

IN THE
Supreme Court of the United States

AMERICAN HOME PRODUCTS CORP. D/B/A WYETH, *et al.*,
Petitioners,

v.

MARCELO A. FERRARI AND CAROLYN H. FERRARI,
Individually and as Parents and Next Friend of
STERAN R. FERRARI,
Respondents.

On Petition for a Writ of Certiorari to the
Supreme Court of Georgia

BRIEF AMICI CURIAE OF THE
AMERICAN ACADEMY OF PEDIATRICS AND
10 OTHER PHYSICIAN AND PUBLIC HEALTH
ORGANIZATIONS IN SUPPORT OF
PETITIONERS

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**STATEMENT OF INTEREST OF
AMICI CURIAE**

Founded in 1930, amicus curiae the American Academy of Pediatrics (“AAP”) is a national, not-for-profit organization dedicated to furthering the interests of children’s health.¹ Since AAP’s inception, its

¹ Pursuant to Sup. Ct. R. 37.2(a), amici curiae note that counsel of record for all parties received notice at least 10 days prior to the due date of AAP’s intention to file this brief. Pursuant to Sup. Ct. R. 37.6, amici note that no part of this brief was authored by counsel for any party. Amici also note

membership has grown from 60 pediatricians to over 60,000 primary care pediatricians, pediatric medical subspecialists, and pediatric surgical specialists. Over the past 79 years, AAP has become a powerful voice for children's health through education, research, advocacy, and the provision of expert advice. AAP has worked with the federal and state governments, health care providers, and parents on behalf of America's children to ensure the availability of safe and effective childhood vaccines, the vast majority of which are administered in pediatricians' offices after careful consultation with parents.

Amicus curiae the AAP Section on Infectious Diseases was founded in 1990. It is comprised of AAP members who have a special interest in pediatric infectious diseases.

Amicus curiae the American Academy of Family Physicians ("AAFP") is a medical specialty society founded in 1947 to promote the science and art of family medicine. It is one of the largest medical organizations in the United States, with more than 94,000 members. The AAFP endorses the concept that all children and adults should have access to all immunizations recommended by the organization.

Amicus curiae the American College of Osteopathic Pediatricians ("ACOP") is the official pediatric organization of the American Osteopathic Association. ACOP's advocacy efforts represent the interests of all U.S. osteopathic pediatricians before Congress and

that no party or counsel made a monetary contribution intended to fund the preparation or submission of this brief, and that no person or entity other than amici or their members made such a monetary contribution. This brief is filed with the consent of all parties.

other governmental bodies as well as in coalition with other organizations that focus on children's welfare.

Amicus curiae the American Medical Association ("AMA"), an Illinois non-profit corporation, is a national association of approximately 240,000 physicians, residents, and medical students. The AMA is the largest medical society in the United States. Its members practice in every state and in every field of medical specialization, including pediatrics. The AMA was founded in 1847 to promote the science and art of medicine and the betterment of public health, and these remain its core purposes. The AMA has long been a vocal advocate of the importance of vaccines in maintaining high standards of public health in the United States.

Founded in 1872, amicus curiae the American Public Health Association ("APHA"), is the oldest and most diverse organization of public health professionals in the world. The association aims to protect all Americans and their communities from preventable, serious health threats and strives to assure that population-based health promotion and disease prevention activities and preventive health services are universally accessible in the United States. APHA represents a broad array of health providers, educators, environmentalists, policymakers, and health officers. APHA has a long-standing policy in support of safe and effective vaccines for children.

Amicus curiae Every Child By Two ("ECBT") is a non-profit health advocacy organization based in the United States and dedicated to protecting children from disease through promotion of vaccinations and raising parental awareness of potential vaccine

benefits. ECBT was founded in 1991 by former First Lady of the United States Rosalynn Carter and former First Lady of Arkansas Betty Bumpers.

Amicus curiae the Immunization Action Coalition (“IAC”) is a non-profit organization that works to increase immunization rates and prevent disease by creating and distributing educational materials for health professionals and the public that enhance the delivery of safe and effective immunization services. IAC also facilitates communication about the safety, efficacy, and use of vaccines within the broad immunization community of patients, parents, healthcare organizations, and government health agencies.

Amicus curiae the Infectious Diseases Society of America (“IDSA”) represents more than 8,500 infectious disease physicians and scientists devoted to patient care, research, prevention, and public health. IDSA’s purpose is to improve the health of individuals, communities, and society by promoting excellence in patient care, education, research, public health, and prevention relating to infectious diseases. Its members care for patients of all ages with serious and life-threatening infections, including vaccine-preventable diseases.

Amicus curiae the Pediatric Infectious Diseases Society (“PIDS”) is the world’s largest organization of professionals dedicated to the treatment, control, and eradication of infectious diseases affecting children. PIDS’s mission is to enhance the health of infants, children, and adolescents by promoting excellence in diagnosis, management, and understanding of infectious diseases through clinical care, education, research, and advocacy.

Amicus curiae the Vaccine Education Center at the Children’s Hospital of Philadelphia was launched in 2000 to provide accurate, comprehensive, and up-to-date information about vaccines and the diseases they prevent to parents and healthcare professionals. The Center communicates facts about vaccines, including how vaccines are made, how and why vaccines work, who recommends them, whether they are safe, whether they are still necessary, and when they should be administered to patients.

Amici—all of whom support the routine vaccination of children against a host of vaccine-preventable infectious diseases—urge this Court to grant the petition and reverse the judgment of the Supreme Court of Georgia below. As explained below, Congress enacted the National Childhood Vaccine Injury Act (“Vaccine Act”), 42 U.S.C. §§ 300aa-1 *et seq.* to avert a public health crisis and thus safeguard the Nation’s vaccine supply. As the Third Circuit recently recognized, Congress achieved that objective in part by expressly preempting “*all* design defect claims, including those based in negligence.” *Brusewitz v. Wyeth Inc.*, ___ F.3d ___, 2009 WL 792468, at *13 (3d Cir. Mar. 27, 2009) (emphasis added).

The Supreme Court of Georgia below reached the exact opposite conclusion, holding that design defect claims are preempted only “if it is determined, on a case-by-case basis, that the particular vaccine was unavoidably unsafe.” Pet. App. 15-16 (emphasis added). That decision—which allows judges *and juries* to decide whether a particular vaccine can be made safer—threatens a resurgence of “the very problems which led to instability in the vaccine market and which caused Congress to intervene

through the passage of the Vaccine Act” in the first place. *Bruesewitz*, 2009 WL 792468, at *13. The petition thus presents an issue of exceptional importance that warrants this Court’s review.

SUMMARY OF ARGUMENT

The public health benefits of childhood vaccines cannot be overstated. Because of vaccines, a number of debilitating and life-threatening infectious diseases have been eliminated or virtually eliminated in this country, providing not only significant savings in direct and indirect costs, but also enhancing the length and quality of life of countless children. It is no wonder that Congress has declared that “[t]he availability and use of vaccines to prevent childhood diseases is among the Nation’s top public health priorities.” H.R. Rep. No. 99-908, at 5 (1986).

In the mid-1980s, the number of vaccine-related lawsuits filed against vaccine manufacturers rose sharply. Although the tort system failed to provide adequate compensation for many children injured by vaccines, the flood of vaccine-related litigation overwhelmed vaccine manufacturers. A genuine threat to the public health emerged as manufacturers abandoned or considered abandoning the vaccine market. As the then-President of the AAP testified: “The threat to our vaccine supply in this country is a real one * * *. We could lose the remainder of our suppliers unless some positive legislative action is taken.” *National Childhood Vaccine Injury Compensation Act of 1985: Hearing Before the S. Comm. on Labor and Human Resources*, 99th Cong. 8 (Dec. 9, 1985) (hereinafter “Dec. 9, 1985 Hearing”) (statement of Martin Smith, M.D., President of the AAP).

Congress responded by passing the Vaccine Act. The Act established a no-fault alternative compensation program intended to provide adequate compensation to children injured by vaccines and to ensure the stability of the vaccine market and thus safeguard the Nation’s vaccine supply. As the Third Circuit recently recognized, the Act furthers that latter objective in part by expressly preempting “all design defect claims, including those based in negligence.” *Bruesewitz*, 2009 WL 792468, at *13 (emphasis added).

In the decision below, the Supreme Court of Georgia reached a contrary conclusion, holding that design defect claims are preempted only “if it is determined, *on a case-by-case basis*, that the particular vaccine was unavoidably unsafe.” Pet. App. 15-16 (emphasis added). As the Third Circuit explained in expressly rejecting that conclusion, if the Vaccine Act is interpreted “to allow case-by-case analysis of whether particular vaccine side effects are avoidable,” then “every design defect claim is subject to evaluation by a court.” *Bruesewitz*, 2009 WL 792468, at *29 (emphasis added). Thus, if allowed to stand, the decision below could precipitate the same crisis that Congress sought to avert in passing the Vaccine Act: “the very real possibility of vaccine shortages, and, in turn an increasing number of unimmunized children, and, perhaps, a resurgence of preventable diseases.” H.R. Rep. No. 99-908, at 7.

REASONS FOR GRANTING THE WRIT

I. VACCINE DEVELOPMENT IS ONE OF THE GREATEST PUBLIC HEALTH ACHIEVEMENTS OF THE TWENTIETH CENTURY.

The “[v]accination of children against deadly, disabling, but preventable infectious disease has been one of the most spectacularly effective public health initiatives this country has ever undertaken.” *Id.* at 4. See also Centers for Disease Control and Prevention, *Ten Greatest Public Health Achievements—United States, 1900-1999*, 48 MMWR 241 (Apr. 2, 1999) (listing vaccination as one of the ten greatest public health achievements of the twentieth century). Indeed, “the sharp and deep reduction in [infectious] diseases * * * is [largely] attributable to the development and employment of effective vaccines.” *Immunization and Preventive Medicine, 1982: Hearing Before the Subcomm. on Investigations and General Oversight of the S. Comm. on Labor and Human Resources*, 97th Cong. 103 (May 7, 1982) (hereinafter “1982 Hearing”) (statement of Vincent A. Fulginiti, Chairman, Committee on Infectious Diseases, AAP).

Because of vaccines, smallpox has been eradicated worldwide, Sandra W. Roush, *et al.*, *Historical Comparisons of Morbidity and Mortality for Vaccine-Preventable Diseases in the United States*, 298 JAMA 2155, 2160 (2007), and polio, diphtheria, and tetanus have essentially been eliminated in the United States. H.R. Rep. No. 99-908, at 5. In 2007, cases of measles, mumps, rubella, and pertussis (whooping cough) were reduced by more than 90% of twentieth century baseline levels. American Academy of

Pediatrics, *Red Book: 2009 Report of the Committee on Infectious Diseases* 2 (forthcoming 28th ed. 2009).

The significance of these developments is beyond dispute: “[C]hildren in the United States enjoy substantial freedom from the ravages of once common communicable infectious diseases and illnesses. These illnesses limited life expectancy and left tens of thousands disabled in their wake.” *1982 Hearing, supra*, at 103 (statement of Vincent A. Fulginiti, Chairman, Committee on Infectious Diseases, AAP). Vaccines have enhanced not only the length and quality of life of countless children who have been vaccinated but also that of others in the community who are unable to be vaccinated but whose risk of exposure to an infectious disease has been reduced accordingly. Institute of Medicine, *Financing Vaccines in the 21st Century: Assuring Access and Availability* 27 (2004). It has been estimated that vaccination with just seven of the routinely recommended childhood vaccines “prevents an estimated 33,000 deaths and 14 million cases of disease in every birth cohort.” Roush, *et al.*, *supra*, at 2160.

Vaccines have also translated into direct savings in medical costs, as well as increased productivity from families that would otherwise be burdened by disease. Institute of Medicine, *supra*, at 27-29. It has been estimated that for every dollar invested in childhood vaccination against nine vaccine-preventable diseases, \$5.80 is saved in direct medical costs; \$17.70 is saved when indirect benefits, such as lost productivity, are taken into account. Walter A. Orenstein, *et al.*, *Immunizations in the United States: Success, Structure, and Stress*, 24 Health Affairs 599, 600 (2005). Overall, “[b]illions of medi-

cal and health-related dollars have been saved by immunizations." H.R. Rep. No. 99-908, at 4.

Of course, "[n]o vaccine is completely safe or effective." Centers for Disease Control and Prevention, *General Recommendations on Immunization: Recommendations of the Advisory Committee on Immunization Practices (ACIP)*, 55 MMWR 1 (Dec. 1, 2006). See H.R. Rep. No. 99-908, at 6 ("There is today no 'perfect' or reaction-free childhood vaccine on the market."). Even when vaccines are properly manufactured, distributed, and administered, a small number of children may suffer rare but serious adverse reactions. See H.R. Rep. No. 99-908, at 4, 6. "Despite these possibilities, public health officials, private physician groups, and parent organizations have repeatedly stated that it is safer to take the required shots than to risk the health consequences of contracting the diseases immunizations are designed to prevent." *Id.* at 6. In other words, the enormous benefits of vaccination vastly outweigh the small risk of injury. See *General Recommendations on Immunization, supra*, at 1 ("[R]ecommendations for vaccination practices balance scientific evidence of benefits for each person and to society against the potential costs and risks for vaccination for the individual and programs."); H.R. Rep. No. 99-908, at 6 ("in light of the overall success of immunization programs, the Federal government continues to support * * * immunizations to children").

II. CONGRESS ENACTED THE VACCINE ACT TO PROVIDE ADEQUATE COMPENSATION TO CHILDREN INJURED BY VACCINES AND TO SAFEGUARD THE NATION'S VACCINE SUPPLY.

In 1986, the Nation faced a public health crisis. Vaccine-related lawsuits against vaccine manufacturers had spiked, and rising litigation and insurance costs threatened to halt vaccine production in the United States. H.R. Rep. No. 99-908, at 4, 6-7. At the same time, however, the tort system had failed to provide adequate compensation for children injured by vaccines. *Id.* at 6. Congress responded by enacting the Vaccine Act, thereby ensuring adequate compensation for children injured by vaccines and safeguarding the Nation's vaccine supply.

A. The Costs Of Vaccine-Related Litigation Had Threatened To Halt Vaccine Production In The United States.

In the mid-1980s, the number of vaccine-related suits filed against vaccine manufacturers increased markedly. *Id.* at 4. According to a 1985 survey of the seven manufacturers producing childhood vaccines,² between January 1980 and March 1985, 299 lawsuits were filed against them seeking compensation for vaccine-related injuries; 84 percent of those suits were related to childhood vaccines. Staff of H. Subcomm. on Health and the Environment of the Comm. on Energy and Commerce, 99th Cong., *Childhood Immunizations* 86 (Comm. Print 1986) (hereinafter

² Two of the manufacturers were operated by State organizations in Michigan and Massachusetts. H.R. Rep. No. 99-908, at 7.

"*Childhood Immunizations*"). About 60 percent of all the suits filed sought damages in the aggregate of \$3.5 billion. *Id.* Between 1983 and 1984 alone, litigation costs nearly doubled—jumping from \$4.7 million to \$9.8 million. *Id.* at 87.

With the deluge of lawsuits, vaccine manufacturers faced rising insurance premiums and a decreasing pool of insurers willing to cover them. H.R. Rep. No. 99-908, at 6-7; *Childhood Immunizations, supra*, at 73; *National Childhood Vaccine-Injury Compensation Act: Hearing Before the S. Committee on Labor and Human Resources*, 99th Cong. 288 (July 18, 1985) (hereinafter "*July 18, 1985 Hearing*") (statement of Stephen White, Vice President of Reed-Stenhouse, Ltd.) (explaining that insurance companies were struggling with losses from pharmaceutical, asbestos, and pollution claims). As litigation and insurance costs soared, "the prices of vaccines * * * jumped enormously," H.R. Rep. No. 99-908, at 4—in some cases as much as 900 percent, *see Childhood Immunizations, supra*, at 90—and "[t]he number of childhood vaccine manufacturers * * * declined significantly." H. Rep. No. 99-908, at 4.

As it became increasingly clear that vaccine prices could not forever keep pace with escalating litigation and insurance costs, *see, e.g., July 18, 1985 Hearing, supra*, at 240 (statement of Robert Johnson, President of Lederle Laboratories) (noting that "vaccine pricing" of the previous year would not "cover the projected costs of liability" for the following year), the few remaining vaccine manufacturers began "to question their continued participation in the vaccine market." H.R. Rep. No. 99-908, at 7. *See July 18, 1985 Hearing, supra*, at 256 (statement of Robert

Johnson, President of Lederle Laboratories) ("If the current trend of spiraling litigation continues or worsens, there * * * is a very real possibility that we will be forced to abandon the vaccine business."); *id.* at 284 (statement of David J. Williams, Vice President and General Manager of Connaught Laboratories, Inc.) ("There is always a possibility that Connaught will be unable to remain in the vaccine business.").

As Congress recognized, "[t]he loss of any of the existing manufacturers of childhood vaccines * * * could create a genuine public health hazard in this country." H.R. Rep. No. 99-908, at 7. At that time, "there [was] only one manufacturer of the polio vaccine, one manufacturer of the measles, mumps, rubella (MMR) vaccine, and two manufacturers of the [diphtheria, pertussis, and tetanus] DPT vaccine." *Id.*³ Thus, as the then-President of the AAP testified: "The threat to our vaccine supply in this country is a real one * * *. We could lose the remainder of our suppliers unless some positive legislative action is taken." *Dec. 9, 1985 Hearing, supra*, at 8 (statement of Martin Smith, M.D., President of the AAP).

B. The Tort System Had Failed To Provide Adequate Compensation For Children Injured By Vaccines.

Ironically, while lawsuits against vaccine manufacturers skyrocketed, some children injured by vaccines failed to receive any compensation. Despite their injuries, some children were simply not deemed

³ Michigan and Massachusetts also produced their own DPT vaccine. H.R. Rep. No. 99-908, at 7.

good “candidates for litigation,” *National Childhood Vaccine-Injury Compensation Act: Hearing Before the S. Committee on Labor and Human Resources*, 98th Cong. 171 (May 3, 1984) (hereinafter “1984 Hearing”) (statement of Andrew Dodd, Attorney at Ward, Dodd & Grant, Torrance, California), because any prospective recovery was not “large enough” to make their cases attractive to an attorney. *Id.* at 146 (statement of Martin H. Smith, M.D., President-elect of the AAP).

Thus, while a few lawsuits reaped multi-million dollar awards, some injured children received no compensation at all. 132 Cong. Rec. H9943, H9952 (Oct. 14, 1986) (statement of Rep. Waxman); *id.* at H9954 (statement of Rep. Biaggi); 1984 *Hearing, supra*, at 4 (statement of Senator Kennedy) (tort system “awards few handsomely and sends others equally aggrieved away penniless”). The tort system was thus aptly described as a “lottery.” 1984 *Hearing, supra*, at 277 (statement of John E. Lyons, President of Merck Sharp & Dohme).

C. The Vaccine Act Provides Adequate Compensation To Children Injured By Vaccines And Ensures The Stability Of The Vaccine Market And The Nation's Vaccine Supply.

Congress responded to the looming crisis by enacting the Vaccine Act. The overriding goals of the Act were two-fold: (1) to ensure adequate compensation for children injured by vaccines, and (2) to stabilize the vaccine market and safeguard the Nation's vaccine supply. H.R. Rep. No. 99-908, at 7.

Congress addressed both of those goals in part by establishing the National Vaccine Injury Compensation Program (“VICP”), a no-fault alternative compensation system under which children injured by certain vaccines would receive “fair and expeditious” compensation for their injuries. *Id.* at 12. See 42 U.S.C. § 300aa-10 *et seq.* Under the VICP, a person seeking compensation for an injury caused by a vaccine covered by the Act must file a petition with the United States Court of Federal Claims, which refers the petition to a “Vaccine Court”—an office within the court of special masters appointed to four-year terms by the court to hear VICP claims. 42 U.S.C. §§ 300aa-11(a)(1)-(2), 300aa-12(c), 300aa-21(a). The Secretary of Health and Human Services is named as a respondent; vaccine manufacturers are not parties to VICP proceedings. *Id.* § 300aa-12(b)(1).

A petitioner is entitled to compensation if he or she has suffered an injury set forth in the “Vaccine Injury Table”—a table of vaccines and the injuries presumed to be caused by those vaccines—unless it can be shown by a preponderance of the evidence that the petitioner's injury was not caused by the vaccine. *Id.* §§ 300aa-11(b), (c), 300aa-13(a)(1), 300aa-14. A petitioner who has not suffered a “Table Injury” may still obtain compensation by proving that his or her injury was in fact caused by a vaccine covered by the Act. *Id.* § 300aa-11(c)(1)(C)(ii). See *Grant v. Secretary of HHS*, 956 F.2d 1144, 1147-48 (Fed. Cir. 1992).⁴ Payment of compensation is made

⁴ The special masters of the Vaccine Court have developed a proficiency in the complex medical and scientific issues involved in causation claims. Indeed, the Court of Federal Claims

from a “Vaccine Injury Compensation Trust Fund”—funded by a manufacturers excise tax on those vaccines covered by the Act, see 26 U.S.C. §§ 4131, 9510—on a no-fault basis. 42 U.S.C. §§ 300aa-13,

recently observed that “instead of being passive recipients of information, such as jurors, special masters are given an active role in determining the facts relevant to Vaccine Act petitions,” and that “special masters have the expertise and experience to know the type of information that is most probative of a claim.” *Doe v. Secretary, HHS*, 76 Fed. Cl. 328, 338-339 (2007).

The expertise of the special masters in evaluating causation claims was recently demonstrated in an Omnibus Autism Proceeding established under the VICP to determine whether there is a causal link between childhood vaccines and autism. Approximately 5,000 cases alleging an association between autism and either the MMR vaccine (which does not contain thimerosal) or vaccines containing the preservative thimerosal, or both, have been filed with the Vaccine Court. See <http://www.hrsa.gov/vaccinecompensation>. On February 12, 2009, three special masters issued voluminous opinions evaluating evidence based on the theory that the MMR vaccine, in combination with vaccines containing the preservative thimerosal, causes autism. See *Cedillo v. Secretary of HHS*, 2009 WL 331968 (Fed. Cl. Feb. 12, 2009); *Hazlehurst v. Secretary of HHS*, 2009 WL 332306 (Fed. Cl. Feb. 12, 2009); *Snyder v. Secretary of HHS*, 2009 WL 332044 (Fed. Cl. Feb. 12, 2009). All three cases rejected the proposition that the vaccines in question caused autism.

In reaching those conclusions, the special masters in each case considered a wealth of scientific evidence. As the special master in *Snyder* observed: “The evidentiary record in this case * * * encompasses, *inter alia*, nearly four weeks of testimony, including that offered in the *Cedillo* and *Hazlehurst* cases; over 900 medical and scientific journal articles; 50 expert reports (including several reports of witnesses who did not testify); supplemental expert reports filed by both parties post-hearing, [and] the testimony of fact witnesses on behalf of [the injured child and his] medical records.” *Snyder*, 2009 WL 332044, at *8.

300aa-14, 300aa-15(i). Since 1989, the Vaccine Court has issued more than 2,200 awards totaling over \$1.7 billion. See National Vaccine Injury Compensation Program, *Statistic Report* (Apr. 3, 2009), available at http://www.hrsa.gov/vaccinecompensation/statistics_report.htm.

After the Vaccine Court has issued a final judgment, a petitioner may accept or reject it. 42 U.S.C. § 300aa-21(a).⁵ Although a party who rejects the Vaccine Court’s judgment may pursue certain *limited* claims in state or federal court, design defect claims are not among them. *Id.* § 300aa-21(a), (b). As the Third Circuit recently recognized, Congress expressly preempted “all design defect claims, including those based in negligence.” *Brusewitz*, 2009 WL 792468, at *13 (emphasis added). See 42 U.S.C. § 300aa-22(b)(1). If an injured person has such a claim, he or she “should pursue recompense in the compensation system, not the tort system.” H.R. Rep. No. 99-908, at 26. The preemption of all design defect claims is critical to Congress’s objective of stabilizing the vaccine market and safeguarding the Nation’s vaccine supply. As the Third Circuit explained: “Congress[] believe[d] that an alternate compensation system would reduce awards and create a stable, predictable basis for estimating liability.” *Brusewitz*, 2009 WL 792468, at *12. Indeed, as the legislative history makes clear, Con-

⁵ The Vaccine Act also authorizes petitioners to “opt out” of a VICP proceeding if a special master has not resolved his or her petition within 240 days or if the Court of Federal Claims has not completed its review of a special master’s decision within 420 days of the date on which the petition was filed. See 49 T.T.C. § 300aa-22(f).

gress “believe[d] that once this system [was] in place and manufacturers ha[d] a better sense of their potential litigation obligations, a more stable childhood vaccine market [would] evolve.” H.R. Rep. No. 99-908, at 7.

III. THE DECISION BELOW—WHICH DIRECTLY CONFLICTS WITH A DECISION OF A FEDERAL COURT OF APPEALS—POSES A THREAT TO THE FUTURE PRODUCTION AND DEVELOPMENT OF VACCINES.

Contrary to all clear indications of congressional intent, the Supreme Court of Georgia held below that design defect claims are preempted only “if it is determined, *on a case-by-case basis*, that the particular vaccine was unavoidably unsafe.” Pet. App. 15-16 (emphasis added). As the Third Circuit recently concluded, that interpretation of the Vaccine Act is simply wrong. See *Brusewitz*, 2009 WL 792468, at *11. As the Third Circuit explained, if the Act is interpreted “to allow case-by-case analysis of whether particular vaccine side effects are avoidable,” then “*every* design defect claim is subject to evaluation by a court.” *Id.* at *12 (emphasis added).

If that were the case, “[e]ach of the objectives extolled [in the Vaccine Act’s legislative history] would be undermined.” *Id.* at 36. Thus, the decision below—which allows judges *and juries* to decide whether a particular vaccine can be made safer—threatens a resurgence of “the very problems which led to instability in the vaccine market and which caused Congress to intervene through the passage of the Vaccine Act” in the first place. *Id.* That threat is extremely palpable, as the recent decisions issued by the Vaccine Court promise to unleash a barrage of

claims in the courts. See *supra* at 16 n.4. If allowed to stand, therefore, the decision below could drive vaccine manufacturers from the market and halt the future production and development of childhood vaccines in this country.

A. Unpredictable Litigation Costs Could Once Again Force Vaccine Manufacturers To Abandon Or Consider Abandoning The Vaccine Market.

By eliminating the threat of most lawsuits, the Vaccine Act has prevented manufacturers from abandoning the vaccine market, thus ensuring a stable supply of vaccines. See Louis Z. Cooper, *et al.*, *Protecting Public Trust in Immunization*, 122 *Pediatrics* 1, 2 (2008). By allowing case-by-case consideration of whether vaccines are unavoidably unsafe, the decision below will “undoubtedly increase the costs and risks associated with litigation and [will] undermine a manufacturer’s efforts to estimate and control costs.” *Brusewitz*, 2009 WL 792468, at *13. Thus, the decision creates the “very real possibility” that vaccine manufacturers will once again abandon or be forced to consider abandoning the vaccine market. H.R. Rep. No. 99-908, at 7.

That is particularly so given the precarious state of the vaccine industry. Today, as in 1986, there continues to be only *one* manufacturer of the MMR vaccine, and only *two* manufacturers of the DTP vaccine. American Academy of Pediatrics, *Status of Licensure and Recommendations for New Vaccines*, Red Book Online (2009), available at <http://aapredbook.aappublications.org/news/vaccstatus.shtml>; *Childhood Immunizations*, *supra*, at 67; H.R. Rep. No. 99-908, at 7. And while the Vaccine

Act has been instrumental in preventing manufacturers from fleeing the vaccine market, the number of vaccine manufacturers has not greatly increased since the Act's passage. See *Status of Licensure and Recommendations for New Vaccines, supra*. The costs of developing and producing vaccines have also increased over the years. Between 1991 and 2003, costs for research and development, postlicensure clinical studies, and production process improvements grew from \$231 million to \$802 million. Stanley A. Plotkin, *et al.*, *Vaccines* 38 (2008).

Thus, vaccine manufacturers today are no better—and, indeed, are perhaps even more poorly—situated to handle the unpredictability and expense of litigation. Yet, as was true in 1986, “the withdrawal of even a single manufacturer would present the very real possibility of vaccine shortages, and, in turn an increasing number of unimmunized children, and, perhaps, a resurgence of preventable diseases.” H.R. Rep. No. 99-908, at 7.

B. The Progress That Has Been Made In Vaccine Development Since The Passage Of The Vaccine Act Could Come To A Halt.

In addition to ensuring the stability of the existing childhood vaccine market, one of Congress's objectives in passing the Vaccine Act was ensure “that a greater number of vaccine products will become available to prevent disease.” H.R. Rep. No. 99-908, at 4. In that regard, the Act has been unquestionably successful. Vaccine development has flourished since 1986 with the number of vaccine-preventable diseases having more than doubled. See *Childhood Immunizations, supra*, at 1; Centers for Disease

Control and Prevention, *Recommended Immunization Schedules for Persons Aged 0 Through 18 Years—United States*, 57 MMWR Q1-Q4 (Jan. 2, 2009).

In 1986, children were routinely vaccinated against seven diseases (diphtheria, measles, mumps, pertussis, poliomyelitis, rubella, and tetanus). *Childhood Immunizations, supra*, at 1. Today, children are also routinely immunized against an additional eight diseases: *haemophilus influenzae* type b (Hib), hepatitis A, hepatitis B, influenza, meningococcal disease, pneumococcal disease, rotavirus, and varicella (chicken pox). *Recommended Immunization Schedules for Persons Aged 0 Through 18 Years—United States, supra*, at Q1-Q4. Research and development of new vaccines is always ongoing. See Immunization Action Coalition, *Vaccine-Related Journal Articles, available at* <http://www.immunize.org/journal/articles/toipoten.asp> (listing, by year, published articles regarding vaccine development).

Vaccine manufacturers face many challenges in bringing new vaccines to market. See Paul A. Offit, *Why Are Pharmaceutical Companies Gradually Abandoning Vaccines?*, 24 Health Affairs 624, 623-629 (2005). In addition to research and development, vaccine manufacturers are also “almost exclusively” responsible for the production and distribution of such vaccines. See Orenstein, *et al.*, *supra*, at 601-603. As noted, by eliminating the threat of most lawsuits, the Vaccine Act has kept manufacturers from abandoning vaccine production. See Cooper, *et al.*, *supra*, at 2. If the decision below is allowed to stand, the prognosis for future vaccine development is extremely poor.

CONCLUSION

For the foregoing reasons, and those stated in the petition, the petition for a writ of certiorari should be granted, and the judgment below reversed.

Respectfully submitted,

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