Novel Sequence Variation Affects GPIbα in Post-diarrheal Hemolytic Uremic Syndrome

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Abstract

Background: Hemolytic uremic syndrome (HUS) is one of the major causes of renal failure in children. In most cases the disease is caused by infection with Shiga toxin-producing Escherichia coli (STEC) and preceded by diarrhea. Only in 15% of cases STEC infection leads to HUS. Genetic predisposition of a patient to develop HUS after STEC infection might play a role, but very few reports on this subject are available. We describe a novel missense mutation in the GP1BA gene encoding platelet-receptor glycoprotein Iba (GPIbα) in a severely affected HUS patient.

Methods: GP1BA was screened by Sanger sequencing. Binding of recombinant GPIbα and von Willebrand factor (VWF) fragments was analyzed using surface plasmon resonance (SPR). The hematological studies using patient blood were performed.

Results: The detected heterozygous mutation p.Pro46Leu is located in the proximity to one of the two GPIbα-VWF binding sites. SPR experiments show that the p.Pro46Leu leads to a small increase in GP Ibα-VWF binding (p<0.001). The hemostatic parameters of patient blood after recovery from HUS show normal values.

Conclusions: The described mutation affects GPIbα interaction with VWF in a mild gain-of-function manner and might have contributed to a prothrombotic state in the patient and to development of HUS.

Keywords: DNA change; GPIbα; Hemolytic uremic syndrome; Platelets; Von Willebrand factor

Introduction

The Hemolytic Uremic Syndrome (HUS) is one of the major causes of renal failure in children. It is characterized by hemolytic anemia, thrombocytopenia and acute renal failure [1]. More than 90% of cases (typical HUS) are preceded by watery or bloody diarrhea and are caused by an infection with Shiga toxin-producing Escherichia coli (STEC), which can be ingested with improperly cooked meat and contaminated vegetables and water [2]. The majority of typical HUS cases affects children and is associated with infection by O157:H7 STEC serotype. The STEC infection leads to typical HUS in approximately 15% of cases [3]. Interestingly, the outbreak in Germany in 2011 was caused by an unusual O104:H4 strain, which mostly affected adults [4]. In the intestine, HUS-causing STEC strains produce Shiga toxins, belonging to type 1 and/or type 2 [5,6]. The toxins pass the intestinal wall and enter the circulation; they are recognized by the glycolipid receptor globotriaosylceramide (Gb3) on the surface of various cell types. The exact mechanism of how exposure to Shiga toxins leads to HUS and the reason why glomerular endothelium of the kidney is most strongly affected in HUS is poorly understood. Shiga toxins are known to inhibit protein synthesis and are involved in the induction of apoptosis [7]. Complement activation by Shiga toxin on endothelial cells was shown in vitro, in a murine HUS model [8] and on platelet-leukocyte complexes and microparticles of HUS patients [9]. In vitro studies showed that Shiga toxin 2 binds complement factor H (CFH), thereby reducing CFH complement inhibiting activity at the cell surface [10]. Interestingly, 85% of individuals that are infected with STEC will not progress to HUS.

In atypical HUS (aHUS), which is not preceded by the STEC infection, 50-60% of patients carry predisposing changes in complement genes encoding CFH, complement factor I (CFI), membrane cofactor protein (MCP), C3, complement factor B (CFB) and thrombomodulin and/or autoantibodies to CFH [11]. Similar to aHUS, genetic predisposition might be important in typical HUS. A single study reported significantly different frequencies of the three missense single nucleotide polymorphisms (SNPs) in the group of typical HUS patients, compared to healthy controls [12]. In the GPIBA gene, which encodes platelet-receptor glycoprotein Iba (GPIbα), a higher frequency of the (human platelet antigen 2) HPA2 SNP was strongly associated with typical HUS. Furthermore, significantly different frequencies among HUS patients were reported for the two SNPs in genes encoding adipoocyte-derived leucine aminopeptidase and factor V. Another report described a single mutation in MCP that was found in a typical HUS patient and also in patients with aHUS, HELLP (hemolysis, elevated liver enzymes, and low platelets) syndrome, and glomerulonephritis with C3 deposits [13]. Gaining further insight into the genetic background in typical HUS is important as it could improve our understanding of pathogenesis of the disease and possibly provide new possibilities for the treatment of patients. Here we describe a novel sequence variation in the GPIBA gene found...
in a post-diarrheal (typical) HUS patient and provide evidence for its functional consequence.

**Methods**

**Genetic analysis**

Genomic DNA was isolated from peripheral blood leukocytes as described before [14]. Coding sequences of GP1BA, C3, CFH, CFI, MCP and CFB were amplified by PCR; primer sequences are available upon request. The amplimers were subjected to double-stranded DNA sequence analysis on an ABI 3130xl Genetic Analyzer (Applied Biosystems). Presence of the novel genetic change was analyzed among 192 ethnically-matched control subjects. To this end, commercially available Human Random Control DNA panels were obtained (Sigma-Aldrich).

**Autoantibodies to CFH**

Patient serum was tested for the presence of autoantibodies to CFH by means of ELISA as described before [15]. A test result was considered positive if obtained extinction value was more than twice the standard deviation above the mean value calculated from 16 controls. The assay was repeated three times; a positive control sample was taken in every measurement.

**Hematological analyses**

Ristocetin induced platelet agglutination was analyzed in Platelet Rich Plasma using 0.5, 0.6, 0.7, 0.8, 0.9 and 1.52 mg/ml of ristocetin. Agglutination was analyzed in a Chronolog Aggregometer. All other hematological analyses were performed according to the standard operational protocols used at the RUNMC; which are available upon request.

**Recombinant protein production**

GP Ibα residues (1-290) containing p.Asn37Gln and p.Asn175Gln mutations to remove N-glycosylation sites, preceded by a cystatin signal peptide and fused to a C-terminal Arg-(His)6 sequence, were cloned into a modified pUPE expression vector. The p.Pro46Leu sequence variation and p.Met255Val were expressed in HEK-293-EBNA1-S cells and purified as previously described [16]. Fully sulfated protein was concentrated to ~1 mg/ml in the gel filtration buffer (50 mM NaCl, 20 mM Tris/HCl pH 8.0).

Wild type von Willebrand factor A1 domain (VWF-A1) and VWF-A1 carrying the p.Arg1306Gln type 2B von Willebrand disease mutation were introduced using the Quick Change method. Proteins were expressed in HEK-293-ERNA1-S cells and purified as previously described [16]. Fully sulfated protein was concentrated to ~1 mg/ml in the gel filtration buffer (50 mM NaCl, 20 mM Tris/HCl pH 8.0).

**Surface plasmon resonance experiments**

Surface plasmon resonance (SPR) binding studies were performed on a Biacore T100 (GE Healthcare). Approximately 9500 response units (RU) of GP Ibα monoclonal antibody 2D4 were immobilized on a series S CM5 sensor chips by amine coupling as instructed by the supplier. A control channel was activated and blocked by using the amine coupling reagents in the absence of protein. Proteins were dialyzed to standard SPR buffer (150 mM NaCl, 0.005% (v/v) Tween-20 and 25 mM HEPES pH 7.4) and analyzed at 25°C. GP Ibα variants (wild type, p.Met255Val and p.Pro46Leu, each 150 nM) were injected for 60 seconds at 10 μl/min, followed by a 60 seconds association phase of wild type VWF-A1 (0–4000 nM) or p.Arg1306Gln VWF-A1 (0–1000 nM) and a 120 sec dissociation phase at 30 μl/min. Proteins were injected until binding equilibrium was reached. Between runs, the sensor chip was regenerated by injections of 50 mM triethylamine, 10 mM sodium formate pH 2.0 containing 150 mM NaCl and another injection of 50 mM triethylamine. Each interaction was analyzed at seven different VWF-A1 concentrations. Data evaluation was performed with BIAcore T100 evaluation software. Affinity constants were derived by steady state analysis. Kinetics analysis was not possible due to the instrument limitations.

**Results**

**Case report**

The female patient is the youngest child in a family with two children of healthy parents. In 1977, at the age of 6.5 years the patient developed HUS. The initial symptoms included diarrhea, extreme paleness, fatigue and vomiting. The laboratory findings demonstrated anemia, with presence of fragmentocytes, and high creatinine and urea serum values. The patient was started on peritoneal dialysis, which was performed six times; heparinization was used, but was stopped after an intestinal bleeding episode. A Scribner shunt was placed in the leg for hemodialysis. Within the two months after first symptoms the patient suffered two cerebral accidents, possibly due to hypertensive encephalopathy. The patient developed hypertension, which was stabilized by medication (Nepresol, Inderal, Catapresan).

At the age of nine years, the patient received a first cadaver kidney transplant. At the age of 17, the kidney graft was removed due to chronic rejection and the patient was referred for hemodialysis for three years. After that, at the age of 20, she received a second cadaver kidney transplant, which was removed 14 years later due to chronic rejection, during this period the patient also suffered from cytomegalovirus infection. Two years later the patient received a kidney graft from a living donor via cross-over transplantation program, where the mother donated a kidney to another patient. At the moment of the study, the patient had a stable kidney graft function for five years. The function of the remaining kidney of the mother is also normal. There was no HUS recurrence reported in any of the grafts.

As the patient in question was diagnosed with HUS in 1977, presence of the STEC infection was not investigated. The STEC involvement in HUS was only demonstrated in 1983 [18] therefore it was not possible to test this patient for the presence of STEC in the acute phase of the disease. Nevertheless, presence of diarrhea, the young age of the patient, and lack of recurrence in kidney grafts support the possibility that the patient had suffered from the STEC-associated typical HUS, rather than from the atypical form of HUS.

**Novel GP1BA alteration found in HUS patient**

The objective of this study was to investigate whether this severely affected patient carries genetic predisposition to the development of HUS. No DNA alterations were found in genes encoding aHUS-predisposing factors CFH, CFI, MCP, C3, CFB and thrombomodulin. Moreover, autoantibodies to CFH were not detected in the patient. Previously, a strong association between typical HUS and GP1BA SNP was described [12]. We, therefore, screened the coding sequence of GP1BA. A single novel heterozygous missense sequence variation c.137C>T (p.Pro46Leu) was identified (Figure 1a). This novel sequence variation was also found in the DNA of patient’s brother (16 month older than the patient) and her father that were both not affected by HUS. The patient did not carry the HPA2 polymorphism. The DNA change was neither described previously, nor found by us in 192
healthy control individuals. The variation affected an evolutionary conserved amino acid residue (Figure 1b) and was suggested to alter the GPIbα protein function by mutation analysis programs SIFT (score 0.00, http://sift.jcvi.org/) and PolyPhen-2 (possibly damaging, http://genetics.bwh.harvard.edu/pph2/). Therefore, the found DNA change was likely to affect GPIbα protein function and might have facilitated the development of HUS in the patient.

**Novel alteration enhances GPIbα interaction with VWF**

The identified novel DNA change affected an amino acid residue in the proximity of one of the two interaction sites between GPIbα N-terminal part and the VWF-A1 domain [16] (Figure 1c). Since formation of microthrombi in capillaries is a prominent feature of HUS, we hypothesized that the p.Pro46Leu change is prothrombotic. The missense alteration might have strengthened the GPIbα and VWF interaction and in this way contributed to the formation of microthrombi in HUS. We, therefore, investigated the impact of the novel sequence variation on VWF binding using purified recombinant proteins. The binding of the GPIbα N-terminal fragment to the VWF-A1 domain was measured by SPR. The p.Pro46Leu change in GPIbα led to a small increase in steady state affinity for the VWF-A1 domain (p<0.001) (Table 1). Approximately 10% decrease in dissociation constant (Kd) was observed for the interaction with wild type VWF-A1 as well as with a variant carrying the p.Arg1306Gln mutation that stabilizes the interaction and is associated with type 2B von Willebrand disease [16]. The increase in affinity for the p.Pro46Leu variant was much smaller than for the gain-of-function p.Met255Val change in GPIbα, which is associated with platelet-type von Willebrand disease and strengthens the interaction more than 30 fold (Table 1).

**Hematological findings**

The SPR experiments suggested a mild increase in GPIbα affinity to VWF. We further performed extensive hematological studies to determine whether hemostasis is affected in the patient. The routine hematological analysis revealed no abnormalities, except for elevated factor-VIII activity (Table 2). In particular, we were interested in
features that could indicate abnormal platelet function: mean platelet volume, associated with higher platelet activity, aggregation and prothrombotic potential; platelet activation markers (CD63, PAC-1, GP53 (CD63), ADP aggregation Normal NA) and in murine model [8].

Recent study demonstrated that *Listeria monocytogenes* binds to platelets in the blood in a manner dependent on GPIb and C3 in mice [23]. This finding might implicate GPIb in complement biology, which might also be relevant in HUS.

The brother and father of the patient carry the same genetic alteration in GPIBA but were never affected by HUS. In particular, the absence of HUS in the brother is interesting because this disease mostly affects children [2]. However, it is possible that the brother was not exposed to STEC.

Various mutations in GPIba were previously described in relation to human disease. Gain-of-function mutations that increase GPIbα-VWF affinity cause platelet- type von Willebrand disease, the well-known mutations are p.Met255Val and p.Gly249Val (often in literature referred to as p.Met239Val and p.Gly233Val, when not counting the signal peptide) (Figure 1c). Effect of p.Pro46Leu, measured by us using SPR, is much less than that of p.Met255Val (Table 1). This is not unexpected, since our patient does not have platelet-type von Willebrand disease.

Our results indicate that genetic changes that affect GPIba interactions might be relevant in typical HUS. Furthermore, it is possible that gain-of-function GPIBA sequence variations might be relevant in the etiology of thrombotic thrombocytopenic purpura, a thrombotic microangiopathy, where decreased ADAMTS13 activity leads to inefficient cleavage of the highly potent ULVWF multimers [24]. Similar deficiencies might be involved in both typical and atypical HUS, such as the MCP mutation mentioned earlier [13]. Therefore, involvement of GPIBA deficiencies in atypical HUS is also possible. We screened this gene in our cohort of 64 aHUS patients [25]. No novel genetic changes or SNP frequencies different than reported for European population (http://www.ncbi.nlm.nih.gov/pubmed/) were observed (data not shown), however due to the relatively small number of patients in our cohort the involvement of GPIBA changes in aHUS pathology still cannot be excluded.

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**References**


