Replacement Therapy in Patients with Von Willebrand Disease—Indications and Monitoring

Die Substitutionsbehandlung des von-Willebrand-Syndroms—Indikationen und Monitoring

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Abstract

In patients with von Willebrand disease (VWD), replacement therapy may be indicated in the case of spontaneous bleeding, surgical interventions and injuries/trauma or as a prophylaxis of spontaneous bleeding episodes. The deficient von Willebrand factor (VWF) is replaced with or without factor VIII (FVIII). Dual VWF/FVIII concentrates can be beneficial in the case of low FVIII level, while repeated dosing may lead to very high FVIII levels, with a potential thrombogenic effect in individual VWD patients. An excessive FVIII:C increase can be limited by using a VWF product with a low level of FVIII, achieving a haemostatic adequate FVIII:C increase after 6 to 12 hours. Replacement therapy in patients with VWD shall be individualised considering VWD type, history and risk of bleeding and risk of thrombosis, as well as indication and the individually variable VWF and FVIII increase. Deviations from the dosages and minimum trough levels mentioned in guidelines or recommendations can be considered in justified cases. The objective of this review is to provide recommendations for specific constellations of replacement therapy based on the VWD-specific guidelines available in Europe, the available evidence, own experiences and the consensus of the interdisciplinary German author group.
Introduction

Von Willebrand disease (VWD) is a quantitative (type 1, type 3) or qualitative (type 2) impairment of von Willebrand factor (VWF).\(^1\) VWD can be congenital or acquired and impairs haemostasis by:

- Interference with VWF-mediated platelet binding to the subendothelium in situations of vascular damage (primary haemostasis).
- The coagulation process caused by a deficiency of circulating FVIII,\(^2\) which is rapidly cleared from the circulation if not bound to VWF (FVIII/VWF ratio: 1:50).\(^3\)

Mediation of platelet adhesion and aggregation depends particularly on high-molecular-weight VWF multimers (5,500–10,000 kDa), while FVIII binding can also be achieved with low-molecular-weight VWF multimers (500–2,500 kDa).\(^4\)

The diagnosis of VWD and subtypes is made based on a conspicuous own or family history of bleeding with confirmation using appropriate laboratory tests as described elsewhere in detail.\(^5\) The diagnostics should include antigen (Ag) and function tests at a plasma and platelet level [VWF:Ag, VWF:Ristocetin cofactor (VWF:RCo) or test variants, VWF:collagen binding (VWF:CB) and factor VIII coagulant (FVIII:Co)], might need further subtyping tests such as assays of VWF multimers and Ristocetin-induced platelet aggregation, and can be substantiated by molecular genetic examinations.\(^6\) The severity of clinical tendency to bleed does not necessarily correlate with the VWD type and laboratory results.\(^7,8\) Patients with VWD should be equipped with an emergency health card that is updated on a regular basis.

Therapeutic Options

In addition to supportive measures (e.g., tranexamic acid especially in the case of mucosal bleeding, oral contraception in the case of menorrhagia, iron replacement), the following treatment options can be considered:

- Mobilisation of stored VWF using desmopressin (1-desamino-8-d-arginine vasopressin, DDAVP).
- Replacement of deficient VWF with or without the addition of FVIII.

Since endogenous FVIII production is intact in the case of congenital or acquired forms, the lack of circulating FVIII can be compensated if sufficient functional VWF is made available in the plasma.

In patients with VWD type 1, DDAVP is effective in most cases. DDAVP can be contraindicated in type 2B and, of note, DDAVP is not useful in type 3 cases. Whenever possible, a DDAVP test infusion should be given prior to the initiation of treatment to investigate the individual response.\(^9\) If DDAVP is repeated after a short interval, tachyphylaxis (reduced efficiency due to empty endothelial stores) and the increased risk of hyponatremia and subsequent seizures especially in young children must be considered, respectively avoided by adequate fluid restriction.

Replacement treatment with VWF can be indicated in patients with VWD in the following situations:

- Spontaneous bleeding.
- Surgical procedures.
- Injury/trauma.
- Prophylaxis of spontaneous bleeding.
The recommendations for replacement treatment in patients with VWD are given below. These are based on a selective literature review (1987 till 2017), considering the guidelines/recommendations and the consensus of the interdisciplinary German author group.

**Licensed Products for Replacement Therapy in Patients with VWD in Germany**

The following products are available for individualised replacement therapy:

- Dual concentrates (FVIII products containing VWF).
- Highly purified VWF products (Willfact) with a very low FVIII content and Veyvondi, a recombinant human VWF product with only traces of FVIII.\(^{10}\)

The use of dual VWF/FVIII products prevails in German-speaking countries. Dual VWF/FVIII products are beneficial when the FVIII levels are low.\(^9\) This might depend on VWF type/subtype and clinical situation, such as acute bleeding or emergency surgery, when usage of free combinations of a VWF and FVIII product does not seem feasible, as well as on-demand treatment of acute bleeding in the self-treatment setting at home or on vacation.

When FVIII levels are normal, the risk of thromboembolism should be taken into consideration if supraphysiological FVIII concentrations are induced by the exogenously added FVIII.\(^9,11\) With repeated administration of dual concentrates, accumulation may occur due to the overlapping of exogenously administered FVIII and the formation of endogenous VWF-stabilized FVIII.\(^9,11,12\) For longer treatment periods (>24–48 hours), a dose reduction and/or prolongation of the dosing interval—considering the clinical efficacy of the VWF level—or switching to a VWF concentrate with a low FVIII portion is recommended.\(^{12–15}\)

To avoid an excessive FVIII:C increase after replacement, a highly purified plasma-derived VWF product with a low FVIII portion is available (\(\rightarrow\) Table 1). A recombinant VWF with only traces of FVIII recently received market authorization.\(^{16}\) If VWF with a low or without FVIII is injected, a haemostatically adequate FVIII:C increase is usually achieved after 6 to 12 hours by stabilization of the patients’ endogenously formed FVIII (\(\rightarrow\) Fig. 1).\(^{17,18}\)

In a prospective study of patients with different types of clinically severe VWD, very good clinical outcomes were achieved with highly purified VWF concentrates for both spontaneous bleeding episodes and surgical interventions. An additional dose of FVIII was given for the first infusion in 38% of the spontaneous bleeding episodes and 12% of the surgical interventions.\(^{22}\)

The products listed in \(\rightarrow\) Table 1 are licensed options available in Germany for replacement therapy in patients with VWD. Important parameters for product selection include:

- Manufacturing process (purification, virus inactivation and elimination).
- VWF to FVIII ratio.
- VWF:RCo to VWF:Ag ratio.
- Storage and stability.

Regarding the terminal half-life of VWF:RCo, the dual preparations show large individual ranges of variation (e.g., 2.8–51.1 hours),\(^{12–15,19}\) which is one of the reasons for individualized dosages.

Depending on the manufacturing process of plasma-derived VWF products, the function of VWF may be impaired by a loss of higher molecular weight proteins. Therefore, products with a VWF:RCo/VWF:Ag ratio > 0.7 should be favoured.\(^{23}\) The recombinant VWF product contains beside physiologically occurring plasmatic multimer structures also ultra-large multimers.\(^{16}\)

The product dosage should be based on the VWF:RCo level.\(^{23}\) In this context, 1 IU of VWF:RCo/kg raises the VWF plasma level by an average of 1.7 to 2.1%.\(^{16–18,24–26}\)

**Monitoring of Replacement Therapy**

Monitoring is indicated during replacement therapy with VWF.\(^{27}\) Historically, the focus was on the sole monitoring of FVIII. Meanwhile, more automated and standardised tests for measuring VWF activity (VWF:RCo or test variants) are available, which are additionally recommended for individual treatment management, although this is discussed controversially.\(^{28,29}\) For major surgical procedures, FVIII and VWF activity should be monitored daily for at least 1 week; at least once for minor surgical procedures. An in vivo recovery determined pre-operatively can predict the FVIII:C and VWF activity levels achieved peri- and post-operatively only to an insufficient extent.\(^{30}\)

The target levels should be defined individually depending on VWD type, underlying disease and clinical situation, as well as other factors which may be considered, such as age and ABO blood type, accounting for varying reference ranges.\(^{31}\) In addition to dosages, guidelines and recommendations also indicate minimum trough levels for orientation (\(\rightarrow\) Table 2).

**Possible Clinically Relevant Side Effects of Replacement Therapy**

Hypersensitivity or allergic reactions occur rarely outside inhibitor development when using coagulation factor products.

Registry data and case series with small sample sizes point out the possible formation of inhibitors, the incidence of which is approximately 4% and almost exclusively in type 3 VWD.\(^{33–35}\) Inhibitors whose titres are sometimes high can be directed against VWF or FVIII\(^{35,36}\) and can lead to an inadequate clinical response. Anaphylactic reactions are possible at the same time as inhibitor development. Genetic predisposition,\(^{36}\) FVIII content, product type and dosage are discussed as factors which may influence inhibitor development.\(^{35}\)

In the normal population, increased FVIII levels > 150% are considered as risk factors for venous thromboembolism.
Table 1 Products containing VWF/FVIII for the prevention and treatment of bleeding and for the treatment of bleeding in the case of surgical procedures in patients with VWD when DDAVP alone is ineffective or contraindicated (descending order of Ratio VWF:RCo/FVIII:C)\textsuperscript{12–16,19,20}

<table>
<thead>
<tr>
<th>Product (manufacturer)</th>
<th>Pharmaceutical form\textsuperscript{a}</th>
<th>Declared factor content</th>
<th>Active concentration per mL</th>
<th>Ratio VWF:RCo/FVIII:C</th>
<th>Origin and virus inactivation/viral depletion</th>
<th>Shelf-life</th>
<th>Duration of demonstrated physicochemical stability from reconstitution</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Veyvond\textsuperscript{b} (Shire)</td>
<td></td>
<td>Only traces\textsuperscript{c}</td>
<td>650 IU\textsuperscript{d}</td>
<td>VWF:RCo</td>
<td>Only traces\textsuperscript{c}</td>
<td>130 IU</td>
<td>–</td>
</tr>
<tr>
<td>Willfact\textsuperscript{16,19} (LFB-Biomedicaments)</td>
<td>≤100 IU\textsuperscript{g}</td>
<td>1,000 IU\textsuperscript{d}</td>
<td>≤10 IU</td>
<td>FVIII:C</td>
<td>100 IU</td>
<td>10 IU</td>
<td>10:1</td>
</tr>
<tr>
<td>Voncento\textsuperscript{13} (CSL Behring GmbH)</td>
<td>1,000 IU\textsuperscript{g}</td>
<td>2,400 IU\textsuperscript{d}</td>
<td>100 IU</td>
<td>VWF:RCo</td>
<td>240 IU</td>
<td>240 IU</td>
<td>2.4:1</td>
</tr>
<tr>
<td>Haemate P\textsuperscript{12} (CSL Behring GmbH)</td>
<td>1,000 IU\textsuperscript{g}</td>
<td>2,400 IU\textsuperscript{d}</td>
<td>66.6 IU</td>
<td>FVIII:C</td>
<td>160 IU</td>
<td>160 IU</td>
<td>2.4:1</td>
</tr>
<tr>
<td>Wilate\textsuperscript{14} (Octapharma GmbH)</td>
<td>1,000 IU\textsuperscript{g}</td>
<td>1,000 IU\textsuperscript{d}</td>
<td>100 IU</td>
<td>FVIII:C</td>
<td>100 IU</td>
<td>100 IU</td>
<td>1:1</td>
</tr>
<tr>
<td>Immunate\textsuperscript{h} (Shire)</td>
<td>1,000 IU\textsuperscript{g}</td>
<td>600 IU\textsuperscript{d}</td>
<td>100 IU</td>
<td>VWF:RCo</td>
<td>60 IU</td>
<td>0.6:1</td>
<td>Human plasma-derived • Pasteurisation (60°C/10 h) • S/D procedure</td>
</tr>
</tbody>
</table>

Abbreviations: Ag, antigen; DDAVP, 1-desamino-8-d-arginine vasopressin; FVIII, factor VIII; RCo, Ristocetin cofactor; S/D procedure, solvent/detergent procedure; VWD, von Willebrand disease; VWF, von Willebrand factor.

Note: The products listed in Table 1 are licensed options available in Germany for replacement therapy in patients with VWD. Products differ substantially regarding FVIII proportion, so dosing according to FVIII:C may lead to differences in VWF dosing.

\textsuperscript{a}Additional pharmaceutical strengths of listed products with the same VWF:RCo/FVIII:C ratio may be available.

\textsuperscript{b}Licensed for adults (≥ 18 y).

\textsuperscript{c}≤ 0.01 IU FVIII/IU VWF:RCo.

\textsuperscript{d}VWF:RCo activity using international WHO standard for VWF concentrates.

\textsuperscript{e}Manufactured by recombinant DNA (rDNA) technology in the Chinese hamster ovary (CHO) cell line without the addition of any exogenous human- or animal-derived protein in the cell-culture process, purification, or final formulation.\textsuperscript{15,16}

\textsuperscript{f}Veyvondi EPAR, 28 June 2018: “Due to the specific nature of the product, implementation of specific steps for non-enveloped virus reduction is difficult. […] The reduction capacity of the rhVWF purification process with respect to non-enveloped viruses is, however, compensated to an acceptable extent by the tight control of the production process with respect to potential virus contamination including virus filtration of cell culture media.”\textsuperscript{21}

\textsuperscript{g}Using chromogenic assay according to European Pharmacopoeia.

\textsuperscript{h}Another indication: VWD with FVIII deficiency: treatment of bleeds if no specific plasma-derived product that is effective in VWD is available.

\textsuperscript{i}Using international WHO standard for VWF for VWD concentrates.
Replacement Therapy in Patients with von Willebrand Disease  Nowak-Göttl et al.
Table 2 Comparison of recommendations in European guidelines/recommendations on VWD including values for replacement therapy for orientation

<table>
<thead>
<tr>
<th>European (^9)</th>
<th>Italian (^12)</th>
<th>Nordic (^23),a</th>
<th>British (^6)</th>
</tr>
</thead>
</table>
| **Spontaneous bleeding** | FVIII:C > 30 IU/dL  
Dose: 20–60 IU/kg single dose or once daily dosing until the bleeding stops, generally 2–4 d | cf. European; “Therefore, when repeated infusions of VWF/FVIII concentrates are necessary, such as during surgical procedures, FVIII:C plasma levels should be measured daily, in order to avoid values in excess of 150 IU/dL.” | VWF:RCo > 30 IU/dL  
Dose: 20–60 IU/kg once daily until the bleeding stops  
Monitoring: clinical | No dose recommendation |
| **Minor surgery (elective)** | FVIII:C > 30 IU/dL  
Dose: 30–60 IU/kg once daily or every other day; generally 2–4 d | cf. European | VWF:RCo > 30 IU/dL  
Dose: 30–50 IU/kg once daily for 1–3 d | VWF:RCo > 50 IU/dL |
| **Tooth extraction or invasive interventions** | FVIII:C > 50 IU/dL for 12 h  
Single dose: 30 IU/kg | cf. European | VWF:RCo > 30 IU/dL for 12 h  
Dose: Usually 1 × 20–30 IU/kg prior to the intervention | VWF:RCo > 30 IU/dL and FVIII:C > 50 IU/dL |
| **Major surgery** | VWF:RCo and FVIII:C > 80–100 IU/dL until 36 h post-op, then >50 IU/dL; duration 5–10 d  
Dose: 50–60 IU/kg once daily until complete wound healing is achieved | FVIII:C > 80–100 IU/dL;  
Dose: 50 IU/kg once daily until complete wound healing is achieved (commonly 5–10 d). | VWF:RCo 50–100 IU/dL  
Level for 3–10 d  
Dose: 40–60 IU/kg initially every 12 h, then once daily until complete wound healing is achieved | VWF:RCo > 50 IU/dL |
| **Long-term prophylaxis, i.e., ≥ 4 d** | To be considered in the case of severe VWD and recurrent bleeding in higher risk localisations (GI, haemarthrosis, epistaxis in children) | cf. European | VWD type 3 joint bleeds or severe epistaxis, GI bleeds, menorrhagia, when other treatments failed: VWF 20–50 IU/kg 2–3 times weekly | VWD type 3 with haemarthrosis, severe epistaxis, angiodysplasia, menorrhagia, and children with joint bleeds: VWD 30–50 IU/kg 2–3 times weekly |
| **Labour/postpartum period** | FVIII:C > 50 IU/dL for 3–4 d  
30 IU/kg once daily for 3–4 d | cf. European | VWF:RCo > 50 IU/dL for 3–4 d  
50 IU/kg once daily for 3–4 days | VWF:RCo > 50 IU/dL for 3–4 d (no dose information) |

Abbreviations: FVIII, factor VIII; GI, gastrointestinal; RCo, Ristocetin cofactor; VWD, von Willebrand disease; VWF, von Willebrand factor.

\(^a\) Nordic guidelines: dose information is based on patients with VWD with strongly reduced FVIII:C and/or VWF:RCo level (<0.10 IU/dL); factor levels can be predicted based on pharmacokinetic data: 40–50 IU/kg (VWF:RCo) increase plasma VWF levels to 80–100% depending on the baseline level and haematocrit.
VWD types, whereas gastrointestinal (GI) bleeding is seen mainly in VWD type 3 and VWD type 2A, which are characterized by partial or complete loss of intermediate- and high-molecular-weight VWF multimers. GI bleeding represents a severe manifestation of bleeding in VWD, angiodysplasia being a major cause. High-molecular-weight structures of VWF appear to influence angiogenesis, and loss of such structures may cause vascular malformation, especially in VWD type 2A and acquired VWD of cardiogenic origin. On the other hand, haematomas and joint bleeds occur mainly in VWD type 3 and in those patients with more severe VWD type 1 characterized by lower FVIII levels.

VWF/FVIII products as well as the highly purified and the recombinant VWF product demonstrated good or excellent efficacy in approximately 89 to 98% of spontaneous bleeding episodes. The dose requirement of replacement therapy is highest in GI bleeding.

**Treatment of Traumatic Bleeding**

The European guidelines or recommendations do not contain separate statements on replacement therapy for trauma-related bleeding. The information on bleeding episodes focus either on spontaneous bleeding or more generally on ‘acute bleeding’. There is no difference between the procedures for spontaneous or traumatic bleeding, except that with traumatic bleeding, surgical intervention may follow.

**Treatment of Acute Bleeding**

In addition to general recommendations of transfusion medicine and administration of anti-fibrinolytics, doses of 20 to 60 IU/kg VWF:RCo depending on the localisation, once or twice daily, are recommended for replacement therapy until the bleeding has stopped. As a rule, a 2- to 4-day replacement therapy is generally recommended. In the case of repeated dosing, the FVIII:C should be monitored to avoid persistent FVIII:C levels > 200%, or a VWF product with a low FVIII level should be considered. In GI bleeding due to angiodysplasia, 40 to 60 IU/kg once or twice daily are recommended until the bleeding has stopped. The duration of treatment is generally longer than those with other VWD-associated bleeds. If GI bleeding is suspected, patients should be hospitalised. A VWF:RCo of >30% does not exclude spontaneous bleeding. Higher VWF activities may be required depending on the VWD type and phenotype of bleeding. In acute spontaneous bleeds or traumatic haemorrhage, the initial treatment must include FVIII in addition to VWF if the FVIII trough levels are inadequate and usage of a dual VWF/FVIII concentrate could be recommended.

**Procedure for Elective Surgical Procedures**

In VWD patients, the need for replacement therapy should be reviewed in cases of planned and elective surgery. In approximately 95% of cases with such indication, replacement therapy was judged to be effective.

Large orthopaedic interventions on target joints require special attention. Target joints are joints with recurrent bleeding due to anatomic modifications (e.g., neovascularisation). Retrospective analysis of a cohort with 126 major joint surgeries revealed bleeding complications in 18% of interventions – the risk of bleeding was higher than average in patients without replacement (33%) and in patients without FVIII monitoring (26%).

**Recommendations in Patients Undergoing Surgery**

Close cooperation between the surgeon and haematologist must be ensured and therapy must be adjusted according to the current guidelines and to the individual and dynamic bleeding situation. For elective procedures, an individualised replacement and monitoring plan (FVIII and VWF activity) should be prepared, with a higher initial dose and subsequent individualised maintenance doses. In emergency surgery requiring replacement therapy, the initial treatment should also include FVIII in addition to VWF to achieve an immediate increase in FVIII levels. In the case of elective surgery, a VWF product with a low FVIII level can be administered if the basal FVIII level is adequate or the first dose is given at least 12 to 24 hours prior to the start of surgery. In this case, an adequate FVIII level can be expected because of endogenous FVIII synthesis. In general, patients with VWD type 3 and 2B in particular require replacement therapy for minor surgical procedures (e.g., biopsies, catheter placements, dental surgery or laparoscopic procedures), whereas in type 1, 2A, 2M and 2N, DDAVP is the treatment of choice if the response is good. Major elective surgical interventions are a possible indication for a VWF concentrate with a low FVIII level, especially if multiple doses are anticipated in patients with an increased risk of thrombosis.

**VWD and Prophylaxis**

Long-term prophylaxis in VWD is being used increasingly common to prevent secondary complications, reduce hospitalization, and improve quality of life. Due to the late onset of clinically relevant bleeding, prophylaxis in patients with VWD is usually secondary prophylaxis, which is started in early adulthood. Bleeding scores can help to assess the severity of bleeding tendency.

Indications for prophylaxis may be severe VWD with recurrent forms of GI bleeding, joint bleeding, severe nose bleeding, menorrhagia and in cases which could not adequately controlled with other therapeutic options.

If joint bleeds occur, an early start of secondary prophylaxis, e.g., at the age of <5 years, is beneficial to prevent arthropathy. A VWF dose of 30 to 50 IU/kg two to three times a week is usually required. In women with type 1 VWD suffering from menorrhagia, replacement therapy is required in approximately 1%. To reduce blood loss due to menorrhagia resistant or not amenable with oral estroprogestin pill, VWF doses of 33 to 95 IU/kg were used in women with VWD for 1 to 6 days per cycle.

Prophylaxis with VWF-containing concentrates is the most efficient method to prevent recurrent GI bleeding, often requiring three or more doses per week. The combination with oral tranexamic acid seems wise in the case of GI bleeding.
Recommendations for Long-Term Prophylaxis

- Long-term prophylaxis
  - should be considered in all patients with VWD and recurrent bleeding\(^5\) and
  - is required in cases of recurrent, clinically relevant bleeding episodes that affect the quality of life.\(^{34,35}\)
- Long-term prophylaxis is a possible indication for a VWF concentrate with a low FVIII level.\(^5,^{32}\)
- Depending on the severity and the impact of VWD on the specific patient, long-term prophylaxis should be initiated with two to three infusions of 20 to 50 IU/kg per week.\(^{23}\)
- In patients with recurrent GI bleeding due to angiodyplasia, doses of 40 to 60 IU/kg two to three times a week are recommended.\(^{50}\)

Procedure during Pregnancy and in the Peri-/Post-natal Period

Due to the physiological increase of pro-coagulant factors during the course of pregnancy,\(^{51}\) pre-existing bleeding conditions in women with VWD type 1 improve in most cases.\(^{59}\)

Thus, clinically relevant bleeding episodes are rare during pregnancy in patients with VWD type 1,\(^{59}\) and women with VWF activities \(>50\) IU/dL prior to pregnancy onset do not automatically need replacement therapy. Levels reached in the third trimester are maintained during labour (important for the anaesthetist) with a decrease reaching baseline non-pregnancy ranges approximately 1 week after birth.\(^{50}\)

Larger studies to evaluate the course of FVIII/VWF levels in the peri-partal and post-partal period are needed. Women with VWD type 2 may reach normal VWF:Ag level, but VWF is dysfunctional and replacement therapy may become necessary, e.g., to avoid intermittent vaginal bleeding, and may be necessary during delivery.\(^{61}\)

A special feature of VWD type 2B is that a previously existing thrombocytopenia can be aggravated by the increased release of abnormal VWF multimers during pregnancy.\(^{61,62}\)

Therefore, continuous monitoring of the platelet count is required. VWD type 3 does not achieve a normal VWF activity apart from bleeding episodes during pregnancy, and replacement therapy is required during labour and delivery.\(^{51,62}\)

The risk of spontaneous abortion in women with VWD is comparable to the rates reported in the normal population,\(^{63,64}\) but in contrast, the rate of recurrent miscarriage in women with VWD is not clear so far.\(^{65}\)

In the view of a multifactorial disease, poor endometrial perfusion during the implantation window is reported to be one of the possible causes of recurrent miscarriage. A meta-analysis reported in 2011 including 18 case–control studies in a total of 2,397 women with recurrent miscarriage revealed that angiogenesis- and vasoconstriction-related genes were involved.\(^{66}\)

Since endothelial VWF regulates angiogenesis\(^{67}\) with deficiency states resulting for example in pathological angiodyplasia and vascular malformations with severe GI bleedings, the latter mechanism has to be discussed as one hypothetical reason also in women with recurrent foetal loss.

Evidence regarding replacement with VWF/FVIII products in pregnancy is limited to case reports and case series.\(^{61,63}\)

In European recommendations/guidelines, a period of prophylaxis of 3 to 4 days after delivery is recommended unanimously with a VWF:RCo target \(>50\) IU/dL.\(^{6,9,23,32}\)

Nevertheless, the optimal duration and frequency of post-partum prophylaxis is unknown.\(^{68}\)

In addition to the risk of bleeding, the pregnancy-related risk of thrombosis must be taken into consideration. Due to the physiological particularities of the balance of coagulation physiology during pregnancy,\(^{31}\) the role of FVIII and D-dimer concentration must be re-evaluated as possible predictors of the risk of thrombosis in this population. The FVIII:C activity can increase to up to approximately 350 IU/dL in healthy women during pregnancy.\(^{31}\)

Therefore, an individual approach, in which only the deficient VWF is replaced, seems to be beneficial in patients with VWD.

General Recommendations Regarding Pregnancy and in the Peri-/Post-natal Period

- History of bleeding and family history should be considered.
- VWF:Ag, VWF activity and FVIII:C should be checked prior to invasive procedures, and in certain patients in vitro bleeding time (e.g., platelet function analyzer, PFA) might be useful.\(^{59}\)
- Evidence regarding FVIII:C and VWF levels, which may require treatment or prophylaxis, is lacking.\(^{61}\)
- Restrictive use regarding DDAVP is necessary due to possible side effects and interactions, e.g., oxytocin receptor (uterus contraction\(^{68,70}\); tranexamic acid as an alternative).
- In the case of VWF replacement in a prothrombotic situation, a FVIII overload must be avoided and preference should be given to a product with a low FVIII content.

Recommendations Regarding Labour/Post-partum Period

- All women with VWD (including type 1) should give birth in close cooperation with an obstetrician, haematologist, anaesthetist and neonatologist and—in the case of severe VWD—be managed in cooperation with a centre with a blood bank and 24-hour monitoring of FVIII, VWF and maybe in vitro bleeding time (PFA).\(^{61,69}\)
- Peri-partal therapy such as replacement is rarely required in women with VWD type 1 and VWF:RCo \(\geq 50\) IU/dL if no severe bleeding is documented in the patient’s medical history.\(^{71}\)
  However, a VWF activity \(>50\) IU/dL in week 34 to 36 might be too low in certain women, e.g., women with additional FXIII deficiency.
- During and after delivery, a VWF:RCo of at least 50 IU/dL and a VWF:RCo dose of 50 IU/kg once daily for 3 to 4 days are to be aimed at (longer in the case of caesarean delivery)—in individual cases, this procedure can be used for further operations.

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Blood loss should be documented up to approximately 14 days after delivery (give instructions to the woman if applicable).

- No spinal anaesthesia, e.g. peridural anaesthesia, without replacement in patients with a VWF:RCo < 50 IU/dL or with a history of severe bleeding (exception: VWD type 1 with normal PFA values and VWF:RCo ≥ 50 IU/dL).

**Procedure in Paediatrics**

No specific recommendations for replacement therapy in children with VWD can be found in European guidelines, and therefore recommendations for adults must be applied. In the overall population, the lower reference values (10th percentile) for FVIII:C are 50% in the first 12 months of life. Between 1 and 18 years, the 10th percentile for FVIII:C were reported as 55% (1–5 years), 58% (5.1–10 years) and 48% (10.1–18 years). These small differences between the age groups are not clinically relevant for issues regarding VWD. No stratification according to blood groups is available for children.

The indications for the plasma-derived VWF products are not restricted to adults or specific age groups in patients with VWD, although clinical data are limited in children and adolescents. The market authorization for the recombinant VWF is currently limited to adults only. Unlike patients with haemophilia A, pharmacokinetics of the VWF/FVIII concentrates were not found to be age-dependent. There is limited data on the prophylaxis in children with VWD compared with prophylaxis in children with haemophilia.

**Recommendations**

- Children who are likely to require replacement therapy should be vaccinated subcutaneously against hepatitis A and B.
- Children with VWD who need to undergo surgery or dental interventions should be treated at hospitals with an on-site haemophilia center.
- Short-term prophylaxis is often needed in children with VWD type 3 but may also be required in children with severe type 1 and type 2.
- In children, depending on the type of VWD, long-term prophylaxis may be applied individually (1–3 times weekly), e.g. in recurrent joint bleeds and possibly in children with recurrent epistaxis. However, long-term prophylaxis should be restricted to intensive bleeding periods and the indication must be re-evaluated on regular follow-up visits.
- Replacement therapy can be administered at the haemophilia centre or self-administered at home.

**Conclusion**

Decisions on replacement therapy in patients with VWD are to be made individually based on VWD type, history and risk of bleeding and risk of thrombosis as well as the indication (treatment or prophylaxis of bleeding, surgical intervention or labour) and the variable individual VWF and FVIII increase. Other factors such as age and ABO blood type may be considered.

Information on dosages and minimum trough levels in guidelines or recommendations are for orientation only and deviations can be considered if justified.

- Clinically relevant side effects of replacement therapy are notably hypersensitivity reactions, the occurrence of inhibitors and thromboembolic complications.
- Dual VWF/FVIII products have advantages when the FVIII levels are low:
  - Acute bleeding or during emergency surgery, when usage of free combinations of a VWF and a FVIII product does not seem feasible.
  - On-demand usage in the self-treatment setting at home or on vacation.
- A VWF concentrate with a low or zero level of FVIII may be the treatment of first choice in the following situations:
  - Repeated infusions in the case of long-term prophylaxis, major bleeding episodes or major surgical interventions.
  - Pre-existing risk of thrombosis (age > 70 years, own history or family history of thromboembolism, obesity, immobility ≥ 3 days, cardiovascular, malignant or inflammatory disease or cases of coagulation activation, thrombophilia or increased FVIII:C level), pregnancy and labour.
- In the case of short-term replacement for elective surgery, a VWD concentrate with a low level of FVIII can be considered as well.

**Conflict of Interest**

U.N.G. declares speaker and/or advisor honoraria. W.M. declares speaker and/or advisor honoraria from Shire, Octapharma, LFB and CSL Behring. J.K. declares speaker and/or advisor honoraria from Aspen, Bayer Health Care Pharmaceuticals, Daiichi Sankyo, Boehringer Ingelheim, CSL Behring, Pfizer, LFB, BMS, Mitsubishi, Roche, Sanofi and Novo Nordisk. C.E.D. declares speaker and/or advisor honoraria from Bayer, Boehringer Ingelheim, Daiichi Pharma, Pfizer and LFB. M.M. declares speaker and/or advisor honoraria from Astra Zeneca, Bayer, CSL Behring, IL-Werfen/TEM International and LFB Biomedicaments. M.v.D.P. declares speaker and/or advisor honoraria from Astra Zeneca, Bayer, CSL Behring, Novartis, Octapharma and Shire. D.W. was an employee of LFB GmbH, Münster, from February 2017 to January 2018. M.S. declares speaker and/or advisor honoraria from Bayer, LFB, Novo Nordisk, Pfizer, Shire, Sobi, IL-Werfen and CSL Behring.

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**References**

16 Baxter Innovations GmbH. Fachinformation Veyvondi, effective February 17, 2019
27 Wissenschaftlicher Beirat der Bundesärztekammer. Querschnitts-Leitlinien zur Therapie mit Blutkomponenten und Plasmaerivaten – Herausgegeben von der Bundesärztekammer auf Empfehlung ihres Wissenschaftlichen Beirats; 4. überarbeitete Auflage; Deutscher Ärzte-Verlag, Köln, 2014
39 Girolami A, Tasinato V, Sambado I, Peroni E, Casonato A. Venous thrombosis in von Willebrand disease as observed in one centre and as reported in the literature. Blood Coagul Fibrinolysis 2015;26(01):54–58
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