Thromboembolic Risks of Non-Factor Replacement Therapies in Hemophilia

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Impressive advances in the field of hemophilia have been reported during the recent congress of the International Society on Thrombosis and Haemostasis (ISTH) in Berlin in July 2017. Two of the major oral abstract presentations presented studies in the field of hemophilia that appeared the same day as full publications in the New England Journal of Medicine (1, 2), reinforcing the importance of hemophilia as a driver of innovation in hemostasis.

The new non-factor replacement therapies promise to overcome two important shortcomings of traditional factor replacement therapy: (i) the need of frequent intravenous administrations; and (ii) the risk of transient or permanent formation of neutralizing antibodies, so-called inhibitors (3, 4).

Hemophilia with inhibitors: state of the art

The mainstay of treatment in congenital hemophilia with inhibitors (CHWI) is immune tolerance induction (ITI), aiming at eradicating inhibitors and inducing tolerance to factor VIII or IX (5). This treatment involves repeated administration of clotting factor, twice daily in the most intense protocols (6). Placement of central venous lines is usually required. The treatment is successful in 60 to 80% after several months of treatment (7), and is often hampered by problems with venous access, including line infection and thrombosis (8). Until tolerance is achieved, bleeds are difficult to prevent and difficult to control, resulting in higher risks of morbidity, mortality, and disability (9–12). FVIII bypassing agents (BPA), including activated prothrombin complex concentrate (APCC) and recombinant activated factor VII (rFVIIa) can restore hemostasis in the absence of factor VIII (FVIII) or IX (FIX) and effectively stop acute bleeds in most patients (13). However, the ability of those BPA to prevent bleeds in a prophylactic setting is limited, partly because of their short half-life (14–16).

These issues set the scene for the development of the novel non-factor replacement therapies (17). Representing diverse mechanisms of action, they have in common (i) efficacy independent of the presence of inhibitors, (ii) long half-life, (iii) subcutaneous administration, and (iv) reduced or absent risk of anti-drug antibodies. However, first reports on thromboembolic events remind us that groundbreaking innovation may not only offer unprecedented success but also unexpected pitfalls and risks.

Emicizumab: FVIIIa-mimetic bispecific antibody

Developed under the name of ACE 910, emicizumab is a humanized bispecific antibody targeting FIX/FIXa and factor X (FX) to support the activation of FX. Emicizumab will replace FVIIIa in hemophilia A with or without inhibitors. It is not useful in hemophilia B.

The clinical study results published so far underline the great potential of the drug (1, 18). The HAVEN 1 study presented on ISTH 2017 was an open-label, randomized controlled study of 109 adult and adolescent patients with hemophilia A and inhibitors (1). Patients previously using on-demand BPA were randomized to prophylaxis with emicizumab (arm A, 1.5 mg/kg weekly, n = 35) or continued on-demand BPA (arm B, n = 18). During the study period of 24 weeks, the number of treated bleeds was 87% lower in patients on emicizumab (mean annualized bleeding rate [ABR] 2.9; 95% confidence interval 1.7–5.0) compared to on-demand BPA (23.3; 12.3–43.9). Patients previously on prophylactic BPA switched to emicizumab without randomization (arm C) and experienced a 79% reduction in bleeds (mean ABR 3.3; 1.3–8.1) compared to the historic control period (15.7; 11.1–22.3).

The drug was generally well tolerated. The most frequent adverse event was injection site reaction in 15% of patients. Four thromboembolic events were reported in the primary analysis. This was amended to a total of 5 events in 5 patients during the ISTH conference oral presentation by Odenburg et al. (results update as of April 2017, [19]): 3 cases of thrombotic microangiopathy (TMA), and 2 venous thromboembolic events, including a sinus vein thrombosis, and a superficial thrombophlebitis. One patient died in the context of rectal bleeding that could no longer be treated with APCC due to the TMA.

Emicizumab: a closer look at thromboembolic events

In HAVEN 1, all thromboembolic events occurred during episodes of treatment with APCC for breakthrough bleeding. The risk appeared to be dose-related: the 5 events occurred during 5 out of 8 treatment episodes of APCC >100 U/kg per day for >24h. In contrast, no events occurred with lower doses of APCC (≤100 U/kg) or shorter treatment durations (≤24h), or when rFVIIa was used for breakthrough bleeding (Fig. 1).
Even though thrombosis was occasionally observed with APCC (20) and rFVIIa (21, 22), TMA is a truly novel form of thromboembolic complication that was seen neither with APCC or rFVIIa alone, nor with emicizumab alone in > 350 patients enrolled altogether in the ongoing study program.

It will be important to better understand the pathomechanism of thromboembolic events during treatment with emicizumab. The drug’s affinity to FIX/IXa and FX/Xa is lower in solution than that reported for FVIIIa on phospholipid membranes (23). The $K_D$-based simulations predict that with therapeutic plasma concentrations of emicizumab, the majority of FIX and FX exist as monomers, followed by emicizumab-FIX and emicizumab-FX dimers, and only a very small fraction as ternary FIX-emicizumab-FX complexes. However, these measurements have not been performed on phosphatidylserine-exposed phospholipid membranes or in the presence of APCC. It is tempting to speculate that active phospholipid membranes and/or increased concentrations of (pro)-enzyme FIX(a) or substrate FX may increase the drug’s ability to support FXa generation and result in excessive thrombin formation.

For the time being, the manufacturer recommended to avoid the combination of emicizumab with APCC and, if APCC was needed, to use it at the lowest dose possible. It remains to be seen whether APCC can be avoided in all CHWI patients receiving emicizumab, given the established experience that some patients respond clinically differently to the BPA and may not be sufficiently treated with rFVIIa.

### Fitusiran: antithrombin silencing

Fitusiran is a small interfering ribonucleic acid (siRNA), coupled to N-acetylgalactosamine (GalNAc). GalNAc-siRNA conjugates can be administered subcutaneously and will be targeted to hepatocytes through uptake by the asialoglycoprotein receptor. Inside the cell, it will turn off the expression of antithrombin (SERPINC1) mRNA to achieve stable and consistent reduction in plasma antithrombin activity to about 20% of baseline (2).

Low antithrombin, in the context of congenital deficiency, has been reported to increase thrombin levels (24). Occasionally, hemophilia patients with a less severe bleeding phenotype have been found to carry inherited thrombophilic ‘risk’ factors including antithrombin deficiency (25). Heterozygous antithrombin deficiency also mitigates the phenotype of FVIII deficiency in the hemophilia A mouse model (26). These observations formed the rationale for the development of antithrombin silencing by RNA interference in hemophilia.

The phase 1 dose-escalation study enrolled 4 healthy individuals and 25 patients with moderate or severe hemophilia A or B (26). According to the June 2017 data update of the phase 2 open-label extension study presented during ISTH (27), 33 patients with hemophilia A or B, with or without inhibitors, have been treated for a median of 11 months. The median ABR for non-inhibitor patients was 1.7 (n = 19), and for inhibitor patients 0 (n = 14). This compared with a historic ABR of 2 (non-inhibitor patients previously on prophylaxis, n = 7), ABR of 12 (non-inhibitor patients previously treated on demand, n = 12), and ABR of 38 (patients with inhibitors, n = 14), suggesting that fitusiran can prevent bleeds in most patients with and without inhibitors.

The drug was generally well tolerated. Injection-site reactions were seen in 18% of patients; abdominal pain, diarrhea, and headache occurred in 9% each. Asymptomatic alanine transferase (ALT) increases $\geq 3x$ upper limit of normal occurred in 11 patients (33%), all of which were hepatitis C antibody positive.

Of note, breakthrough bleeds in non-inhibitor patients were managed with low doses of FVIII (18 bleeds; mean 17 [range 5–31] IU/kg for a mean of 1.1, [range 1–2] doses) or FIX (7 bleeds; 18 [9–27] IU/kg for 3.9 [1–8] doses). All patients used less than or same amount of factor per bleed as prior to fitusiran. In inhibitor patients, APCC (56 bleeds; 27 [14–37] U/kg for 1.5 [1–3] doses) and rFVIIa (3 bleeds; 56 [37–62] µg/kg for 2.7 [2–3] doses) were also dosed less intensely in most patients. No thromboembolic events were reported in these patients.

### Fitusiran: first thromboembolic event

Most recently, a first thromboembolic event with fitusiran was reported (28). According to the study sponsor’s press release, this was a patient, who treated an exercise-induced hip pain with three doses of FVIII (31–46 IU/kg). On the last day of dosing, he developed severe headache, was diagnosed with subarachnoid hemorrhage on CT scan, and received further treatment with FVIII. Over a 14-day hospitalization his condition worsened and he died from...
cerebral edema. Post-hoc review of the initial CT scan, however, confirmed that the initiating event was a sinus vein thrombosis rather than subarachnoid hemorrhage. The sponsor suspended fitusiran dosing in all patients currently enrolled in the trial program.

Comment: where do we go from here?

Although both drugs hold great promise as a potential treatment for patients with and without inhibitors, the thromboembolic events observed during the trial programs raise concerns about their safety. We must concede that thrombosis used to be a rare event in the hemophilia population, even in elderly patients and during treatment with BPA. Of note, the thromboembolic events seen in emicizumab and fitusiran trial programs were mostly atypical in nature, including sinus vein thrombosis and TMA. Deep vein thrombosis, the most frequent form of venous thromboembolism in the general population, has so far not been observed. This suggests a distinct pathophysiology of thromboembolism in patients treated with the non-factor replacement therapies that needs to be explored carefully.

First, more basic research is needed to understand mechanisms of action. How do emicizumab and APCC interact with each other? How can hemostasis be limited to the site of injury in the presence of emicizumab? What happens if emicizumab dosing is increased, either intentionally or accidentally?

Second, we need dose finding studies for BPA and factor concentrates in patients treated with emicizumab or fitusiran. Thrombin generation experiments and occasional observations in patients treated in the trial programs indicate that lower doses and shorter treatment durations of BPA or FVIII may be adequate to restore hemostasis in the event of breakthrough bleeds. However, this needs to be studied more carefully at the clinical level. For the time being, the combination of emicizumab with APCC >100 U/kg for >1 day appears to be clearly contraindicated. However, it may not be feasible to abstrain from APCC use in all inhibitor patients. Surgery and major bleeds may require the use of BPA for prolonged periods of time, posing significant uncertainties on how to dose APCC, but also FVIIa, FVIII and FIX. Laboratory monitoring of coagulation activation markers or fragmented red blood cells (schistocytes) could potentially help to recognize thromboembolic risks earlier, but this needs systematic investigation in the ongoing and forthcoming studies.

Third, the development of antitides should be considered. An antitode against emicizumab is currently not available, but might be warranted in certain clinical scenarios, very much like the antitides against the direct oral anticoagulants. Antithrombin lowering by fitusiran can theoretically be reversed by infusion of antithrombin concentrate. However, the clinical scenario where this might be useful must be carefully considered and experience should be systematically collected in studies and registries.

Conclusion

The non-factor replacement therapies are groundbreaking developments addressing a major unmet medical need. They have the potential to revolutionize hemophilia treatment, both for patients with and without inhibitors. However, in the field of hemostasis and thrombosis we had to realize often that hemostasis and thrombosis are two sides of the coin that cannot be easily separated. Bleeding has always been the major concern of antithrombotic therapies; and thrombosis turns out to be the major issue with the new hemophilia treatments. Again, we are facing how difficult it is to keep the balance.

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