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Prevention of Congenital Disorders and Care of Affected Children
A Consensus Statement

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As the Sustainable Development Goals are adopted by United Nations member states, children with congenital disorders remain left behind in policies, programs, research, and funding. Although this finding was recognized by the creation and endorsement of the 63rd World Health Assembly Resolution in 2010 calling on United Nations member states to strengthen prevention of congenital disorders and the improvement of care of those affected, there has been little to no action since then. The Sustainable Development Goals call for the global health and development community to focus first and foremost on the most vulnerable and those left behind in the Millennium Development Goal era. To maximize the opportunity for every woman and couple to have a healthy child and to reduce the mortality and severe disability associated with potentially avoidable congenital disorders and their consequences for the children affected, their families and communities, and national health care systems, we propose priority measures that should be taken urgently to address this issue.

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The ICBDs were established to focus attention on the need to improve the prevention and care of congenital disorders given that the proportion of deaths in children younger than 5 years due to congenital disorders is rising in many low- and middle-income countries. In addition, rates of disability resulting from congenital disorders are increasing in many countries where newborn survival is improving but quality health care is lagging. Congenital disorders and their associated disabilities result in a substantial emotional, social, and economic toll on affected individuals, their families, and the communities in which they live. The Dar es Salaam ICBD was assembled to agree on actions to reduce this toll, especially in vulnerable, marginalized populations. A draft consensus statement was presented and discussed by the audience on the last day of the conference, with major changes incorporated at that time. The final document was created by the authors and circulated to all conference participants, with 76 providing input and supporting the consensus statement. A working group was also established to explore additional ways to put this statement into action.

We support accelerating the prevention of congenital disorders and improvement of care of affected individuals, recognizing that:

• An estimated 7.9 million children are born each year with a genetic or partially genetic (multifactorial) congenital disorder, and several hundred thousand more are born with congenital disorders due to in utero insults after conception, such as infections, and exposure to teratogens, such as alcohol.2,3
• Of the 2.7 million newborns who die annually, more than 1 in 10 die of a congenital disorder, and, overall, there are an estimated 484,000 deaths due to congenital disorders among children younger than 5 years.4 This number is likely a gross underestimate, however, because many deaths due to congenital disorders, such as heart defects and metabolic disorders, go undetected.

• It is likely that more than 192,000 of the 2.6 million annual stillbirths may result from an underlying congenital disorder.5 The percentage of stillborn children affected by congenital disorders is likely much higher than for live births, and many efforts to prevent stillbirth will help reduce the occurrence of congenital disorders among stillborn children.

• Most newborns with a serious congenital disorder who survive face a lifetime of severe disability.2,3

• An estimated 94% of newborns with 1 or more congenital disorders are born in low- and middle-income countries, placing an additional severe burden on families, communities, and national health care systems.2,3

• Up to an estimated 70% of congenital disorders are preventable or their effect can be substantially mitigated and quality of life improved, but these preventive and mitigating actions are occurring almost exclusively in high-income settings.2,3

• The fact that children with congenital disorders have been left behind in policies, programs, research, and funding was recognized by the creation and endorsement of the 63rd World Health Assembly Resolution,6 calling in 2010 for United Nations member states to (1) raise awareness of congenital disorders as a cause of child morbidity and mortality; (2) develop and strengthen birth registration and surveillance for birth defects; (3) strengthen evidence on etiologic factors, diagnosis, and prevention of major birth defects; and (4) develop national plans for implementation of effective interventions to prevent and manage birth defects. This call by the World Health Organization, however, has gone unheeded except in pockets such as Southeast Asia, where a strategic framework7 is guiding efforts to prevent and control congenital disorders in 12 countries of the region.

The conference participants agreed that, with the advent of the SDGs, greater emphasis must be placed on more holistic approaches, including preventive care and, beyond survival, optimization of children’s developmental potential. This emphasis aligns with the SDGs’ call for equality in social inclusion and in opportunities for education, employment, and the ability for all human beings to fulfill their potential and enjoy prosperous, productive lives. Furthermore, the SDGs call for giving priority in policy and action to the most vulnerable, specifically including those with disabilities, and those currently most left behind.8 Congenital disorders can be considered the first chronic disease experienced in life and are encompassed in SDG goal 3 that calls for a reduction, by 2030, by one-third in premature mortality from noncommunicable diseases through prevention and treatment and the promotion of mental health and well-being.7

Call to Action
To maximize the opportunity for every woman and couple to have a healthy child; to reduce the consequences of potentially avoidable congenital disorders for those affected, their families, the health care system, and the wider society; and to promote the well-being of children who have a congenital disorder, there are many measures that should be taken urgently to address this issue. In this context and in order that no child is left behind, we pledge an initial focus that supports the following:

Improving data quality:
1. Building consensus on and widespread use of a standardized definition of congenital disorders, such as “abnormalities of structure or function, which are present from birth,”25 to facilitate data comparison and ensure that the contribution of congenital disorders to the burden of disease is comprehensively represented.

2. Establishing registries and surveillance systems and their integration, where possible, into existing data platforms to monitor the toll and risks of congenital disorders and evaluating the outcome of interventions for prevention and care. Consideration should also be given to the collection of pertinent data available from existing registries and surveillance systems in other countries.

Reducing risk:
1. Promoting family planning, allowing women and couples to choose when they have their first child, space their pregnancies, plan family size, define the ages at which they wish to complete their family, and reduce the proportion of unintended pregnancies.

2. Ensuring a healthy, balanced diet for girls and during a woman’s reproductive years through an adequate intake of macronutrients (protein, carbohydrates, and fats) and a broad range of micronutrients. Special attention should be given to adding 400 μg of synthetic folic acid daily to the diet through fortification and supplementation while also promoting a diet rich in food folates, correcting iodine deficiency through fortification, and ensuring iron sufficiency through fortification, supplementation, and therapy for those with deficiencies.

3. Removing teratogenic substances from the diet, the most important of which is alcohol, and minimizing environmental contaminants in foods.

4. Controlling infections in women of reproductive age, including rubella and syphilis, and optimizing maternal health through detection and management of chronic illnesses associated with an increased risk of congenital disorders, such as type 2 diabetes mellitus and epilepsy, which require teratogenic medications.

Improving care:
1. Training physicians, nurses, allied health care professionals, and workers in the fundamentals of the recognition, causes, and care of children with congenital disorders and ensuring physical examinations of all newborns by trained health care professionals before discharge from the hospital or clinic.

2. Aligning medical and social services to provide timely treatments for congenital disorders, including surgery, medications, dietary modifications, and rehabilitation services when needed.

3. Providing emotional and practical support for parents to enable them to understand and manage their risk of congenital disorders and to help families in supporting the growth and development of children with congenital disorders.

Empowering the public and civil society:
1. Educating the public about congenital disorders and the steps mothers and fathers can take with their health care professionals to maximize the chances of a healthy pregnancy.
2. Strengthening civil society—including patient and parent support groups, faith-based groups, and nongovernmental organizations—to advocate for improved prevention of congenital disorders and access to high-quality, family-centered patient care, including facilitating community and professional awareness and education and advocating for increased funding for research on the causes of congenital disorders.

The following additional actions should be taken by countries as priorities and circumstances allow:

1. Training physicians, nurses, and allied health care professionals in the essentials of medical genetics. This training should include diagnosis of common congenital disorders before and at birth; means of treatment where possible in the primary health care setting; knowing when to refer a patient for more specialized treatment; basic genetic counseling, including best practices in communicating unfavorable health information to parents; and support for families who have a child or are at risk of having a child with a congenital disorder. Genetic counseling aims to empower those who are counseled to make autonomous decisions regarding their health in ways that are consonant with their religious and ethical beliefs and circumstances and to support them in their decisions.

2. Establishing periconception medical services to assist women and their partners in attaining optimal physical and mental health and well-being and to facilitate a healthy pregnancy and delivery of a healthy infant. These services include rubella immunization; screening for the risk of genetic, partially genetic, and teratogenic congenital disorders; and mental health counseling, including identification and support for depression.

3. Implementing preconception or prenatal medical screening to identify women and couples at risk of having a baby with hemoglobin disorders; Down syndrome; blood type incompatibility; congenital sexually transmitted infections such as syphilis, human immunodeficiency virus, and herpes simplex virus; and structural malformations, particularly neural tube defects.

4. Establishing newborn screening to identify congenital hypothyroidism, phenylketonuria, galactosemia, sickle cell disease, glucose-6-phosphate dehydrogenase deficiency, and other metabolic disorders.

5. Supporting research into the diagnosis, prevention, etiologic factors, and treatment of congenital disorders to enable improved outcomes for children into the future.

The conference participants call for concerted action by international government policymakers and donor organizations to explore how these recommended actions can be funded through more cost-effective and rational integration of policy, funding, and interventions across the reproductive, maternal, newborn, child, and adolescent continuum. Systems are required that encourage more effective partnership among the many existing organizations and agencies whose missions address common risk factors and outcomes and who would benefit financially and programatically from better integration of policy, research, and action at international and national levels.

Conclusions

The Seventh ICBD in Dar es Salaam, Tanzania, was a pivotal opportunity to build consensus and commitment for accelerated prevention of congenital disorders and improvement of care of affected children in low- and middle-income countries. In conjunction with the newly launched SDGs and building on the 63rd World Health Assembly resolution calling on United Nations member states to strengthen prevention and care for congenital disorders, immediate action on the plan outlined above will save newborn and child lives, reduce disability rates and improve quality of life in survivors, and substantially lessen the current emotional and economic toll of these conditions on affected individuals, their families, and the communities in which they live.

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