Federal Hemophilia Programs – Ensuring Access to Care, Education and Research

Request: Support full funding for the federal hemophilia programs at the Maternal and Child Health Bureau (MCHB) and Centers for Disease Control and Prevention (CDC) to ensure access to hemophilia treatment centers (HTCs) and other critical education, research and surveillance activities.

About Bleeding Disorders and Hemophilia Treatment Centers (HTCs)

- Hemophilia is a rare, chronic bleeding disorder affecting about 20,000 people in the US. Von Willebrand Disease (VWD) is a related bleeding disorder that affects up to 1 million Americans.
- People with hemophilia require life-long infusions of expensive clotting factor therapies that replace missing or deficient blood proteins, thus preventing debilitating and life-threatening internal bleeding.
- Since 1974, Congress has authorized and funded HTCs to ensure access to comprehensive, multidisciplinary, specialized care for people with bleeding disorders.
- CDC studies show that mortality and hospitalization rates are 40% lower in people who use HTCs compared with those who do not, even though more severely affected patients are more likely to be seen in HTCs.
- A patient needs assessment survey completed in 2013 found that 90% of the 4000 survey respondents felt it was important to receive care at a federally-funded HTC and that the four HTC core services (hematology, nursing, social work and physical therapy) met their needs.

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Health Resources and Services Administration (HRSA) Maternal and Child Health Bureau (MCHB)

- With $4.9 million in funding from HRSA, HTCs provide critical, comprehensive multidisciplinary services not typically covered by insurance, such as physical therapy assessments, social work and case management services.
- In keeping with HRSA’s emphasis on patient-centered, integrated care, HTCs serve as "specialty medical homes" for people with bleeding disorders, from childhood through end of life. HRSA recognizes the centers' role in providing comprehensive disease management and considers them a model for chronic disease care.

CDC Division of Blood Disorders

- The CDC Division of Blood Disorders program includes $5 million for HTC research, surveillance, and prevention activities. This funding level must be maintained to monitor blood and blood product safety within the hemophilia population and to improve treatment and prevention strategies.
- CDC supports outreach and education programs provided by the national patient organizations in hemophilia and other bleeding and clotting disorders. These programs received significant reductions in 2014 when funding for the Division was reduced by nearly $6 million. Funding for these important programs should be restored to 2013 levels to ensure that important patient education and outreach activities in hemophilia, von Willebrand Disease, and women with bleeding disorders are maintained.

 Federal hemophilia programs must be adequately funded to ensure that patients with bleeding disorders continue receiving high-quality care.