

Novel Mutations in the GPIIb and GPIIIa Genes in Glanzmann Thrombasthenia

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Keywords

Glanzmann thrombasthenia · $\alpha_{IIb}\beta_3$ ·
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Summary

Background: Glanzmann thrombasthenia (GT) is an inherited autosomal recessive platelet disorder characterized by a complete or partial lack, or mutation, of the GPIIb/IIIa complex (integrin $\alpha_{IIb}\beta_3$) on the thrombocytes' surface, leading to a severe bleeding syndrome. **Material and Methods:** Molecular genetic analysis was performed in patients with suspected GT. The aim of the present study was the identification of new natural variants, their impact on platelet function, and their relation to the risk of bleeding. **Results:** Expression of the platelet integrin $\alpha_{IIb}\beta_3$ was determined by flow cytometry. Mutations were identified through sequencing of cDNA and genomic DNA. In addition, platelet function studies (PAC-binding, aggregations) were implemented. The study included 25 patients revealing 13 mutations (GPIIb: n = 9; GPIIIa: n = 4). Two of the 13 mutations were previously described (T207I; L214P). The remaining mutations have not been published yet, whereas 1 mutation in 2 unrelated families was identical (3062 T→C). **Conclusion:** All patients with less than 25% of present $\alpha_{IIb}\beta_3$ have a medical history of bleeding.

Schlüsselwörter

Thrombasthenia Glanzmann · $\alpha_{IIb}\beta_3$ ·
Molekulargenetische Analyse ·
Plättchenfunktionsuntersuchungen

Zusammenfassung

Hintergrund: Die Thrombasthenie Glanzmann (GT) ist eine angeborene, autosomal rezessiv vererbte Störung der Plättchenfunktion, die durch einen völligen oder partiellen Mangel bzw. eine Mutation des GPIIb/IIIa-Komplexes (Integrin $\alpha_{IIb}\beta_3$) auf der Thrombozytenmembran verursacht wird und zu einer schweren Blutungsneigung führt. **Material und Methoden:** In der vorliegenden Studie wurden bei Patienten mit Verdacht auf GT molekulargenetische Analysen durchgeführt. Ziel der Studie war die Aufklärung neuer natürlicher Varianten, deren Einfluss auf die Funktionsfähigkeit der Thrombozyten und ihre Relation zum Blutungsrisiko. Die Expression des GPIIb/IIIa-Komplexes auf Thrombozyten wurde mittels Durchflusszytometrie überprüft. Die Identifikation der Mutationen erfolgte über die Sequenzierung von cDNA und genomischer DNA. Weiterhin wurden Plättchenfunktionsuntersuchungen (PAC-Messung, Aggregationen) durchgeführt. **Ergebnisse:** Es wurden 25 Patienten untersucht, insgesamt wurden 13 Mutationen identifiziert (GPIIb: n = 9; GPIIIa: n = 4). Zwei dieser Mutationen wurden bisher beschrieben (T207I; L214P). Die übrigen Mutationen waren bisher noch nicht bekannt, wobei eine Mutation bei zwei nicht verwandten Patienten detektiert wurde (3062 T→C). **Schlussfolgerung:** Patienten mit einem Rezeptorbesatz unter 25% hatten positive Blutungsanamnesen.

Introduction

Glanzmann thrombasthenia (GT) is a rare, recessively inherited, bleeding syndrome affecting the megakaryocyte lineage [1, 2]. Affected patients exhibit a lifelong moderate to severe bleeding tendency with mainly mucocutaneous bleeding. The thrombasthenic phenotype is linked to quantitative and/or qualitative abnormalities in the platelet fibrinogen receptor, the $\alpha_{IIb}\beta_3$ integrin (glycoprotein (GP) IIb/IIIa, CD41/CD61), which mediates the incorporation of platelets into an aggregate or thrombus at sites of vessel injury [3]. Laboratory parameters in severe GT show normal platelet count, prolonged bleeding time, absence of platelet aggregation in response to multiple physiologic agonists, and impaired adhesion to siliconized glass. The more common heterozygous GT patients show almost normal or slightly diminished platelet aggregation.

The α_{IIb} (ITGA2B) and β_3 (ITGB3) genes are located on chromosome 17 (q21–22). ITGA2B spans 17 kb and comprises 30 exons, ITGB3 spans 46 kb and has 15 exons. The fibrinogen binding site of the receptor complex seems to be a discontinuous surface made up of various regions of both subunits. Fibrinogen binding to activated $\alpha_{IIb}\beta_3$ induces further sequential conformational changes in the receptor complex, inducing exposition of cryptic ligand-binding sites with outside-in signal transduction processes and cytoskeletal rearrangement [4]. The exact course of events is not known but various regions in the α_{IIb} and β_3 subunits are known to be involved in fibrinogen binding [5–12].

GT is classified into 3 subtypes, depending on the level of present $\alpha_{IIb}\beta_3$. Patients with type I or classical GT are homozygous for the disease and have a virtual absence of $\alpha_{IIb}\beta_3$ (<5% of normal level). Type II GT patients can have up to 25% of the normal level. In the variant type, $\alpha_{IIb}\beta_3$ levels are near normal but functionally impaired, leading to defective binding of fibrinogen [13]. Heterozygotes for type I GT have approximately 50% of the normal level of platelet $\alpha_{IIb}\beta_3$, without bleeding problems. Since the cloning of the α_{IIb} and β_3 genes, a number of mutations have been found to be associated with GT. A continually updated database is available on the Internet (sinaicentral.mssm.edu/intranet/research/glanzmann); it currently lists 103 records of mutations in the α_{IIb} and 68 records of mutations in the β_3 gene. The types of mutations identified in both genes include minor and major deletions, insertions, inversions, and mostly point mutations. The molecular and functional characterization of many of them has provided important information about the biosynthesis and structure-function relationships of the $\alpha_{IIb}\beta_3$ complex as well as the biology of other molecules of the integrin family [14, 15].

Most of the documented single nucleotide substitutions are located in the coding sequence and cause missense or nonsense substitutions at the amino acid level, producing either normal-sized non-functional or truncated proteins [16–20].

Splice site defects are also widespread, and mutations that alter mRNA splicing are frequently nonsense mutations [21, 22] or mutations directly affecting the standard consensus splicing signals, and typically lead to skipping of the neighboring exon [23]. By and large, mutations are specific for each family. GT occurs in high frequency in certain ethnic populations with an increased incidence of consanguinity, such as Indians, Iranians, Iraqi Jews, Palestinian and Jordanian Arabs, and French Gypsies [24–27]. To elucidate the molecular basis of GT, we investigated 25 GT patients for gene mutations within α_{IIb} and β_3 .

Material and Methods

Study Subjects

25 Patients, belonging to 12 unrelated families, with a diagnosis of GT, and their first-degree relatives if available were the subjects of the study (table 1). The investigated patients had different genetic backgrounds (Caucasian: n = 13; Asian: n = 9; African: n = 3). GT was diagnosed on the basis of clinical and hematologic parameters. The patients' phenotypes and genotypes were studied to perform carrier studies in their families. Bleeding symptoms were evaluated by examining available hospital records. Mild bleeders were defined as those who had minor symptoms, such as epistaxis or gum bleeds, and moderate bleeders suffered, in addition, from bleeding complications after surgery and trauma. Severe bleeders were defined as those with a history of spontaneous or life-threatening hemorrhages, such as gastrointestinal bleeding, or who had repeated episodes requiring platelet transfusion. Blood samples for the studies described below were taken from the patient after informed consent was obtained. The study was approved by the Ethics Committee of the Landesärztekammer Hessen.

Blood Sampling and Processing

Blood was taken in citrated buffer or in ethylenediaminetetraacetic acid (EDTA) from the patients. Citrated platelet-rich plasma (PRP) was obtained by centrifugation at $180 \times g$ for 10 min, platelet-poor plasma (PPP) by centrifugation at $2,500 \times g$ for 20 min. PRP was used for flow cytometry, platelet aggregation, and binding experiments. DNA extraction from white blood cells was carried out in blood samples containing EDTA.

Flow Cytometric Analysis of Platelets

These studies were carried out as recently reported with some modifications [28]. Monoclonal antibodies (mabs) against CD42a (clone SZ1), CD42b (clone SZ2), $\alpha_{IIb}\beta_3$ (clone P2), CD36 (clone FAG.152), and mouse IgG isotypes were purchased from Beckman Coulter (Krefeld, Germany). Mab PAC-1, specific for activated $\alpha_{IIb}\beta_3$, was obtained from BD Bioscience (Heidelberg, Germany). To obtain information relating to the receptor count, 200 μ l of PRP were fixed using 2% formaldehyde in PBS at room temperature for at least 1 h. The cells were sedimented by centrifugation at $600 \times g$ for 3 min and resuspended in 200 μ l HEPES buffer (20 mol/l HEPES, 150 mol/l NaCl with 60 mg/ml bovine serum albumin (Sigma-Aldrich, Steinheim, Germany); pH 7.5). Aliquots of platelet suspension were then incubated for 30 min in the dark with appropriate fluorescence-marked monoclonal antibodies. The samples were washed (2,400 rpm, 3 min) using CellWash (BD Bioscience), and resuspended in 400 μ l of CellWash. Fluorescence-labeled isotype matched IgG antibodies were used as negative control. Fluorescence intensity was measured with a FACScan flow cytometer and analyzed with CellQuest 3.1 software (BD Bioscience) which allowed the parallel measurement of 3 different fluorescences. Quantum fluorescence mi-

Table 1. Clinical characteristics and flow cytometry analysis in GT patients

Patient (sex)	Bleeding severity	Bleeding time, min	CD41 anti- α IIB, %	CD42a anti-GPIbIX, %	CD42b anti-GPIb α , %
TA (M) ⁺	severe	33	<1	99	105
TG (M) ⁺	severe	> 30	<1	107	111
TN (F) [°]	mild	9	90	109	110
TF (M) [°]	none	6.5	78	94	93
GE (F)	severe	21	<1	108	87
CM (F) [°]	mild		58	99	93
SB (F)	moderate	>15	23	120	125
YF (M)	moderate		3	n.a.	73
TE (M)	severe		<1	110	122
GI (F) [°]	mild		97	102	103
TA (M) [°]	mild		90	99	109
DZ (F)			0	103	110
LN (F)			<1	98	96
LH (F) [°]			67	130	129
LB (M) [°]			82	101	104
HM (M)	moderate		0	normal	normal
HS (F) [°]			n.a.	n.a.	n.a.
HA (M) [°]			n.a.	n.a.	n.a.
OK (F)	severe	>30	4	101	114
MM (M)	severe		<1	98	97
MU (F) [°]			82	94	86
WC (M)	severe		0	119	113
WV (F) [°]			68	121	113
WA (M) [°]			78	101	84
PJ (M)	severe		<1	normal	normal

F = Female; M = male; n.a. = not available; ⁺ = siblings; [°] = parents.

crobeads (Calibrite beads; BD Bioscience) were used each day for standardization of the instrument settings. To study platelet GPIIb/IIIa activation and fibrinogen binding by flow cytometry, platelets in PRP were activated with either adenosine diphosphate (ADP, 10 μ mol/l) or thrombin receptor-activating peptide (TRAP-6, 10 μ mol/l) in the presence of saturating concentrations of FITC-conjugated PAC-1 (directed against activated $\alpha_{IIb}\beta_3$) or anti-fibrinogen mab. Following incubation for 15 min at room temperature, platelets were fixed for 15 min, washed with CellWash, and analyzed in a final volume of 400 μ l for cell-bound fluorescence in the flow cytometer. The background was measured using FITC-conjugated anti-mouse IgG.

Light Transmission Aggregometry

Light transmission aggregometry (LTA) was performed on the ATRACT 4S Plus aggregometer (LABiTec, Ahrensburg, Germany). Platelet counts were not adjusted, and the baseline optical density was set with PPP. The used inducers were ADP (final concentrations: 2.5, 5, 10 μ mol/l), collagen (0.5, 1, and 2 μ g/ml; Horm[®] Nycomed Austria GmbH, Linz, Austria), epinephrine (10 μ mol/l final concentration), and arachidonic acid (0.75 mol/l final concentration). Optical density changes were recorded photoelectrically for 7 min as platelets began to aggregate. The aggregation response is given as percentage of maximal light transmission (Amax).

DNA and RNA Extraction

DNA isolation from whole blood was carried out by using the QIAamp DNA Blood mini kit (Qiagen, Hilden, Germany) according to the manufacturer's instructions. Total RNA in platelets was isolated from PRP as previously described [28].

Amplification of Genomic DNA by PCR and Synthesis of cDNA

Amplification of all individual exons of the α_{IIb} and β_3 genes, including splicing sites, was performed by using primers located in the flanking intron regions. Table 2 provides information about the length and location of all used primers. According to the different primers, polymerase chain reactions (PCRs) were performed using different annealing temperatures varying from 50 to 62 °C for 30 s to optimize the amplification products. Extension was performed at 72 °C for 60 s using Hot Start Taq Plus polymerase (Qiagen) for all exons. The amplification contained 35 cycles and was followed by a 5-min final extension step at 72 °C. Amplified DNA fragments were purified using QIAquick PCR Purification kit (Qiagen) according to the manufacturer's instructions. Platelet mRNA was reverse transcribed using the cDNA Synthesis System Plus (Amersham International, Little Chalfont, UK) as previously described [28].

Nucleotide Sequence Analysis

Cycle sequencing of the entire α_{IIb} and β_3 DNA was accomplished by using the primers from table 2 and the BigDye[®] Terminator v1.1 Cycle Sequencing kit (Applied Biosystems, Foster City, CA, USA) according to the manufacturer's instructions. cDNA sequencing of α_{IIb} and β_3 was performed with overlapping primers (sequences not shown). The cycle sequencing products were purified by using DyeEx 2.0 Spin kit (Qiagen) pursuant to the manufacturer's instructions. Analysis of the sequencing products was executed in ABI Prism 310 (Applied Biosystems) sequencer utilizing Performance Optimized Polymer 6 (Applied Biosystems) and ABI Prism Genescan software (Applied Biosystems). The regions containing the mutation were sequenced 3 times in both directions.

Table 2. List of primers used in the present study

Exon	Sense location, bp	Sequence	Antisense location, bp	Sequence
<i>αIIb</i>				
1	(-)-79-(-)58	GCAAGTACTTGGGGTTCCAG	307-287	CGCAAATGGGAAACTCGAAGC
2	3324-3343	GCCTGGGATACGCTGGAATC	3547-3526	GCAGTCCACGTCCCTCTGAC
3	3519-3538	AAGGTTGGGGTCAGAGGGAC	3731-3710	AGTGTCCCTGCCCCGATTG
4	3739-3758	AGGAGGAGCCCAAGTCTCGC	4054-4033	CAGATCCAAAGCAAGGGCTG
5	4127-4145	CTCCCTCACCCCTGGGCTG	4294-4273	GGAAGGGAAGTCTGAGGGT
6	4220-4239	GGCGAGTAGGGAGCAAAGC	4378-4359	GCGAGAAGGGAGGGAGGTGT
7	4387-4406	GAGACCGCTTTGGGCTTCAC	4598-4579	TGACCCTCGGGGTGCTGGAA
8	4886-4905	CCCTGGAAAAGACTAATTTG	5034-5015	AGATTTCAGGAAGGCGCTCC
9	5097-5166	AAGTGGGTAGGTTCTAAGGC	5243-5224	AGAAGTGGGATAAGGGGCTT
10	5307-5326	CCTGGAGTGGGAGGTTGCTT	5481-5461	TTATTCTGAAGTCTCAGTTCC
11	5506-5525	CTTAAAGAGGATGCTTGTC	5656-5637	ATTTGGGACCCAACTGGGTA
12	5767-5786	GTCCAGTCCCATGTAACCAC	6048-6029	CCTCTGCAGCAAGTAGGGCT
13	8409-8428	AACAATCAGCCACTTCCTTT	8662-8643	CTTGGGCATTTCTAGCTGGA
14	8826-8845	ATCGCCAATTCTGACCCATT	8962-8943	GCCTCCTCTGATGGCACAG
15	8981-9000	AATGGCAAGCCTACCCCATC	9163-9144	GAGGTCCCAGATCCTTTAAG
16	9145-9183	AGAAAGGCTCCAACCCCTGA	9300-9281	TCCAAGCCCACCTCCCTCCT
17	9317-9336	CCAGGGAGGTCCTGACTCTT	9559-9540	GTCCCAGTGGGTAAGTTCTAC
18	9658-9677	TAGCAAGATGGCCTGACTCT	9869-9849	ATTTGCTATCAGGGGTCTCTG
19	10730-10750	CCAAGTACCCGCACACCCC	10918-10899	GCAGGTATGATAGGCAGAAA
20	10962-10981	CCAGGCTCCCTGGCTTCACT	11179-11160	GGTCCCAGGACTGTTCCCA
21	11681-11700	TCTGTAATTTCTTTCTTGGA	11844-11825	TGGTGGAGACCCGGTACCAC
22	12382-12401	ACTTGGGCAGTGACCTTGGC	12537-12518	GGGGAAGGGTGGTGGGTAGG
23	13084-13103	CTGGCCCTGTTTCTCCTCAT	13233-13214	AATGCCATCTCCCTTCTCCA
24	13287-13306	ATCACACTCTCTGGGGGT	13471-13452	AGTTCTGAGGACCCGCTCAC
25	13484-13504	TTAAGTCCCCACACCTGCC	13707-13688	GCCTTCCCAGGTCTTTCTTC
26 (1)	13719-13738	AACCACCGGGGCACCTCTGT	13974-13955	ATCCCCTCTGCCCGTGGGT
26 (2)	13755-13774	GAGACCTGGGCCTGACCACT	13975-13955	ATCCCCTCTGCCCGTGGGT
27	14336-14355	TGGGGGATGATGGGGTGATG	14561-14542	AAAGTGGTCCCCGCCAGAA
28	14711-14730	TGGGTGTAAACGGCTTTCAA	14881-14862	AGGACTGGTCTCTGCTCCAT
29	14988-15007	AAGGCAGGTGTCAAGGTGA	15207-15188	AGGGCAGCGCAAGCCTGTG
30 (1)	17023-17042	TGTGCTCTGGGGCCAGCAAA	17385-17366	TCCAAAACATTCCTTCGGTC
30 (2)	16975-16995	TTCTGCGCTGGTCCAGGGAGG	17333-17313	GCTCAGCATGAGGGCTCAGTC
<i>β3</i>				
1	(-)-113-(-)93	CGTTGCGTCCCACCCACCGCG	151-131	CGCGGTGGGTGGGACGCAACG
2	20490-20510	AGGATGAGGCAGGCAAGTACC	29752-20694	CTTCATGGGTCTGTTTCTGCAC
3	29427-29447	CTGCAGGAGGTAGAGAGTCGC	29752-29732	CCTCCACCTTGTGCTCTATGC
4	30523-30543	CTGCTTATTCAATCTTGGTGG	30898-30878	GAACCAGGACTTGGACCTTCC
5	32344-32363	CATGCTGCCTTTTCCATGAAG	32637-32617	CCCAGAGTTGAGTCACTTCT
6	33141-33161	TCTTACCAGTGACATGGCTG	33431-33411	CTAGATTAGGGCAACCCTCAC
7	35760-35780	CTTTAGACCTTAGAGATGTAC	35989-35969	TGCTAGCACAGGCTGGCTACC
8	36232-36252	CATCTCAGGACTCTCAGTGGG	36471-36451	AAGCCAAATGAGGTGGACTCTCCGC
9	37035-37059	GCATTTCCCGTTTCTTTCAGTTCA	37269-37246	CACAGATGCTCCAGGACAAAGGCC
10	38227-38252	AGTGTTAACTGGGCCAACTGTGTCT	38773-38749	TCCCAGTGGTTGCAGGTATATGAGGG
10 (intra)	38408-38432	AGGCCAAAGCTGAACCTAATAGCC		-
11	45387-45410	GCTGCCATGGAGTGGAGCTCTCG	45720-45698	GATCCTCTCCTACCTCCCAGCCT
12	46536-46556	TGCATGGAGATCAGAGCTGGA	46758-46736	GGTCTGGGATTGAATTAGAAAC
13	48793-48815	GGTTTGCAGTGGTCCCATCTTCC	49047-49025	ATTCTGCTTAAACATGGTTCTCC
14	53552-53573	AGGTTCAAGTGACTCCTGCTTCA	53839-53816	TGCTGCTTCCACTTAAGAGTCTC
15	56218-56240	TTCAGGGTAGGGAAGGACTTAAG	56410-56389	ACATGATGGCAGGGACTCCTGC

Table 3. Mutations in the α_{11b} gene identified in GT patients

N	Mutation	Location	Patient	Genotype	Phenotype	Effect
1	620C→T	exon 5	OK	homozygous	missense	T207I
2	641T→C	exon 6	MM MU	homozygous heterozygous	missense missense	L214P L214P
3	1754T→C	exon 17	DZ	homozygous	splicing site	alternative splicing
4	1878G→C	exon 18	HM HS HA	homozygous heterozygous heterozygous	splicing site splicing site splicing site	alternative splicing alternative splicing alternative splicing
5	2051T→G	exon 20	GE CM	heterozygous heterozygous	missense missense	L684R L684R
6	2232Gins	exon 22	WC WV	heterozygous heterozygous	ins. out of frame ins. out of frame	frameshift frameshift
7	3060G→A	exon 29	TA TG TN TF	homozygous homozygous heterozygous heterozygous	splicing site splicing site splicing site splicing site	alternative splicing alternative splicing alternative splicing alternative splicing
8	3062T→C	exon 29	LN LH LB	homozygous heterozygous heterozygous	splicing site splicing site splicing site	alternative splicing alternative splicing alternative splicing
9	3062T→C	exon 29	SB	heterozygous	splicing site	alternative splicing

N = Consecutive numbering of the found mutations; ins. = insertion.

Table 4. Mutations in the β_3 gene identified in GT patients

N	Mutation	Location	Patient	Genotype	Phenotype	Effect
1	856G→A	exon 6	TE TA	heterozygous heterozygous	missense missense	G286R G286R
2	1129Ains	exon 9	TE GI	heterozygous heterozygous	ins. out of frame ins. out of frame	frameshift frameshift
3	1550Gdel	exon 10	YF	homozygous	del. out of frame	frameshift
4	2068delGT	exon 13	PJ	homozygous	del. out of frame	premature termination

N = Consecutive numbering of the found mutations; ins. = insertion; del. = deletion.

Results

Molecular Analysis of α_{11b} and β_3 Transcripts

To determine the molecular basis underlying the abnormalities of $\alpha_{11b}\beta_3$ receptor in the patients, full-length α_{11b} and β_3 (1–2 kb fragments) from the patients' platelets were amplified using PCR and sequenced by a modified Sanger method. A list of mutations identified within the α_{11b} and β_3 genes is shown in tables 3 and 4. We found 8 different mutations of the α_{11b} gene in 20 patients and 4 mutations of the β_3 gene in 5 patients. Four missense mutations, 4 splicing site mutations, 2 insertions of a single nucleotide, 1 deletion of a single nucleotide, and a deletion of 2 nucleotides were found. Overall, 9 patients were homozygous for a mutation in the α_{11b} gene ($n = 7$) or in the β_3 gene ($n = 2$), whereas only 1 patient was

found to be compound heterozygous with both mutations located in the β_3 gene (table 5). In 14 patients, only a mutation in the heterozygous state was found and no additional gene defect was identified. We assume that 3 out of those 14 heterozygotes are compound heterozygous (GE, SB, WC), however the second mutations could not yet be located. To exclude that the mutations found in our patients represent common polymorphisms, 50 randomly chosen samples from different ethnic or geographic backgrounds were screened for all mutations. None of these subjects were found to carry one of the mutations.

Missense Mutations ($n = 4$)

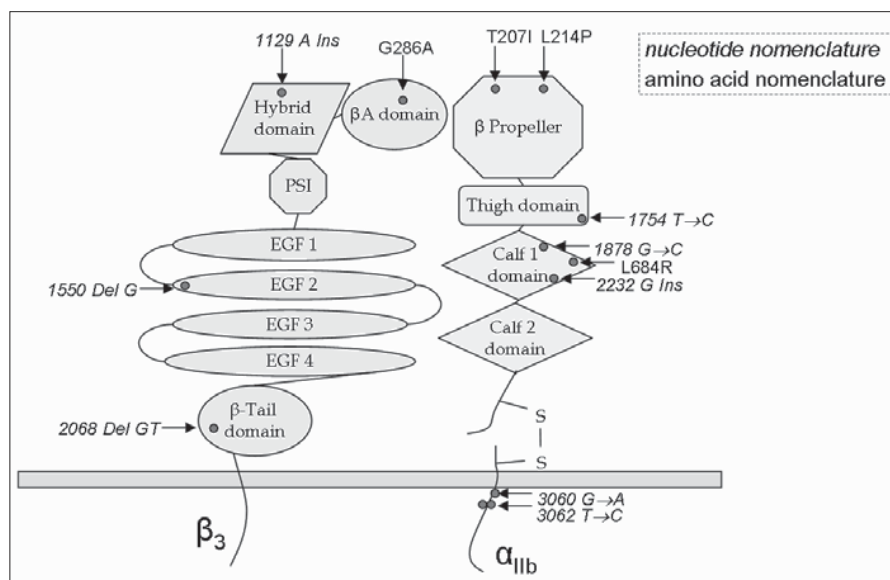
Two missense mutations were detected within the α_{11b} gene (table 3). In exon 5, a homozygous Thr 207 to Ile (ACT→ATT)

Table 5. Genetic findings in GT patients

Patient (sex)	Genotype	Gene	First mutation		Second mutation	
			location	effect	location	effect
OK (F)	homozygous	α_{11b}	exon 5	T207I	–	–
MM (M)	homozygous	α_{11b}	exon 6	L214P	–	–
MU (F)	heterozygous	α_{11b}	exon 6	L214P	–	–
DZ (F)	homozygous	α_{11b}	exon 17	splicing site	–	–
HM (M)	homozygous	α_{11b}	exon 18	Q626H	–	–
HS (F)	heterozygous	α_{11b}	exon 18	Q626H	–	–
HA (M)	heterozygous	α_{11b}	exon 18	Q626H	–	–
TA (M)	homozygous	α_{11b}	exon 29	splicing site	–	–
TG (M)	homozygous	α_{11b}	exon 29	splicing site	–	–
TN (F)	heterozygous	α_{11b}	exon 29	splicing site	–	–
TF (M)	heterozygous	α_{11b}	exon 29	splicing site	–	–
LN (F)	homozygous	α_{11b}	exon 29	splicing site	–	–
LH (F)	heterozygous	α_{11b}	exon 29	splicing site	–	–
LB (M)	heterozygous	α_{11b}	exon 29	splicing site	–	–
SB (F)	(compound) heterozygous	α_{11b}	exon 29	splicing site	unknown	–
GE (F)	(compound) heterozygous	α_{11b}	exon 20	L684R	unknown	–
CM (F)	heterozygous	α_{11b}	exon 20	L684R	–	–
WC (M)	(compound) heterozygous	α_{11b}	exon 22	ins. out of frame	unknown	–
WV (F)	heterozygous	α_{11b}	exon 22	ins. out of frame	–	–
WA (M)	heterozygous	α_{11b}	unknown	unknown	–	–
TE (M)	compound heterozygous	β_3	exon 6	G286A	exon 9	frameshift
TA (M)	heterozygous	β_3	exon 5	G286A	–	–
GI (F)	heterozygous	β_3	exon 9	frameshift	–	–
YF (M)	homozygous	β_3	exon 10	G517DelG frameshift	–	–
PJ (M)	homozygous	β_3	exon 13	premature termination	–	–

F = Female; M = male; ins. = insertion.

Fig. 1. Schematic illustration of $\alpha_{11b}\beta_3$. Shown is the distribution of all mutations (red dots) onto the different domains of α_{11b} and β_3 . Nucleotide nomenclature is italicized and refers to splicing site and frameshift mutations whereas the amino acid nomenclature is non-italicized referring to point mutations causing an amino acid exchange.



substitution was identified, and in exon 6, a homozygous T-to-C transition leading to a Leu 214 to Pro (CTT→CCT) substitution. A heterozygous Gly 286 (GGA→AGA) to Arg mutation was detected within the β_3 gene (table 4) of patient TE. Analysis of the parents' DNA showed that the missense mutation was inherited from the father (TA). All detected

missense mutations are non-conservative substitutions that involve strictly conserved residues among vertebrates in the α_{11b} as well as in the β_3 gene. The Thr 207 to Ile substitution allows for a non-polar instead of a polar and aliphatic amino acid in the β -propeller domain of the N-terminus part of the mature protein (fig. 1). The Leu 214 to Pro substitution (Patient MM)

Table 6. Flow cytometric analysis and agonist-induced platelet aggregation in GT patients

Patient	FACS analysis			Platelet aggregation		
	CD41 anti- α_{IIb} , %	PAC-1, %		ADP (5 μ mol/l)	collagen (1 μ g/ml)	ristocetin (1.3 mg/ml)
		10 μ mol/l ADP	10 μ mol/l TRAP			
TA (M)	<1	<1	<1	∅	∅	N
TG (M)	<1	<1	<1	∅	∅	N
TN (F)	90	37	55	(▼)	(▼)	N
TF (M)	78	38	25	(▼)	(▼)	N
GE (F)	<1	<1	<1	∅	∅	N
CM (F)	58	44	51	(▼)	▼	N
SB (F)	23	13	12	▼	(▼)	N
YF (M)	3	n.a.	n.a.	n.a.	n.a.	n.a.
TE (M)	<1	<1	<1	∅	∅	N
DZ (F)	0	n.a.	n.a.	n.a.	n.a.	n.a.
LN (F)	<1	<1	n.a.	∅	∅	N
HM (M)	0	<1	<1	n.a.	n.a.	n.a.
OK (F)	4	<1	<1	∅	∅	N
MM (M)	<1	<1	<1	▼	▼	(▼)
MU (F)	82	57	42	n	N	N
WC (M)	0	<1	<1	∅	∅	N
WV (F)	68	35	25	▼	▼	N
WA (M)	78	66	22	∅	∅	(▼)
PJ (M)	<1	n.a.	n.a.	∅	∅	n

N = Normal; ∅ = absent; ▼ = inhibited or reduced; (▼) = slightly inhibited; n.a. = not analyzed.

occurs also in the β -propeller domain and involves substitution from an aliphatic amino acid to heterocyclic amino acid. In the β_3 gene, the heterozygous missense mutation (Gly 286 Arg) lies, according to the known crystal structure [29], in a highly conserved position within a proposed intramolecular disulfide loop, between Cys 232 and Cys 273 [30]. Furthermore, a heterozygous Leu 684 (CTG→CGG) to Arginin substitution was found in 2 related patients (GE, CM).

Deletions (n = 2)

Two mutations were detected within the β_3 gene (table 3). A homozygous deletion at Gly 517 in exon 10 (1550delG) leads to changes of the corresponding amino acids of exon 10. In addition, a homozygous deletion of 2 nucleotides in exon 13 (2068delGT) is predicted to lead to changes of the corresponding amino acids of exon 13 and to a premature stop codon at position 696.

Insertions (n = 2)

Two mutations were detected within the α_{IIb} gene (table 3). A heterozygous insertion at Ala744 (2232Gins) in exon 22 is predicted to lead to changes in the corresponding amino acids, and was detected in 2 affected blood relatives (WC, WV). One insertion mutation was located within the β_3 gene (table 3), and was found in the heterozygous state at Ile377 in exon 9 (1129Ains) in 2 related patients. This mutation would lead to changes in the corresponding amino acids encoded by the subsequent portion of exon 9.

Splicing Site Mutations (n = 4)

All splicing site mutations were found in the α_{IIb} gene (table 3). Sequencing of exon/intron regions using genomic DNA revealed a homozygous splicing donor site mutation (3060G→A) in exon 29 in 2 siblings (TA, TG). Analysis of the entire cDNA showed a homozygous deletion of exon 29 in the α_{IIb} subunit, the putative transmembrane region. As expected, both parents (TN, TF) were heterozygous carriers of that mutation. A further homozygous splicing donor site mutation (3062T→C) was found in exon 29 (LN) and as heterozygous condition in the patients' parents (LH, LB). In addition, this splicing site mutation (3062T→C) was also detected, as heterozygous condition, in an additional patient (SB). Although no blood relationship was reported, these 2 patients were from the same region in Germany. Moreover, homozygous splicing donor site mutations in exon 17 (1754T→C) (DZ) or exon 18 (1878G→C) (HM) were identified.

Polymorphisms (n = 5)

Relating to GT, there are various single nucleotide polymorphisms known in α_{IIb} and β_3 , which can lead to an amino acid exchange but have no impact on the function of the thrombocytes in terms of aggregation or adhesion ability. The investigation of all patients revealed previously described polymorphisms in homozygous and heterozygous forms concerning several patients (DZ, WC, TE, YF, PJ). The polymorphism Ile843Ser in the α_{IIb} subunit could be identified in 2 patients (WC: heterozygous; TE: homozygous). The β_3 subunit contains

more polymorphisms. Patients DZ and YF show the Leu33Pro polymorphism in the homozygous and patient SB in the heterozygous form. Furthermore, the polymorphisms Val381Val (TE: heterozygous; PJ: homozygous) and Arg489Gln (TE: heterozygous; PJ: homozygous) could be ascertained. The possibility of a founder effect for the mutations V585G, 2232Gins, V1021A, G286A + 1129Ains, 1550delG, and 2068delGT seems unlikely but cannot be excluded because the patients carrying polymorphisms all seem to suffer from GT type I, except for one. However, a coherency is not established.

Platelet Function Studies

The results of flow cytometric and platelet aggregation studies are summarized in table 6. As expected, platelets from patients with type I GT (0–5% of $\alpha_{IIb}\beta_3$ on platelets: TA, TG, GE, TE, LN, HM, OK, MM, WC, PJ) were unable to aggregate in response to ADP (5 $\mu\text{mol/l}$) and collagen (1 $\mu\text{g/ml}$). Platelets of these patients failed to aggregate even with higher concentrations of ADP (10 $\mu\text{mol/l}$) and collagen (2 $\mu\text{g/ml}$) (data not shown). Furthermore, analysis of $\alpha_{IIb}\beta_3$ activation-dependent epitope using flow cytometry showed that mab PAC-1 was unable to bind after activation with TRAP or ADP (table 6). However, a larger variability was seen between the different tests in patients with >20% of $\alpha_{IIb}\beta_3$ on platelets. For example, the heterozygous parents of TE and TG had a normal amount of $\alpha_{IIb}\beta_3$ and slightly inhibited platelet aggregation but showed distinct inhibition of PAC-1 binding. The parents of patient WC also showed normal $\alpha_{IIb}\beta_3$ expression on platelets. PAC-1 binding was markedly decreased with a more pronounced effect for the mother's platelets. Surprisingly, the father's platelets were unable to aggregate in response to ADP and collagen. In the mother, we observed a markedly decreased platelet aggregation response to ADP and collagen. Patient SB had a residual $\alpha_{IIb}\beta_3$ surface expression of 23%. Functional analysis of $\alpha_{IIb}\beta_3$ revealed low PAC-1 binding capacity of about 10% after activation with ADP or TRAP. These results indicate that both a quantitative and qualitative abnormality in the $\alpha_{IIb}\beta_3$ receptor exist in this patient.

Discussion

GT is a rare congenital bleeding syndrome which results in deficient platelet aggregation due to the absence or dysfunction of the fibrinogen receptor $\alpha_{IIb}\beta_3$. Due to the dimension and diversity of the occurring mutations in $\alpha_{IIb}\beta_3$, it is rather complex to identify mutations that are responsible for GT. In the present study, 10 novel and 2 previously described gene alterations that are known to cause GT were identified in 12 unrelated patients (table 3) and their first-degree relatives. To ascertain the mutations, we sequenced the coding regions of the α_{IIb} and β_3 subunits including the exon/intron boundaries. Our study revealed 4 point mutations which are all non-conserva-

tive substitutions in strictly conserved residues among vertebrates in both genes. Two of the point mutations in α_{IIb} , Thr 207 to Ile and Leu 214 to Pro, are located in the FG-GAP 3 of the β propeller. Leu 214 to Pro has already been described by Grimaldi et al. [31]. It has been shown that the FG-GAP is important for ligand binding, so the Leu 214 to Pro mutation disrupts the structural conformation and impairs the ligand-binding properties of the heterodimeric complex. Additionally, the mutation seems to confer susceptibility to proteolysis [31, 32]. Concerning the Thr 207 to Ile mutation, it has been described to impair fibrinogen binding and affect the expression of $\alpha_{IIb}\beta_3$ on the platelet surface [28]. Further, we found a point mutation at position 684 leading to a Leu to Arg substitution. This mutation is located in the calf-1 domain of exon 20, also in the α_{IIb} subunit. The calf-1 domain creates minor interfaces with the β_3 subunit [33]. Therefore, conformational changes due to this amino acid substitution could impede contact between α_{IIb} and β_3 resulting in a failure of complex-formation. If the subunits do not form a complex, the protein subunits are not protected from intracellular proteolysis, and therefore they cannot be anchored in the cell membrane.

Gly 286 to Arg is located in the VWFA domain, also called βA domain, of the β_3 subunit. The βA domain is a ligand-binding head region of the β subunit. Conformational changes in VWFA have an impact on the ligand-binding activity, and ligand recognition also causes conformational changes transduced from this domain [34].

In 5 unrelated individuals and their relatives we detected splicing site defects. Three of those splicing site mutations are located in the splicing donor site in exon 29, and 2 of them (3062T→C; SB, LN) are even identical although no apparent consanguinity was reported. However, both patients were from the same region in Germany. Both of those splicing site mutations lead to a complete loss of exon 29, which was proven by cDNA sequencing. The mutations are located in the cytoplasmic domain proximal to the transmembrane region in the N-terminal area of the GFFKR motif. Regions proximal to the transmembrane domains of the α and β subunits contain highly conserved sequences, such as the GFFKR motif referring to the α_{IIb} subunit, and are found to be important for maintaining the inactive low-affinity state of the integrin. It is confirmed that the cytoplasmic domains are essential for the formation of the heterodimer, and the GFFKR motif appears to be required for association of the α and β subunits [35, 36]. The 3062T→C substitution at position V1021 is located at only 1 amino acid prior to the start of the GFFKR motif and leads to an alternative splicing. Therefore, it could be assumed that due to the alternative splicing, N-terminal to the motif, the conformation of the protein is changed in such a way that it could be impossible for the above-mentioned motif to associate with the corresponding motif in the β_3 subunit.

The other two splicing site defects were detected in exons 17 and 18 of the α_{IIb} subunit. The 1754T→C (DZ) substitution situated in the splicing donor site of exon 17 is located in the thigh

domain of the α_{IIb} subunit. The thigh domain has substantial contact with the β -propeller. Losonczy et al. [37] previously suggested that this contact might be important for the proper orientation of the β -propeller to interact properly with the β_3 subunit. An alteration in this domain could therefore change the orientation of the β -propeller and inhibit complex formation with the β_3 subunit [37]. The homozygous splicing donor site mutation 1878G→C (HM) affecting exon 18 is situated in the calf-1 domain of the GPIIb subunit. The calf-1 and calf-2 domains form minor interfaces with GPIIIa [36]. Rosenberg et al. [33] formerly published that significant changes in the calf-1 and calf-2 domains do not avert complex formation of $\alpha_{IIb}\beta_3$ but affect its transport to the cell surface, probably by its retention and subsequent degradation in the endoplasmic reticulum [33]. We assume that due to the found splicing site defects, the affected exons are non-existent and the concerned domains are thereby sustainably influenced in their conformations.

In 4 GT patients, we found frameshift mutations caused by insertions or deletions. Three out of 4 are situated in the β_3 subunit. Insertions or deletions lead to a shift in the reading frame and often cause a premature stop codon resulting in premature termination of the protein or an abnormal protein product with an incorrect amino acid sequence.

In 1 patient, we could identify a heterozygous insertion of G at nucleotide position 2232. This frameshift does not result in a premature stop codon but the amino acid sequence is severely altered, probably resulting in an abnormal protein. Since this mutation is also situated in the calf-1 domain of the α_{IIb} , we assume once more that complex formation is impaired resulting in a premature intracellular proteolysis.

A compound heterozygous patient and his mother showed a heterozygous insertion of A in exon 9 at nucleotide position 1129. This insertion results in a stop codon at amino acid position 380, probably leading to premature termination. Here, the hybrid domain is affected.

Concerning frameshift mutations we could additionally

identify 2 patients with deletions within the β_3 subunit. A deletion of 1 nucleotide (1550delG) was detected in exon 10 of β_3 . This deletion is situated in the EGF-2 domain and does not lead to a premature stop codon, nevertheless we assume that the nearby located disulphide bond can not be formed due to a modified amino acid sequence. The EGF domains are cysteine-rich modules, containing multiple disulphide bonds. In particular, the EGF-2 region is wedged in the bent between integrin stalk and headpiece domains, poised to communicate structural rearrangements between the cytoplasmic domains and the ligand-binding headpiece [38]. Also located in β_3 , a 2-nucleotide deletion (2082delGT) was identified in exon 13, affecting the β -tail domain (β -TD). This mutation leads to premature termination of the translation at amino acid position 696 and therefore to a truncated protein.

Several of these patients were carrier of diverse polymorphisms (table 6). We cannot exclude founder effects of polymorphisms regarding the newly revealed mutations in α_{IIb} and β_3 but it seems improbable. Solely in 3 heterozygous patients, a second mutation forming the compound heterozygous status could not yet be detected. A genotype/phenotype correlation is only vaguely viable. Therefore, a profound genotype/phenotype coherence could not be implemented. However, we came to know that the patients with splicing site defects suffer from mild to moderate bleeding, whereas point mutations and insertions/deletions tendentially result in moderate to severe bleeding. In conclusion, our investigation of several patients, and in some cases including their first-degree relatives, revealed a number of different mutations within the α_{IIb} and β_3 subunits associated with causing GT. This vast number of different mutations indicates the genetic heterogeneity in the studied group as well as in GT in general.

Disclosure

The authors declared no conflict of interest.

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