

Ensuring Lasting Smiles Act (Pending Reintroduction; HR. 1916, S. 754 in the 117th Congress)

Please support and become an Original Cosponsor for reintroduction of the bipartisan, bicameral Ensuring Lasting Smiles Act (ELSA), led by Senate sponsors Senator Tammy Baldwin (D-WI) and Senator Joni Ernst (R-IA) and House sponsors Congresswoman Anna Eshoo (D-CA-16) and Congressman Drew Ferguson (R-GA-3). Please contact Senator Baldwin's or Congresswoman Eshoo's office to discuss and receive updates about the bill.

In the 117th Congress, ELSA gained strong bipartisan support from 62 U.S. Senate cosponsors, 317 U.S. House cosponsors, and it passed in the U.S. House of Representatives with a 310-110 vote. In preparation for ELSA's reintroduction in the 118th Congress, the bill sponsors and a broad coalition of patient-focused and professional medical organizations have worked together to strengthen ELSA's language by addressing feedback and incorporating recommendations from federal agencies through technical assistance. ELSA is endorsed by over 70 national health care professional and patient advocacy organizations (see back).

Summary: ELSA would require a group health plan, and a health insurance issuer offering group or individual health insurance coverage, to provide coverage for outpatient and inpatient items and services related to the diagnosis and treatment of a congenital anomaly or birth defect, including any medically necessary item or service to achieve normal body functioning, and it clarifies that coverage includes adjunctive dental, orthodontic, and prosthodontic support. ELSA explicitly excludes coverage of cosmetic surgeries.

Background: About four percent of children in the U.S. are born with congenital anomalies that affect the way they look, develop, or function. Many born with congenital anomalies suffer from severe oral defects (such as cleft lip or palate, hypodontia, enamel hypoplasia), skeletal defects (such as craniosynostosis), vision defects (such as congenital cataracts or aphakia), hearing defects (such as microtia), or other loss of bodily functions. Individuals who do not receive timely, continuous care for their congenital anomalies face long-term physical and psychological injuries.

Problem: Most group and individual health plans state that they include coverages for congenital anomalies, and all 50 states require health insurers to provide coverage for the treatment of congenital anomalies. Despite this, health plans systematically and routinely deny or delay claims and appeals for treatment of many congenital anomalies by wrongfully categorizing certain treatments or body parts as cosmetic or not medically necessary. This common practice leaves families with the burden of paying 100% of the cost for their child's medically necessary treatments, despite maintaining private health plans.

Solution: The Ensuring Lasting Smiles Act will close loopholes and ensure that group and individual health plans do not deny or delay medically necessary treatments of congenital anomalies, therefore improving the health of countless Americans.

Contact Information: To learn more and cosponsor, please contact (Senate) Sen. Baldwin's or Sen. Ernst's office or (House) Rep. Eshoo's or Rep. Ferguson's office.

List of Professional and Patient Organizations Supporting ELSA

(Hyperlinks are active when viewed on computer)

American Association for Dental, Oral, and Craniofacial

Research

Academy of General Dentistry

American Academy of Dermatology Association

American Academy of Facial Plastic and Reconstructive

<u>Surgery</u>

American Academy of Neurology

American Academy of Ophthalmology

American Academy of Oral and Maxillofacial Pathology

American Academy of Oral and Maxillofacial Radiology

American Academy of Pediatrics

American Academy of Pediatric Dentistry

American Association of Oral and Maxillofacial

Surgeons

American Association for Pediatric Ophthalmology and

Strabismus

American Association of Orthodontists

<u>American Association of Women Dentists</u>

American Behcet's Disease Association

American Cleft Palate-Craniofacial Association

American College of Prosthodontists

American College of Surgeons

American Dental Association

American Prosthodontic Society

American Society for Dermatologic Surgery Association

American Society of Dentist Anesthesiologists

American Society of Maxillofacial Surgeons

American Society of Pediatric Otolaryngology

<u>American Society of Plastic Surgeons</u>

<u>Association of Dental Support Organizations</u>

Barth Syndrome Foundation

Born a Hero, Research Foundation

Bridge the Gap - SYNGAP Education and Research

Foundation CCD Smiles

CDH International

Children's Hospital of Wisconsin

Colorado Rare

Costello Syndrome Family Network

Crane Dental Laboratory, Inc.

Dermatology Nurses' Association

Derma Care Access Network

Ear Community, Inc.

EveryLife Foundation for Rare Diseases

Face-To-Face Colorado

FACES: The National Craniofacial Association

FD/MAS Alliance

Foundation for Ichthyosis and Related Skin Types, Inc.

(FIRST)

Genetic Alliance

Georgia Prosthodontics

Gillette Children's Specialty Healthcare

International Pemphigus and Pemphigoid Foundation

Karen S McAndrew DMD, MS, PLC

Lawrence C. Wright Craniofacial Center at John R.

Oishei Children's Hospital

Lupus and Allied Diseases Association

Lymphedema Advocacy Group

M-CM Network

March of Dimes

Moebius Syndrome Foundation

MyFace

National Association of Dental Laboratories

National Organization for Rare Disorders

Noah's Hope - Hope4Bridget

Operation Smile

Ozark Prosthodontics

Pathways for Rare and Orphan Studies

Project Accessible Oral Health

Rare & Undiagnosed Network

rareLife Solutions

Smile Train

Soft Bones: The US Hypophosphatasia Foundation

Sujey Morgan, DDS - Boston Medical Center

SunnyStrong

The APS Type 1 Foundation Inc.

The Marfan Foundation

The XLH Network, Inc.

Usher 1F Collaborative

The Sturge-Weber Foundation

Virginia Council of Nurse Practitioners

Wisconsin Speech-Language Pathology and Audiology

Association (WSHA)