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The Need for Comprehensive Care for Persons with Chronic Immune Thrombocytopenic Purpura

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Abstract: Chronic platelet disorders (CPD), including chronic immune thrombocytopenic purpura (cITP), thrombotic thrombocytopenic purpura (TTP) and platelet function disorders are among the most common bleeding disorders and are associated with morbidity and mortality. The clinical phenotype and complexity of cITP is much like that of hemophilia. In cITP and hemophilia, bleeding is problematic for many, complicating employability, insurability and overall quality-of-life (QoL). While myriad drug therapies are available for cITP and hemophilia, each are variable in their effectiveness, very few (except for clotting factor concentrates for hemophilia) alter the natural history of the disorder and sometimes contribute to specific morbidities and mortality. Like in hemophilia, the management of cITP is not solely based on access to effective treatment but also includes accurate diagnosis and comprehensive care by a multidisciplinary team of specialists trained in the management of bleeding disorders. The model of comprehensive care in Hemophilia Treatment Centers (HTCs) has been recognized as highly effective, improving life expectancy for persons with hemophilia. cITP, and other CPDs, are complex disorders requiring specialized care. However, an integrated care model with a systematic and reliable population-based surveillance program does not exist. Extending the Comprehensive Care model with all its related benefits to the community of persons with cITP is sorely needed. This review will focus on cITP as a prototype chronic platelet disorder that could benefit greatly from the Comprehensive Care model.

Keywords: hemophilia, hemorrhage, idiopathic thrombocytopenic purpura, platelet, primary immune thrombocytopenia, thrombocytopenia

Historical Perspectives of Hemophilia and ITP

The analogous history of cITP and hemophilia cannot be overlooked; with similar advancements in recognition, disease characterization and therapy occurring along comparable timelines. In the nineteenth century, both hemophilia and ITP were recognized; ITP was described as morbus maculosus hemorrhagicus and hemophilia was noted to be a bleeding disorder in males of certain families.1,2 Further advancement was made with the use of plasma therapy for hemophilia and splenectomy for treatment of cITP in the first part of the twentieth century.3,4 In the 1950s–1960s, the distinction between Hemophilia A and B was made and cITP was determined to be an autoimmune disorder.5,6 This led to further progress in therapies with cryoprecipitate used for hemophilia A and corticosteroids as treatment for cITP.7,8 In the 1970s-1980s, the discovery of concentrated plasma derived clotting factor allowed for home
treatment of hemophilia. However, this advancement was dampened by the transmission of human immunodeficiency virus (HIV), hepatitis C virus (HCV) and hepatitis B virus (HBV) to persons with hemophilia. This led to widespread fatalities and subsequent improvement in screening methods for blood donors and pathogen attenuation steps in the manufacturing process. During the same era, intravenous immunoglobulin (IVIG) was shown to be an effective treatment for persons with ITP in the 1980s. As IVIG is derived from pooled donor plasma, reports of pathogenic virus transmission (HCV, HBV) emerged between the mid-1980s and the mid-1990s. This risk was mitigated with the introduction of pathogen attenuation steps during the production of IVIG.

Advances in the treatment of hemophilia from the last decade of the twentieth century through the first decade of the twenty-first century largely refined the quality of clotting factor concentrates. Recombinant clotting factors emerged in the mid-1990s and are still considered by many to be the standard of care for persons with hemophilia. Most recently, non-factor therapies, that either mimic the effect of factor VIII or “rebalance” the hemostatic defect, have benefited patients, especially those with neutralizing antibodies (inhibitors) to clotting factor. Gene therapy to correct the inherited defect is the next anticipated treatment advancement.

For cITP, treatment options during the later 1990s, until the late 2000s remained largely unchanged, except for the discovery that a hybridized anti-CD20 monoclonal antibody raised the platelet count in around half of the recipients. The 2000s saw affirmation to the longstanding theory that many patients with cITP not only have accelerated platelet destruction but also impaired platelet production. The thrombopoietin receptor agonists (TPO-RAs) have subsequently become a vital second-line therapy for adults and children with cITP. Today, targeted immunotherapy is a major focus of drug development for persons with cITP (see Figure 1).

**Epidemiology**

Hemophilia is almost always caused by hereditary factor VIII or factor IX deficiency that creates a lifelong bleeding
disorder. Hemophilia affects approximately 400,000 persons worldwide with more recent estimates suggesting the prevalence is over 1 million. In the United States, the prevalence of hemophilia is estimated to be approximately 20,000, based on expected births and deaths since 1994. Hemophilia A occurs in 1 in 5000 live male births and is four times as common as hemophilia B. The overall life expectancy of persons with hemophilia is now comparable to that of the general population.

cITP is an acquired autoimmune disorder that causes isolated thrombocytopenia and related bleeding. The prevalence of cITP in the United States is approximately 60,000 persons; nearly three times more prevalent than hemophilia. The diagnosis of cITP is based primarily on the exclusion of other causes of isolated thrombocytopenia and there are no clinical or laboratory parameters to establish its diagnosis. The incidence of cITP is difficult to establish due to limited studies and variable symptomology, the mildest of which falls short of the threshold for platelet count testing. However, the strongest estimate of the incidence of acute ITP in children is between 1.9 and 6.4 per 100,000 children/year and in adults, is between 1.6 and 3.9 per 100,000 adults/year.

**Shared Clinical Challenges in Hemophilia and cITP**

Hemophilia and cITP are bleeding disorders with a spectrum of phenotypes and much clinical overlap. The clinical phenotype of hemophilia is directly related to the severity of the factor deficiency, but with large variability in the bleeding phenotype among patients with any severity of hemophilia. The clinical manifestations of hemophilia include spontaneous bleeding as well as delayed bleeding after trauma. Beginning in infancy, patients may present with a cephalohematoma, subgaleal or intracerebral hemorrhage. As they grow into toddlers and then children, they often present with hemarthrosis, intramuscular hemorrhage, soft tissue hematomas, hematuria, and, less commonly, hematoma, hemaemorhage and intracranial hemorrhage. Intracranial hemorrhage (ICH) remains the main cause of fatal bleeding in persons with hemophilia.

Bleeding associated with hemophilia and cITP are very similar, with the exception that persons with cITP rarely have bleeding in their joints (Table 1). Mucocutaneous, gastrointestinal and genitourinary bleeding are common, fortunately CNS hemorrhage is less common, especially in the young. The severity of thrombocytopenia correlates directly with the bleeding risk but not entirely, many patients with very low platelet counts have only mild bleeding such as easy bruising. Comparatively, persons with severe bleeding most often have platelet counts less than 10,000–20,000/μL.

Like hemophilia, the main cause of fatal bleeding in patients with ITP is intracranial hemorrhage (ICH), the incidence of which is 0.1–0.5% in children, 1% in adults with acute ITP, and 5% in adults with cITP. Severe and life-threatening bleeding affects a subset of persons with hemophilia and cITP which may alter the lifespan of persons with either disorder.

Table 1 Comparison of Clinical and Non-Clinical Aspects of Hemophilia and cITP

<table>
<thead>
<tr>
<th></th>
<th>Hemophilia</th>
<th>cITP</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Age Affected</strong></td>
<td>Beginning at Birth</td>
<td>Any Age</td>
</tr>
<tr>
<td><strong>Duration</strong></td>
<td>Lifelong</td>
<td>Typically Lifelong Once Diagnosed</td>
</tr>
<tr>
<td><strong>Main Sites of Bleeding</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Joints</td>
<td>Common</td>
<td>Rare</td>
</tr>
<tr>
<td>• Mucocutaneous</td>
<td>Common</td>
<td>Common</td>
</tr>
<tr>
<td>• Gastrointestinal</td>
<td>Common</td>
<td>Common</td>
</tr>
<tr>
<td>• Genitourinary</td>
<td>Common</td>
<td>Common</td>
</tr>
<tr>
<td>• CNS</td>
<td>Uncommon</td>
<td>Uncommon</td>
</tr>
<tr>
<td><strong>Treatment costs</strong></td>
<td>Very expensive</td>
<td>Very expensive</td>
</tr>
<tr>
<td><strong>Health-related Quality of Life</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pain, hemarthrosis, anxiety, ADL limitations, work/ school absenteeism, social functioning, treatment related burden/concerns</td>
<td>Fatigue, anxiety, depression, ADL limitations, work/ school absenteeism, social isolation, treatment related burden/ concerns</td>
<td></td>
</tr>
<tr>
<td><strong>Advocacy</strong></td>
<td>International: WFH</td>
<td>National: NHF, HFA, HA</td>
</tr>
<tr>
<td></td>
<td>National: NHF, HFA, HA</td>
<td>State: National Organization</td>
</tr>
<tr>
<td></td>
<td>Chapters</td>
<td>Chapters</td>
</tr>
<tr>
<td></td>
<td>Local: HTCs</td>
<td>Local: HTCs</td>
</tr>
<tr>
<td><strong>Comprehensive Care Centers</strong></td>
<td>National network of HTCs</td>
<td>None</td>
</tr>
</tbody>
</table>

**Abbreviations:** cITP, chronic immune thrombocytopenia purpura; ICH, intracranial hemorrhage; CNS, central nervous system; ADL, activities of daily living; WFH, World Federation of Hemophilia; NHF, National Hemophilia Foundation; HFA, Hemophilia Federation of America; HA, Hemophilia Alliance; HTCs, Hemophilia Treatment Centers; PDSA, Platelet Disorder Support Association.
related to cITP is left untreated, a 70-year-old woman loses 5 quality-adjusted life years (QALY), whereas a 25-year-old woman will lose approximately 15 QALY of potential life expectancy.33

Improving Quality of Life for Persons with cITP
At an early age, persons with hemophilia must learn to adapt to the essential therapies and the overall complexity of the disorder. As treatments and therapies continue to advance, there has been an emphasis on health-related quality of life (HRQoL) as an outcome measure.36 The long-term complications of hemophilia have had effects on HRQoL, revealing that persons with hemophilia experience depression and anxiety more often than the general population.36 In addition, the presence of one or more target joints has a major impact on HRQoL for persons with severe hemophilia.36

Like in hemophilia, it is important to understand how cITP can impact HRQoL (Table 1). The effects of cITP on a person's HRQoL should not be overlooked; including emotional and functional health, work life, social and leisure activities, and reproductive health.37 Patients characterize the emotional toll by depression, anxiety and report it affecting their overall mental health. In addition, fatigue, fear of bleeding, and the need to limit daily activities are pervasive sequelae of cITP.38 cITP during childhood may negatively impact the HRQoL of the child with cITP and their parents.38 However, when children receive platelet-enhancing therapy to maintain a platelet count in the hemostatic range, HRQoL in children and parental burden improve.39 Although studies of HRQoL have helped in our understanding of the clinical and sociological burden for persons with cITP, more short- and long-term studies are needed to fully appreciate the struggle experienced by many patients.

The Evolution of Comprehensive Care for Hemophilia
Comprehensive Care for persons with hemophilia is defined as the continuous supervision of all medical and psychological aspects affecting the patient and their family.40 Early detection, diagnosis, prevention, treatment, along with psychosocial and educational support are all factors needed to provide optimal management for patients with bleeding disorders.41 Recognized since the 1960s, comprehensive clinical management of hemophilia and related inherited bleeding disorders require a multidisciplinary integrated approach at specialized establishments called Hemophilia Treatment Centers (HTCs).42 Over the past 60 years, this model has been expanded to include patients with other rare inherited or acquired bleeding disorders such as von Willebrand disease, Hereditary Hemorrhagic Telangiectasia, and acquired hemophilia, to name a few.

The approach implemented by Comprehensive Care clinics includes focused education incorporating the basics of hemophilia and other bleeding disorders. These include training about bleeding prevention, managing complications, QoL improvement and self-advocacy.41 In addition, advocacy organizations exist on the local, state, national, and international levels; providing education, empowering individuals, and aiding in financial burdens by actively pursuing long-term relationships with individuals and organizations that align with the tenets of Comprehensive Care.33,44 For patients with hemophilia, HTCs provide multidisciplinary expertise in bleeding disorders, individualized management plans, readily available access to multiple medical disciplines, with a focus on preventative medicine.45 Collectively, this design optimizes effectiveness and efficiency of health care management.

The model of care in comprehensive HTCs has become the standard for persons with hemophilia worldwide. It has improved access to care for many thousands of patients while providing the opportunity for family involvement which improves therapy adherence.40 The HTCs have advocated progress through basic, translational and clinical research. In addition, care at HTCs reduces health care costs, by decreasing emergency room visits, decreasing absenteeism from work or school and preventing life-altering sequelae of hemophilia.45 In another CDC-sponsored study, HTCs were shown to reduce mortality among persons with hemophilia by 40% as well as decrease healthcare resource utilization.45,46 Comprehensive care is one of the essential principles in the model for the development of a national healthcare program for patients with hemophilia.31 HTC have been established to guarantee patients and families access to all services essential for the management of their bleeding disorder.31 Within these comprehensive HTCs, the core team members typically include hematologists trained in the management of bleeding disorders, nurses, clinical specialists, nurse practitioners, laboratory staff, physical therapists, medical social workers, and may include dentists, genetic counselors, dieticians, clinical research coordinators, psychologists as well as other specialists that aid in the coordination of care.
Current standard of care for patients with hemophilia now provides safe and effective treatment, allowing growth into the adult life with minimal disability, a normal lifestyle and employment opportunities. In the 1990s, approximately 67% of persons with hemophilia received care at HTCs at least once a year. More recently, in one US HTC, approximately 81.7% of persons with hemophilia receive care at least once a year. Those that benefit from care at HTCs have a reduced rate of emergency department (ED) utilization and an increased rate of prophylaxis and self-infusion compared to those receiving care outside of the HTC network.

**Potential Value of Comprehensive Care for cITP**

Despite advances in medical therapy, the long-term morbidity and mortality of cITP remains. In the United States, a limited number of hematologists have focused expertise in cITP. To improve health outcomes and HRQoL, a network of medical centers that focuses on the diagnosis, management and prevention of complications of cITP is needed. The contributing centers would adopt or extend the Comprehensive Care model within an existing Hemophilia Treatment Center. This would include disease-specific education, outreach and advocacy programs, and conduct longitudinal surveillance on the complications of treatment. The Comprehensive Care model utilized in persons with hemophilia is well-suited for patients with platelet disorders. This model already incorporates integrated care for patients with inherited platelet disorders, including disorders of platelet number and/or function. This includes Bernard Soulier syndrome, Glanzmann Thrombasthenia, MYH9 disorders and Platelet Storage Pool Defect, to name a few.

The core members of the Comprehensive Care cITP team should be the same as those in the Comprehensive Care model for hemophilia: hematologists, nurse practitioners, nurses, dentists, laboratory staff, physical therapists, medical social workers, and clinical psychologists, but may include immunologists, dieticians, and clinical research coordinators. Patients with cITP may be broadly affected by the disease, specifically the clinical consequences of bleeding episodes, fear of bleeding, fatigue, decreased emotional health, social isolation, and reproductive health issues (see Table 1).

The model of Comprehensive Care in HTCs has been recognized to improve many aspects of health, including life expectancy. An expert team would be equally as valuable to persons with cITP. However, an integrated care model with a reliable population-based surveillance program does not exist for cITP. Although advocacy groups, like Platelet Disorder Support Association (PDSA), have provided an invaluable resource for persons with cITP for the past 20 years, integration with the site of clinical care is usually lacking. The PDSA would be a logical liaison to the ITP treatment centers for patient education and advocacy, similar to the relationship of the National Hemophilia Foundation (NHF) to HTCs. Given the similarities between cITP and hemophilia and the success of comprehensive care centers in caring for persons with hemophilia, the Comprehensive Care model for hemophilia should be extended to persons with cITP.

In late 2017, the Bleeding and Clotting Disorders Institute (BCDI) in Peoria, Illinois launched the first ever Comprehensive Care Clinic in the US for persons with cITP. Beyond the Comprehensive Clinic team (hematologist, nurse practitioner, nurse coordinator, physical therapist, pharmacist, dentist, and medical social worker), each cITP patient is also seen by an immunologist and dietician. The addition of the immunologist has proven to be particularly beneficial and has led to the diagnosis of additional immunological disorders in some patients, such as common variable immune deficiency (CVID) and various autoimmune disorders.

The ITP program at BCDI provides care for approximately 200 patients from 16 US states and Mexico. Of these, more than fifty have been served on at least one occasion in the Comprehensive Care Clinic. The implementation of comprehensive care transcends the formal clinic encounter and, in some cases results in more than 200 interactions per year between the patient and various members of the care team. The response to the Comprehensive Care Clinic for cITP at BCDI has been tremendously positive and patients have reported an overall improvement in ease of their activities of daily living. Formal studies of clinical, psychosocial and economic outcomes are needed and in process at BCDI.

**Conclusions**

The complex phenotype of cITP is much like that of hemophilia, necessitating specialized care. A better understanding of the natural history of cITP, including its impact on HRQoL and drug safety in the treatment of cITP is urgently needed. To accomplish this, the Comprehensive Care model used for persons with hemophilia should be
extended to include adults and children with cITP by adapting the infrastructure used for HTCs in the US to serve this patient population.

Disclosure
Michael Tarantino reports The Bleeding and Clotting Disorders Institute - salaried position, CEO, CMO, personal fees from Angen, Principia, Genentech, Grifols, HemaBiologics, Novo Nordisk, Octapharma, Takeda, and Spark Therapeutics, outside the submitted work. The authors report no other potential conflicts of interest for this work.

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