits. Alternatively, clinicians may be less likely to identify autism in girls than in boys because of differences by sex or gender in diagnostic procedures. Genetic analyses have also shown a higher frequency of deleterious mutations among girls with autism compared with boys with autism, which could contribute to a higher prevalence of profound autism among girls than among boys.

The finding that self-injurious behavior was associated with profound autism is consistent with previous studies.<sup>27</sup> One explanation may be that this behavior is a form of communication when cognitive and communication functions are impaired. Aggression was not associated with profound autism, in contrast to previous findings that showed a relationship between aggression and lower cognitive and language measures.<sup>28</sup> Further research is needed to better understand clinical and behavioral characteristics among those with profound autism.

### Limitations

This study had several limitations. First, ADDM Network sites changed across study years and may not be representative of their entire state or the United States, although they are representative of large, geographically defined populations and less subject to selection bias than clinical samples. Second, data on nonverbal status were not captured during the 2012 and 2014 study years, resulting in a gap in our longitudinal analytic data. Third, we used 48 months as a lower bound to assess nonverbal/minimally verbal status to allow time for verbal ability to develop, while balancing inclusion of an adequate number of years to capture more recent evaluations of verbal ability in surveillance records; children who were classified as nonverbal/minimally verbal may have become verbal at later ages.<sup>29</sup> In contrast, we classified children as verbal if they had no verbal ability information in their records, which might have led to underascertainment of nonverbal/minimally verbal status; furthermore, the ADDM case definition for nonverbal was stricter than the case definition for minimally verbal from other research.<sup>30,31</sup> As is true for data from the entire ADDM Network, it may be harder to distinguish the presence of autism for children with lower IO levels than for children with average or above-average IO levels.<sup>32</sup> Next, 3 of the socioeconomic variables were based on census tract-level data versus family-level data and,

thus, may not reflect the status of a child's family; however, similar findings were observed for the single family-level socioeconomic variable (maternal education status). Lastly, the definition of profound autism originally defined by the Lancet Commission is relatively new, and additional studies are needed to further characterize this population and understand how profiles can inform decision-making for supports and services in addition to prognostic utility; with further exploration, the definition of profound autism may evolve.

### **Conclusions**

Overall, more than one-quarter of 8-year-old children with autism met the criteria for profound autism. As the population of children with autism continues to change, describing and quantifying the profound autism population is critical for planning to ensure this group receives the services and supports needed during the life course and is appropriately represented in research and intervention studies. Further research on the barriers children with profound autism may face as they transition to adolescence and to adulthood will be important to ensure they are included in overall ASD services planning.

# **Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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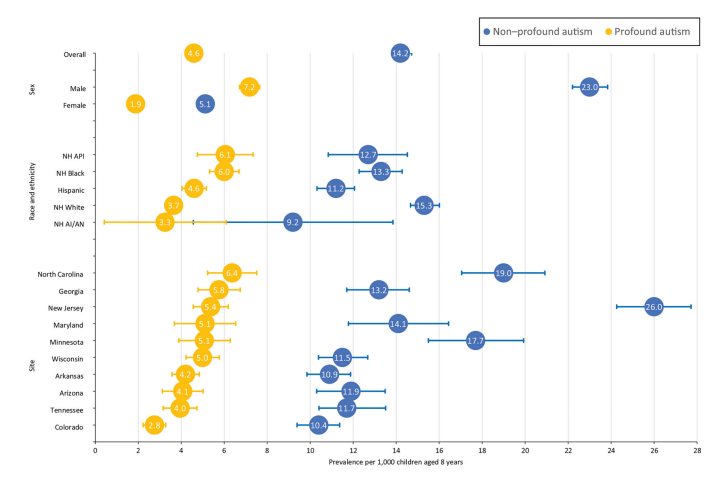


Figure 1.

Prevalence of profound and non–profound autism among children aged 8 years, by sex, race and ethnicity, and site, Autism and Developmental Disabilities Monitoring Network, 10 sites, United States, 2016. Error bars indicate 95% CIs. Data source: Autism and Developmental Disabilities Monitoring Network. Abbreviations: AI/AN, American Indian/Alaska Native; API, Asian/Native Hawaiian/Other Pacific Islander; NH, non-Hispanic.

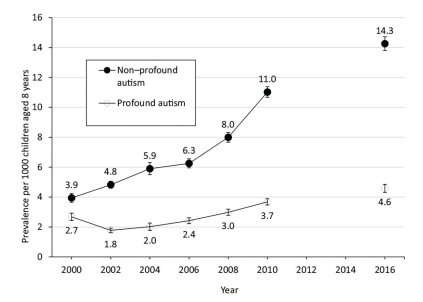


Figure 2.
Prevalence of profound and non–profound autism among children aged 8 years, by year,
Autism and Developmental Disabilities Monitoring Network, 15 sites, United States, 2016.
Data from 2012 and 2014 were unavailable and were excluded from analyses. Error bars indicate 95% CIs. Data source: Autism and Developmental Disabilities Monitoring Network.<sup>8</sup>

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Table 1.

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Characteristic	Total, $b$ no. (%)	Non-profound autism, $^{\mathcal{C}}$ % (n = 14 755)	Profound autism, <sup><math>d</math></sup> % (n = 5380)	Prevalence ratio (95% CI)
Total	20 135 (100.0)	73.3	26.7	
Sex				
Male	16 524 (82.1)	74.4	25.6	1 [Reference]
Female	3611 (17.9)	68.1	31.9	1.25 (1.18–1.32)
Race and ethnicity				
Non-Hispanic White	11 643 (57.8)	7.8.7	21.3	1 [Reference]
Non-Hispanic Asian/Native Hawaiian/Other Pacific Islander	764 (3.8)	67.1	32.9	1.55 (1.38–1.73)
Non-Hispanic American Indian/Alaska Native	110 (0.5)	71.8	28.2	1.33 (0.97–1.82)
Non-Hispanic Black	4312 (21.4)	62.6	37.4	1.76 (1.67–1.86)
Hispanic	2829 (14.1)	68.1	31.9	1.50 (1.40–1.61)
Non-Hispanic multiple races/other	479 (2.4)	77.6	22.4	1.05 (0.88–1.26)
ADDM study year				
2000	1246 (6.2)	59.5	40.5	1 [Reference]
2002	2366 (11.8)	73.1	26.9	0.67 (0.60–0.74)
2004	1151 (5.7)	74.5	25.5	0.63 (0.55-0.72)
2006	2286 (11.4)	72.1	27.9	0.69 (0.62–0.76)
2008	3216 (16.0)	72.8	27.2	0.67 (0.61–0.74)
2010	4975 (24.7)	75.0	25.0	0.62 (0.57–0.68)
2016	4895 (24.3)	75.7	24.3	0.60 (0.55-0.66)
Site				
Colorado	1130 (5.6)	79.0	21.0	1 [Reference]
Utah	683 (3.4)	78.7	21.3	1.01 (0.83–1.23)
Maryland	1722 (8.6)	9.77	22.1	1.05 (0.90–1.23)
Minnesota	313 (1.6)	7.77	22.3	1.06 (0.83–1.35)
New Jersey	2486 (12.3)	76.8	23.2	1.10 (0.96–1.27)
Arizona	2530 (12.6)	75.4	24.6	1.17 (1.02–1.34)
Tennessee	405 (2.0)	74.8	25.2	1.20 (0.97–1.48)
North Carolina	2209 (11.0)	72.8	27.2	1.30 (1.13–1.49)

Characteristic	Total, <sup>b</sup> no. (%)	Non-profound autism, <sup>c</sup> % $(n = 14.755)$	Profound autism, $d$ % (n = 5380)	Prevalence ratio (95% CI)
West Virginia	257 (1.3)	72.2	27.8	1.33 (1.03–1.70)
Wisconsin	1702 (8.5)	71.5	28.5	1.36 (1.17–1.57)
Arkansas	1514 (7.5)	70.5	29.5	1.41 (1.22–1.63)
Georgia	3306 (16.4)	70.2	29.8	1.42 (1.24–1.62)
Alabama	678 (3.4)	68.6	31.4	1.50 (1.26–1.77)
Florida	327 (1.6)	66.3	33.7	1.60 (1.31–1.97)
South Carolina	873 (4.3)	61.7	38.3	1.82 (1.57–2.12)
Associated behavioral and clinical features Self-injurious behaviors				
No	14 491 (72.0)	76.4	23.6	1 [Reference]
Yes	5645 (28.0)	65.2	34.8	1.48 (1.41–1.55)
Aggressive behaviors				
No	9722 (48.3)	73.8	26.2	1 [Reference]
Yes	10 413 (51.7)	72.8	27.2	1.04 (0.99–1.09)
Seizure or seizure-like behaviors				
No	15 058 (74.8)	75.3	24.7	1 [Reference]
Yes	5077 (25.2)	67.2	32.8	1.33 (1.26–1.40)
Adaptive functioning score				
70	11 015 (54.7)	61.2	38.8	1 [Reference]
71–85	6155 (30.6)	85.8	14.2	0.37 (0.34–0.40)
>85	2965 (14.7)	92.0	8.0	0.21 (0.18–0.24)
Socioeconomic status <sup>e</sup>				
Education, tertile				
Lowest	5259 (26.1)	0.99	34.0	1 [Reference]
Middle	7170 (35.6)	73.0	27.0	0.79 (0.75–0.84)
Highest	7705 (38.3)	78.5	21.5	0.63 (0.60–0.67)
Poverty, tertile				
Highest	5487 (27.3)	65.9	34.1	1 [Reference]
Middle	7071 (35.1)	73.1	26.9	0.79 (0.75–0.84)
Lowest	7578 (37.6)	78.8	21.2	0.62 (0.58–0.66)
Median annual household income, tertile				

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Characteristic	Total, b no. (%)	Non-profound autism, <sup><math>c</math></sup> % (n = 14 755)	Profound autism, $^d$ % (n = 5380)	Prevalence ratio (95% CI)
Lowest	5365 (26.6)	65.8	34.2	1 [Reference]
Middle	6980 (34.7)	72.6	27.4	0.80 (0.76–0.85)
Highest	7790 (38.7)	79.1	20.9	0.61 (0.57–0.65)
Maternal education status				
High school diploma	8711 (43.3)	69.2	30.8	1 [Reference]
Bachelor's degree	9106 (45.2)	75.6	24.4	0.79 (0.75–0.84)
>Bachelor's degree	2319 (11.5)	7.67	20.3	0.66 (0.59–0.74)
Birth characteristics				
$Preterm^f$				
No	16 986 (84.4)	73.8	26.2	1 [Reference]
Yes	3149 (15.6)	70.7	29.3	1.12 (1.02–1.22)
Low birth weight $^{\mathcal{G}}$				
No	17 730 (88.1)	74.2	25.8	1 [Reference]
Yes	2406 (11.9)	66.5	33.5	1.30 (1.20–1.41)
Special education eligibility $h$				
Autism	8452 (42.0)	69.8	30.2	1 [Reference]
Developmental delay/preschool	634 (3.1)	74.1	25.9	0.86 (0.75–0.99)
Emotional disturbance	502 (2.5)	94.1	5.9	0.19 (0.14–0.28)
Hearing or visual impairments	35 (0.2)	63.4	36.6	1.21 (0.76–1.91)
Intellectual disability	1025 (5.1)	43.3	56.7	1.88 (1.76–2.00)
Multiple disabilities	478 (2.4)	50.6	49.4	1.63 (1.47–1.81)
Orthopedic impairments	71 (0.4)	79.0	21.0	0.69 (0.43–1.11)
Unknown/other <sup>j</sup>	6739 (33.5)	7.97	23.3	0.77 (0.73–0.82)
Other health impairments	1271 (6.3)	91.1	8.9	0.29 (0.25–0.35)
Specific learning disabilities	916 (4.5)	6.68	10.1	0.33 (0.27–0.41)
Traumatic brain injury	12 (0.1)	37.5	62.5	2.06 (1.30–3.28)

 $<sup>^{2}</sup>$ Data source: Autism and Developmental Disabilities Monitoring (ADDM) Network.  $^{8}$ 

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bSample sizes may not sum to 20 135 because of rounding the average number in each group for the 20 imputed datasets.

 $<sup>^{\</sup>mathcal{C}}$ Non-profound autism was defined as children with autism who did not meet the profound autism criteria.

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 $q_{\rm pro}^{\prime\prime}$  Profound autism was defined as children with autism who were either nonverbal or minimally verbal or had an (intelligence quotient) IQ <50.

e Socioeconomic status was based on census tract-level indicator rank values. Within each site, census tracts were ranked among all other census tracts in the study population and divided into tertiles, so an approximately equal proportion of the population was categorized into each tertile. The 3 socioeconomic indicators were (1) education: percentage of population aged 25 years with high school diploma or General Educational Development (GED); (2) poverty: percentage of families with children aged <19 years living below the federal poverty level; and (3) median annual household income: percentile rank for census tract.

fBorn at <37 weeks gestation.

 $^{\mathcal{G}}$ Born weighing <2500 g.

hecial education categories were based on the most recent special education eligibility classification; if a child was served under >1 eligibility, the child would be classified as having "multiple disabilities."

j Other/unknown means either Individualized Educational Program eligibility information was not collected or the eligibility listed was not one of the choices.

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Table 2.

Prevalence of profound and non-profound autism among children aged 8 years, by sex, race and ethnicity, and site (N = 4895), ADDM Network, 10 sites, United States, 2016<sup>a</sup>

	Profo	Profound autism	Non-pro	Non-profound autism
Characteristic	Prevalence $^b$ (95% CI)	Prevalence $^b$ (95% CI) Prevalence ratio (95% CI)	Prevalence <sup>b</sup> (95% CI)	Prevalence ratio (95% CI)
Overall	4.59 (4.31–4.85)		14.26 (13.80–14.72)	
Sex				
Female	1.88 (1.62–2.13)	1 [Reference]	5.12 (4.72–5.52)	1 [Reference]
Male	7.18 (6.71–7.65)	3.83 (3.30–4.44)	23.03 (22.21–23.84)	4.50 (4.13–4.90)
Race and ethnicity				
Non-Hispanic White	3.65 (3.31–3.98)	1 [Reference]	15.34 (14.67–16.01)	1 [Reference]
Non-Hispanic Asian/Native Hawaiian/Other	6.05 (4.75–7.34)	1.66 (1.31–2.09)	12.68 (10.84–14.52)	0.83 (0.71–0.96)
Pacific Islander				
Non-Hispanic American Indian/Alaska Native	3.25 (0.42–6.09)	0.89 (0.37–2.13)	9.20 (4.56–13.85)	0.60 (0.36–1.00)
Non-Hispanic Black	6.00 (5.32–6.69)	1.65 (1.42–1.91)	13.28 (12.28–14.28)	0.87 (0.79–0.94)
Hispanic	4.60 (4.04–5.17)	1.26 (1.08–1.47)	11.19 (10.32–12.06)	0.73 (0.67–0.80)
$\mathrm{Site}^\mathcal{C}$				
Colorado	2.76 (2.23–3.28)	1 [Reference]	10.38 (9.39–11.37)	1 [Reference]
Tennessee	3.95 (3.16-4.73)	1.43 (1.09–1.88)	11.73 (10.41–13.51)	1.13 (0.98–1.31)
Arizona	4.07 (3.12–5.02)	1.48 (1.09–1.99)	11.90 (10.30–13.49)	1.15 (0.97–1.35)
Arkansas	4.20 (3.57-4.84)	1.52 (1.20–1.94)	10.86 (9.85–11.88)	1.05 (0.92–1.20)
Wisconsin	4.99 (4.22–5.77)	1.81 (1.42–2.31)	11.53 (10.39–12.68)	1.11 (0.97–1.27)
Minnesota	5.08 (3.89–6.28)	1.84 (1.36–2.50)	17.72 (15.50–19.93)	1.71 (1.46–2.00)
Maryland	5.11 (3.68–6.54)	1.85 (1.32–2.60)	14.10 (11.78–16.44)	1.36 (1.12–1.65)
New Jersey	5.37 (4.56–6.19)	1.95 (1.53–2.48)	25.99 (24.26–27.72)	2.50 (2.23–2.81)
Georgia	5.76 (4.78–6.74)	2.09 (1.62–2.69)	13.15 (11.70–14.61)	1.27 (1.10–1.47)
North Carolina	6.37 (5.23–7.51)	2.31 (1.78–3.00)	18.98 (17.05–20.92)	1.83 (1.59–2.10)

<sup>&</sup>lt;sup>a</sup>Profound autism was defined as children with autism who were either nonverbal or minimally verbal or had an (intelligence quotient) IQ <50. Non-profound autism was defined as children with autism who did not meet the profound autism criteria. Data source: Autism and Developmental Disabilities Monitoring (ADDM) Network.<sup>8</sup>

 $<sup>^{</sup>b}$  Prevalence per 1000 children aged 8 years.

 $^{\mathcal{C}}$  Five sites did not participate and contribute data to the 2016 surveillance year and were excluded.

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Table 3.

Characteristics of children aged 8 years with profound autism and non-profound autism among children with autism spectrum disorder (N = 20 135), ADDM Network, 15 sites, United States, 2000-2016<sup>a</sup>

Characteristic	Non–profound autism, no. $^{b}$ (%) (n = 14 755) Profound autism, no. $^{c}$ (%) (n = 5380)	Profound autism, no. <sup>c</sup> (%) $(n = 5380)$
Profound autism case criteria		
Intelligence quotient (IQ) <50 only	I	2295 (42.7)
Nonverbal/minimally verbal only	I	2049 (38.1)
IQ <50 AND nonverbal/minimally verbal	I	1036 (19.3)
Associated behavioral and clinical features Self-injurious behaviors		
No	11 077 (75.1)	3414 (63.5)
Yes	3678 (24.9)	1966 (36.5)
Aggressive behaviors		
No	7178 (48.6)	2546 (47.3)
Yes	7577 (51.4)	2834 (52.7)
Seizure or seizure-like behaviors		
No	11 343 (76.9)	3715 (69.1)
Yes	3412 (23.1)	1665 (31.0)
Adaptive functioning $score^d$		
70	6745 (45.7)	4271 (79.4)
71–85	5283 (35.8)	871 (16.2)
>85	2727 (18.5)	237 (4.4)

apofound autism was defined as children with autism who were either nonverbal or minimally verbal or had an intelligence quotient (IQ) <50. Non-profound autism was defined as children with autism who did not meet the profound autism criteria. Data source: Autism and Developmental Disabilities Monitoring (ADDM) Network.<sup>8</sup>

b Sample sizes may not sum to 14 755 because of rounding the average number in each group for the 20 imputed datasets.

 $<sup>^{\</sup>mathcal{C}}$ Sample sizes may not sum to 5380 because of rounding the average number in each group for the 20 imputed datasets.

dAmong those who met the definition of profound autism by verbal status alone (ie, IQ >50), 64% had an adaptive score of 70, 27% had an adaptive score of 71–85, and 9% had an adaptive score of >85.