Glanzmann thrombasthenia: Genetic basis and clinical correlates

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Glanzmann thrombasthenia: genetic basis and clinical correlates

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**Abstract**

Glanzmann thrombasthenia (GT) is an autosomal recessive disorder of platelet aggregation caused by quantitative or qualitative defects in integrins αIIb and β3. These integrins are encoded by the ITGA2B and ITGB3 genes and form platelet glycoprotein (GP)IIb/IIIa, which acts as the principal platelet receptor for fibrinogen. Although there is variability in the clinical phenotype, most patients present with severe mucocutaneous bleeding at an early age. A classic pattern of abnormal platelet aggregation, platelet glycoprotein expression and molecular studies confirm the diagnosis. Management of bleeding is based on a combination of hemostatic agents including recombinant activated factor VII with or without platelet transfusions and antifibrinolytic agents. Refractory bleeding and platelet alloimmunization are common complications. In addition, pregnant patients pose unique management challenges. This review highlights clinical and molecular aspects in the approach to patients with GT, with particular emphasis on the significance of multidisciplinary care.

**Introduction**

The Swiss pediatrician Eduard Glanzmann was the first to describe in 1918 a type of purpura that presented with normal platelet count and size in patients with absent/decreased clot retraction and prolonged bleeding time.1 In 1962, Caen and Cousin described the lack of platelet aggregation with multiple agonists,2 and a few years later correlated the platelet aggregation result with decreased platelet fibrinogen and a slight reduction in clot retraction in individuals with Glanzmann thrombasthenia (GT).3 The lack of one of the three major platelet surface glycoproteins in GT was reported by Nurden and Caen in 1974;4 several other scientists subsequently identified this as the glycoprotein (GP) IIb/IIIa complex. Additional studies with more advanced glycoprotein imaging techniques provided evidence for the definition of two disease groups: type I with absent IIb/IIIa expression (<5% of normal) and type II with reduced expression (5-20% of normal GP IIb/IIIa).5 A third type (type III) has normal levels of integrin but the protein is non-functional.

Glanzmann thrombasthenia is considered a rare disease, defined in the United States as those affecting less than 200,000 individuals.6 The exact incidence has been difficult to calculate, but is estimated at one in 1,000,000. With an autosomal recessive inheritance, males and females are affected equally. There is a world-
wide distribution; however, a large proportion of the cases have been described in selected populations such as the French Romani, South Indian Hindus, Israeli Jews, and Jordanian nomadic tribes, all of which consanguinity is common. Type I is the most common subtype and accounts for around 78% of patients with GT type II and type III (functional variant in receptor) constituting around 14% and 8% of cases, respectively.

GPIIb/IIIa (integrin αIIbβ3) is a heterodimeric receptor present in large quantities in the plasma membrane of platelets. Activation of this integrin and binding of soluble ligands [primarily fibrinogen, but also von Willebrand factor (vWF) and fibrinectin] are essential for platelet aggregation. In the resting state, the integrin has low affinity for ligands. During platelet activation, driven by exposure to soluble agonists or the subendothelial matrix, “inside out” cellular signaling generates a conformational change in GPIIb/IIIa that allows high affinity binding for fibrinogen, which serves as a “bridge” to other activated platelets, eventually forming the platelet plug. Protein kinase C (PKC), diacylglycerol-regulated guanine nucleotide exchange factor I (CaldAG-GEFI or RasGRF2), and phosphoinositide 3-kinase (PI3K) participate in this signaling pathway. Subsequent “outside in” signaling triggers additional granule secretion, cytoskeletal interactions (that allow for platelet spreading), stabilization and clot retraction to consolidate the fibrin clot. Kindlins (including kindlin-3) and talins are key regulators of integrin activation.

Glanzmann thrombasthenia is usually caused by decreased or absent expression of αIIb or β3, abnormalities in protein folding, defective post-translational processing or transport of either integrin subunit causing decreased surface expression, or abnormalities affecting protein function. Other defects change integrin function by altering the ligand binding pocket (interface between αIIb and β3), which modifies the cytoplasmic domain and affects binding of regulators, or locks the integrin in the activated form.

Molecular basis of Glanzmann thrombasthenia

The ITGA2B and ITGB3 genes are located on chromosomes 17q21.31 and 17q21.32, respectively, and are independently expressed. GT is caused by pathogenic variants in both alleles of either of the two genes; concomitant pathogenic variants in both genes, but affecting only one allele of each, is not known to cause GT. Due to the autosomal recessive inheritance, compound heterozygosity is frequent, except in selected ethnic groups where homozygosity is more likely due to consanguinity. A higher percentage of pathogenic variants occur in ITGA2B, likely because of the larger size of this gene with 30 exons encoding 1,039 amino acids, compared to ITGB3 which is composed of 15 exons with 788 amino acids. The clinical phenotypes associated with either gene are indistinguishable.

Pathogenic nonsense, missense and splice site variants are common and large deletions and duplications, although rare, have also been described. Pathogenic missense variants impair subunit biosynthesis in megakaryocytes or inhibit the transport of the pro-αIIbβ3 complexes from the endoplasmic reticulum (ER) to the Golgi apparatus or the export of the mature complexes to the cell surface. A large proportion of variants affect the β-propeller region of αIIb and the epithelial growth factor domains of β3.

A different type of variant affecting specific regions of these genes has been more recently described to cause a mild autosomal dominant macrothrombocytopenia by interfering with proplatelet formation. These “gain of function” variants cause spontaneous activation of GPIIb/IIIa (αIIbβ3) by affecting the cytoplasmic domains or the membrane proximal residues in the extracellular domains. The majority of these are in ITGB3 and affect the MIDAS (metal ion dependent adhesion site), ADMIDAS (adjacent to MIDAS) or SyMBS (synergistic metal ion binding site) regions. A small proportion have been reported in ITGA2B and affect the conserved intracellular GFFKR sequence.

Clinical manifestations and diagnosis

Bleeding phenotype

With integrins αIIb and β3 participating in primary hemostasis, the bleeding manifestations are typically purpura, epistaxis (60-80%), gum bleeding (20-60%), and menorrhagia (60-90%). Gastrointestinal bleeding in the form of melena or hematochezia is present in 10-20%, and 1-2% develop intracranial hemorrhage. Mucocutaneous bleeding can be spontaneous or occur after minimal trauma. Epistaxis is the most common cause of severe bleeding, especially in the pediatric population, and risk of severe nosebleeds decreases with age as the septal arterial plexus becomes less friable and children grow out of the habit of nose picking. Menorrhagia is highly prevalent in affected females and there is a higher risk of severe bleeding at the time of menarche due to the prolonged estrogen influence on the proliferative endometrium that occurs during anovulatory cycles. Bleeding complications during pregnancy are uncommon; however, the risk of obstetric hemorrhage at the time of delivery and postpartum is high. Hematuria and spontaneous hemarthrosis have been described in some cases but are not usually part of the bleeding phenotype.

Several bleeding scores have been developed with the goal of standardizing the assessment of bleeding and facilitating the diagnosis of patients with a suspected inherited bleeding disorder. These are useful tools that assist in communication of the bleeding phenotype in the clinical and research setting; however, they have not been widely validated for patients with inherited platelet function disorders. Therefore, specific cutoffs to define a positive bleeding score have not been established for this population. This is of particular relevance in GT, because while the types of bleeding are consistent among individuals, the degree of bleeding is highly variable. Given the severity of this disorder, historically, most patients have been diagnosed in childhood (before 5 years of age), but there are some patients that reach adulthood without having severe bleeding. In general, the severity of bleeding (except for menorrhagia and pregnancy-associated bleeding) decreases with age.

Laboratory phenotype

Complete blood count

There should be normal platelet count and size with normal granularity on evaluation of the peripheral blood smear by light microscopy. If bleeding is severe and/or...
chronic, patients can have low hemoglobin, microcytosis and increased red cell distribution width from secondary iron deficiency. Other abnormalities of the complete blood count (CBC), suggest an alternative diagnosis.

**Coagulation screening tests**

Routine tests ordered in the workup of a patient with abnormal bleeding, such as prothrombin time (PT), activated thromboplastin time (aPTT) and fibrinogen, are usually normal, unless a patient is being evaluated in the setting of significant acute hemorrhage and has evidence of consumptive coagulopathy.

**Platelet function screening tests**

The platelet function analyzer (PFA)-100 provides a measure of platelet function under high shear. It is convenient as it uses low volume whole blood samples and is widely available to clinicians. Very prolonged closure times (>300 seconds) are compatible with GT but not specific, as other disorders, such as severe von Willebrand disease, Bernard Soulier syndrome and afibrinogenemia, can produce the same result. However, a normal PFA-100 has a very high negative predictive value for GT and virtually excludes this diagnosis.19

**Platelet light transmission aggregometry**

Despite its limited availability and the need for larger-volume samples and immediate processing, platelet light transmission aggregometry (LTA) remains the gold standard in the clinical diagnosis of GT. It is based on the decreased turbidity generated by platelet agglutinates or aggregates in platelet rich plasma after exposure to different agonists. Decreased/absent aggregation (<10%) with all physiologic agonists, together with a normal agglutination response to ristocetin (mediated by GPIb-IX-V), is the classic pattern observed in patients with GT.20 Due to the large variability in platelet aggregation results and the significant effect of pre-analytical variables on this test, confirmation of the findings in a second sample is recommended.

**Whole blood impedance aggregometry**

This is available in several centers worldwide. While it can be performed in whole blood samples and using lower volumes, there is not enough evidence to support equivalent sensitivity and reproducibility when compared to LTA.21 There is some clinical utility to this test in cases in which access to LTA is difficult; however, patients should preferably be referred for evaluation at a center that has LTA capacity at least once to confirm the diagnosis.

**Platelet glycoprotein expression study by flow cytometry**

This assesses quantitative platelet surface glycoprotein deficiencies using fluorescent-conjugated antibodies that are specific towards GP.22 This test can be performed in low sample volumes shipped for analysis; however, it will not identify type III (functional) defects that are caused by qualitative but not quantitative defects in GPIb/IIa (Figure 1).

**Differential diagnosis**

**Leukocyte adhesion deficiency type III**

Leukocyte adhesion deficiency type III (LAD-III) is an autosomal recessive disorder caused by pathogenic vari-
Molecular confirmation
Genetic analysis is clinically useful for confirmation of the diagnosis, identification of at-risk carriers, reproductive risk counseling for a given couple/family, and for definitive prenatal or preimplantation genetic diagnosis. Genetic counseling plays an essential role in the genetic testing process, obtaining informed consent regarding the unique considerations, benefits, and limitations of genetic testing, and addressing the complexities of clinical application or molecular findings, unexpected results, variants of uncertain significance, and familial implications for medical risks or biological relationships. Genetic coun-

**Figure 2. Diagnostic flow chart for the laboratory evaluation of a patient with suspected Glanzmann thrombasthenia (GT).**

*Can be performed simultaneously or as reflex depending on laboratory and technique used
VWF: von Willebrand Factor
selors can support clinicians by providing pre- and post-test genetic counseling to patients and families in the clinical setting, and can also aid clinicians in test selection and ordering, and in navigating the process of obtaining insurance approval for payment. This is particularly relevant in healthcare systems with multiple payers and multiple testing options. As commercial laboratories offer different services, the clinician should be familiar with the testing methodology and any related limitations.

Given the highly specific phenotype of GT, after platelet aggregation studies and flow cytometry have been performed, genetic testing of ITGA2B and ITGB3 only, as opposed to a panel approach that includes multiple other genes, is appropriate (Figure 2). Sequence analysis will detect the vast majority of pathogenic variants; when sequencing fails to identify both pathogenic variants in a patient with GT, specific deletion/duplication analysis should be considered.

Targeted variant analysis, which is expected to save time and money, can be performed in populations with a known pathogenic variant(s) and high rates of consanguinity; however, with this approach, a second GT variant in the same gene may go undetected. Targeted variant analysis is best adopted when the pathogenic variant in each allele has been identified in an affected individual.

**Variant curation and initiatives for standardization**

In recent years, several initiatives have been taken to provide guidance for the analysis and reporting of molecular findings, specifically in terms of variant classification when attributing pathogenicity. In the United States, the American College of Medical Genetics and Genomics (ACMG) and Association for Molecular Pathology (AMP) published guidelines in 2015 for the interpretation of variants, providing a framework for clinical molecular laboratories. While these guidelines have been extremely useful in the laboratory practice, they are not disease specific, and significant challenges remain when applied to particular disease states. In an effort to address this limitation, and to improve the quality, consistency and access to clinical genomic data, the National Institutes of Health (NIH) funded resource ClinGen is leading initiatives including the formation of Variant Curation Expert Panels (VCEP). Each VCEP consists of a cross-institutional team of experts from complementary specialties, the fields of hematology and genetics, from academia and industry, working collaboratively with the goal of adapting the ACMG criteria for specific genes or disorders.

The Platelet Disorder Variant Curation Expert Panel (VCEP) created in 2018 and supported by the American Society of Hematology, is composed of 28 international scientists and clinicians with expertise in hematology and genetics, and has been working on the adaptation of the ACMG rules for the interpretation of variants in ITGA2B and ITGB3. The goals of this collaboration are to produce high quality variant curation data that are publicly available and create a framework for GT variant curation that allows others to systematically and comprehensively approach genetic data encountered in clinical and research settings.

The VCEP has used estimates of prevalence and population data to define criteria for allele frequency that ultimately serve as standalone, strong or supporting evidence to classify variants as benign, which is a difficult task in rare disorders for which the precise incidence data are not available. Clinical expertise was key in defining the phenotype from a clinical presentation (bleeding phenotype) and clinical laboratory standpoint (platelet aggregation and glycoprotein expression studies), which together are unique to GT. Knowledge of the molecular biology of the disorder allowed for elimination of codes that do not apply for this disease state and points for segregation analysis were modified taking into account the disease inheritance, low frequency and specific clinical phenotype. Special consideration was given to defining the type of assays that provide quality functional evidence in different *in vivo* and *in vitro* systems and models. The disease specific rules were tested in a subset of variants which will be uploaded to ClinVar with a 3-star rating. Variants assessed by approved ClinGen VCEP also receive a US Food and Drug Administration (FDA) approval label. A detailed description of the rule specifications for GT will soon be available to the public.

**Management**

Patients with GT benefit from being managed at a center with expertise in inherited bleeding disorders, with access to staff who are able to provide recommendations and

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Clinical situation</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Local measures</strong></td>
<td>Mild bleeding at a visible/compressible site</td>
</tr>
<tr>
<td>• Compression</td>
<td></td>
</tr>
<tr>
<td>• Gauze or sponge dipped in antifibrinolytic or topical thrombin</td>
<td></td>
</tr>
<tr>
<td>• Nasal packing</td>
<td></td>
</tr>
<tr>
<td>• Gel foam soaked in antifibrinolytic or topical thrombin</td>
<td></td>
</tr>
<tr>
<td><strong>Antifibrinolytics</strong></td>
<td>As monotherapy in mild bleeding</td>
</tr>
<tr>
<td>• Tranexamic acid</td>
<td></td>
</tr>
<tr>
<td>• Aminocaproic acid</td>
<td>As combination therapy in moderate and severe bleeding</td>
</tr>
<tr>
<td><strong>rFVIIa</strong></td>
<td>Moderate to severe bleeding</td>
</tr>
<tr>
<td></td>
<td>Alone for minor surgical procedures or in combination with antifibrinolytics and/or platelet transfusions for major procedures and severe or life-threatening bleeding</td>
</tr>
<tr>
<td><strong>Platelet transfusion</strong></td>
<td>In combination with rFVIIa and/or antifibrinolytics in severe bleeding and major surgical procedures</td>
</tr>
</tbody>
</table>

rFVIIa: recombinant activated coagulation factor VII. *Risk of lack of efficacy in the setting of alloimmunization.
treatment any time of day, and also benefit from being provided directly with medical alerts, and emergency medical contact and treatment information to present when seeking urgent care from clinicians who are not familiar with their case history. Education on avoidance of over-the-counter medications that increase bleeding risk such as non-steroidal anti-inflammatory and aspirin products should be provided. Prescription medications that can affect hemostasis should be carefully monitored.

### Hemostatic management

The treatment of bleeding in patients with GT includes management of acute or chronic bleeding and prevention of hemorrhagic complications around the time of procedures. The choice of treatment depends on the severity of bleeding (Table 1), availability of products, and prior responses to therapy, and is similar in some aspects to the management of patients with other severe bleeding disorders. Note that desmopressin (DDAVP), commonly used in von Willebrand Disease and other milder disorders of platelet function, is of limited clinical utility in the treatment of GT (Table 1).

Recombinant activated factor VII (rFVIIa) is approved by the European Medicines Agency (EMA) and the FDA for the treatment of patients with GT. The approval of this drug has changed the landscape of treatment of GT and has allowed for better hemostatic outcomes in all patients, especially those who do not respond to platelet transfusions. Optimal dosing and interval of doses vary by center and clinical situation. Typical doses for acute bleeding are 90 mcg/kg intravenous (IV) every 2-6 hours until hemostasis is achieved (at least 3 doses). Perioperative dosing is 90 mcg/kg immediately before surgery and every two hours during the procedure, with doses every 2-6 hours postoperatively to prevent postsurgical bleeding. Effectiveness of this drug in the treatment of acute bleeding is high when started early in the course of bleeding and used in combination with other hemostatic treatment. It is also useful for perioperative management. Although cases of thrombembolism during rFVIIa therapy in GT patients have occurred, it remains a safe medication in this population, with low rates of adverse reactions.

### Transfusion

Patients with GT have a high risk of development of isoantibodies, with up to 30% of patients developing anti-GPIIb/IIIa or anti-HLA antibodies after platelet transfusion. Patients with pathogenic variants causing premature stop codons and leading to absent GPllb/IIIa are at the highest risk of anti GPllb/IIIa alloimmunization compared to those with other types of variants (81% vs. 25%). Once these antibodies develop, the patient may no longer respond to platelet transfusion. For this reason, platelet transfusions should be reserved for only major surgeries, life-threatening bleeding, and significant bleeding that does not respond to the above measure. Transfusions in women of reproductive age should ideally be avoided as the antibodies can cross the placenta and affect the fetus.

### Bone marrow transplant

Allogeneic stem cell transplantation has been successfully performed in selected patients with severe recurrent bleeding using reduced intensity conditioning with good clinical outcomes. The presence in the recipient of antiplatelet antibodies affecting the graft remains a challenge in this patient population.

### Pregnancy

Pregnant women with GT have a high rate of complications and are best managed in a specialized center with a multidisciplinary team. While most complications relate to bleeding and occur at the time of delivery, management of the pregnant GT patients should start in the prenatal period. Counseling of the pregnancy-associated risks and screening of the father in consanguineous families to identify at-risk fetuses is important. Identification of HLA or GPllb/IIIa antibodies during pregnancy, which are present in up to 70% of patients, is key for planning delivery. In general, regional anesthesia is contraindicated and support with rFVIIa and antifibrinolytics is given for vaginal deliveries with the option of adding platelet transfusion for cesarian sections. Primary post-partum hemorrhage is common, and a large proportion of women will require red cell transfusion. Clinicians should be aware that, given the phenotypic variability of this disorder, around half the women with GT who are pregnant may not be aware of the diagnosis.

### Future directions

Gene therapy is highly promising in providing a cure for patients with GT, with significant progress made using different techniques, vectors and model organisms. However, further advances that allow safe transgene delivery and stable expression in human models are still required.

### Ongoing challenges

Despite advances in the understanding of the pathophysiology, the relative ease of access to clinical laboratory techniques, and recent improvements in treatment options (such as rFVIIa), many challenges remain in the care of patients with this rare disorder. Access to clinicians and laboratories with the required expertise and resources to diagnose and treat GT is difficult, especially in areas of the world with limited resources. Large-scale data analysis and clinical research are problematic given the limited numbers of patients available and the highly constrained funding for rare disorders. Initiatives through the World Federation of Hemophilia for clinical care and multinational collaborations including the Glanzmann Thrombasthenia Registry for analysis of clinical outcomes, and sharing of molecular data through creation of publicly accessible databases have taken important steps in bridging the gaps. However, more investment is needed to guarantee timely access to quality care, until a true cure for the disease is developed.