Laboratory Monitoring of Factor VIII and Factor IX in Hemophilia Patients

Learning Objectives:

- 1. List and describe functional assays for hemophilia A and B.
- 2. Explain assay discrepancies for factor replacement therapy.
- 3. Detail the identification of inhibitory antibodies for factor replacement therapy.

Abstract:

This review describes and discusses the different types of factor assays used in the clinical laboratory for the diagnosis, as well as during therapeutic management, of hemophilia patients. After diagnosis, infusion of factor replacement concentrates is the standard of treatment for patients with hemophilia to prevent or treat bleeding episodes. Monitoring of the factor activity is necessary to assure proper treatment. Several assays are used to assess the factor activity at the time of diagnosis and during therapeutic management. The pros and cons of each assay are discussed in detail. With the introduction of many new replacement products it is important the clinical laboratory understands the advantages and disadvantages of each assay with respects to each of the replacement products available.

Abbreviations: APTT - activated partial thromboplastin time, CaCl₂ - calcium chloride, CSA - chromogenic substrate assay, CV - coefficient of variance, EHL - extended half-life, FVIII - factor VIII, FIX - factor IX, FIXa - activated factor IX , FX - factor X, FXa - activated factor X, HA - hemophilia A, HB - hemophilia B, PEG - polyethylene glycol, PL - phospholipids, *p*NA - *para*-nitroanaline

Index Term: one-stage assay, two-stage assay, chromogenic substrate assay, hemophilia A, hemophilia B, replacement products

Hemophilia A (HA) and B (HB) are congenital bleeding disorders associated with bleeding into joints and muscle tissues and are characterized by absent or reduced quantity or dysfunction of coagulation factor VIII (FVIII) and factor IX (FIX), respectively. The severity of the disease is closely correlated with the factor's plasma activity level. Administration of replacement factor concentrate forms the standard treatment in patients with severe hemophilia (<1% factor activity), with nearly all hemophiliac boys in the U.S., Canada, Australia and Northern Europe receiving prophylactic treatment, resulting in reduced morbidity and mortality. The past decade has seen an explosion of new treatment options, including recombinant clotting factors engineered to extend their half-lives. Several of these extended half-life (EHL) FVIII and FIX concentrates have recently been approved and others are in clinical trials. While coagulation factor activity is measured for initial diagnosis, it is mostly measured in the clinical laboratory for clinical management at regular follow-ups, post infusion of factor concentrate, during preand post-operative procedures and screening of inhibitor development. The one-stage assay (OSA) employing the activated partial thromboplastin time (APTT) is the most widely-used assay in clinical laboratories. The two-stage chromogenic substrate assay (CSA) is an alternative assay which has been available since the 1980s for FVIII and more recently available for FIX. The modifications applied to these molecules have introduced variations in their activity measurement in routine factor assays. Performance characteristics of the OSA and CSA for quantitation of FVIII and FIX at the time of diagnosis and post-infusion of the plasma-derived factor concentrate (pdFVIII and pdFIX) as well as recombinant factor concentrates without (rFVIII and rFIX) or with extended half-life (EHL-rFVIII and EHL-rFIX) have been reported, with many questions still outstanding. Thus, the clinical laboratory is now challenged to

appropriately, accurately and reproducibly measure factor activity levels for the infusion products given to patient. This article will focus on the measurement of FVIII and FIX activity in the clinical laboratory setting, with particular emphasis at diagnosis and in postinfusion plasma.

Laboratory Measurement of FVIII and FIX

Functional Assays

The two different functional assays used for quantification of FVIII and FIX are the OSA and the CSA.¹ The OSA measures the ability of patient plasma to reduce the APTT clotting time of FVIII or FIX deficient plasma (obtained from congenitally deficient patients or immunodepleted normal pooled plasma (Figure 1). Different dilutions of patient plasma are added to deficient plasma and pre-incubated with an APTT reagent containing phospholipid (PL) and a contact activator (e.g. silica, ellagic acid, kaolin, polyphenols). After a set incubation time calcium chloride (CaCl₂) is added and the time to fibrin clot formation is measured according to the analyzer's methodology and algorithm. The result is derived from a standard curve generated with a reference plasma of known FVIII and FIX concentrations. The large number of APTT reagents, deficient plasmas, reference plasmas, instruments, technical standardization procedures

The CSA for FVIII is a two-stage assay (Figure 1). In the first stage, a reagent containing

purified activated Factor IX (FIXa), FX and thrombin, as well as PL and CaCl₂, are added in optimal concentrations to the patient's plasma, resulting in the generation of activated FX (FXa). This step does not require additional coagulation factors due to the direct activation of FVIII by thrombin. In the second stage, the amount of generated FXa is measured by cleavage of a FXaspecific chromogenic substrate, releasing para-nitroanaline (pNA) which causes a color change detected spectrophotometrically at 405nm.^{1,5,6} Available commercial CSA kits for FVIII differ in source of purified proteins, activation process for FVIII, sample diluent, plasma dilution, CaCl₂ concentration and incubation times (Table 1). There are limited studies comparing CSA kits for sensitivity of FVIII measurement in different subgroups of HA.^{7,8}

and combinations thereof results in CVs of up to 40% and underestimation of factor activity by

specific reagents.²⁻⁴

The CSA for FIX is also a two-stage assay (Figure 1). In the first stage, diluted patient plasma is mixed with a reagent containing purified FVIII and FX. The reaction is initiated with the addition of a mixture of activated FXI (FXIa), thrombin, PL and CaCl₂. FXIa activates FIX to FIXa and thrombin activates FVIII to activated FVIII (FVIIIa). FIXa and FVIIIa in the presence of PL and calcium activates FX to FXa. In the second stage, the amount of generated FXa is measured by its specific activity on a FXa chromogenic substrate which is cleaved, releasing *p*NA which causes a color change detected spectrophotometrically at 405nm.^{1,4-6} Available commercial CSA kits for FIX are listed in Table 1.^{1,5,6}

The two-stage clotting assay is a labor-intensive assay that is generally performed manually. It is presently only performed in very few specialized laboratories and will not be discussed.^{1,2}

Assay discrepancy in plasma samples

In a large proportion of untreated patients with moderate or mild forms of HA a systemic discrepancy between the OSA and the CSA for the diagnosis of HA has been reported.^{5,9-18} Several reports have demonstrated discrepant results associated with a specific causative FVIII gene mutations. $^{1,16,17,19-24}$ The incidence of discrepant results has been reported to be 12 – 50%.¹⁶⁻²¹ Discrepancy has not been found between OSA and CSA assays in patients with severe hemophilia.¹ As an example, in a study of 163 HA hemophiliac, 32 (20%) belonging to 16 families had discrepant FVIII levels, defined as either the OSA or CSA being at least 1.5 times higher than the other. Twenty-four patients (15%; 10 families) showed OSA results greater than CSA results, and 8 patients (5%; six families) presented CSA results greater than OSA results. Six of the patients in the latter group had CSA results within the normal range. In patients with discrepant results, other members of the family showed similar laboratory data. The bleeding tendency and the FVIII and/or DDAVP consumption correlated with OSA result. Mutation analysis showed a strong relationship between the phenotype and the mutation, here and elsewhere, suggesting a genetic basis for the discrepancy between OSA and CSA results.¹⁹⁻²¹ This study shows if only the OSA is used for screening of patients with mild HA, some patient will seem to have normal FVIII activity levels and not be diagnosed as having HA or will be assigned to a milder bleeding risk than is the case, increasing the risk of bleeding complications in a subsequent surgery without proper hemostatic treatment. Conversely, on rare occasions patients with normal FVIII activity in vivo will be diagnosed as having HA, risking cancellation or delay of necessary surgery or exposure to unnecessary factor concentrates or blood products.¹

In mild HB, the CSA is higher than OSA, with the CSA result more closely correlating with bleeding phenotype.²⁶ For HB only a single report could be found investigating the discrepancy between OSA and CSA on HB patient plasma samples. In this report from a single center 50 plasma samples from 36 patients were analyzed. Patients with severe HB showed no discrepancy in results. Among the 44 plasma samples from 32 patients in 18 families with mild or moderate HB, 15 samples from 8 patients (25%) showed a CSA at least two-fold greater than the OSA. Of these 15 samples, 14 were from 7 individuals from 5 families with 2 different nucleotide substitution at the same amino acid (6 with p.Arg191His and 1 with p.Arg191Cys) in the N-terminal cleaving site of the activation peptide. The mutations were not observed in patients with non-discrepant results. For patients with the Arg191His mutation the OSA method result was 2% for all and a range of 5 - 8% (p =0.001) for the CSA method. A detailed review of the medical records for these patients revealed 5 patients on on-demand treatment and one younger patient on prophylaxis with no history of spontaneous bleeding episodes. Only one of these patients has experienced a spontaneous bleed (hematuria) requiring replacement therapy. The single patient with the Arg191Cys mutation had FIX levels of 1% and 5% by OSA and CSA, respectively. This patient was reported as being currently on on-demand treatment, but has been on short-term prophylaxis due to episodes of epistaxis. The authors interpreted the bleeding frequency for these patients as low and indicative of a mild bleeding phenotype and state their findings imply assay discrepancy occurs frequently and both OSA and CSA assays are needed for correct diagnosis and classification of HB.²⁵

Replacement products for clinical treatment

In addition to the diagnosis of hemophilia, factor activity measurement is needed for monitoring of factor levels in prophylactic therapy and during treatment of acute bleeds, determination of the pharmacokinetics, and successful perioperative management of factor replacement therapies. Prophylactic therapy is presently the standard of care in the developed world for the severe form of hemophilia and consists of frequent infusion of a factor replacement product, most commonly plasma-derived (pd) or recombinant (r) full-length factor concentrates. It has proven to be efficient in controlling bleeding episodes of patients with hemophilia; however, it requires frequent infusions (three to four times per week for FVIII and two to three times per week for FIX) to maintain factor values above the threshold of efficacy (generally >1%). It represents a serious burden that significantly impacts the patient's quality of life.²⁸ The last decade has seen the development of numerous modified products (EHL factor protein replacements) as well as alternative treatment modalities (alternative factor bypassing therapeutic products and gene therapy) for the treatment of hemophilia. The EHL factor protein replacements products are intended to allow a reduction in the frequency of infusion and improve the standard of care for people with hemophilia. Some of these products have received U.S. Food and Drug Administration (FDA) approval (Table 2) while others are in clinical trials.²⁶ Due to substantial interpatient variability in response to a replacement product infusion adjustment of frequency and level of dosing is often required, resulting in the need for repeated assessments of factor levels in the clinical laboratory.

Assay discrepancy for FVIII assays in FVIII replacement products

The OSA and CSA both accurately assess FVIII activity for pdFVIII. Assay discrepancies between OSA and CSA have been reported for full-length rFVIII products, with the CSA values 8 - 20% higher than the OSA values²⁷⁻³² with the exception of B-domain deletion (BDD) rFVIII. This discrepancy is not considered clinically significant (i.e. within the accepted variation of \leq 20%).⁶ Reports on BDD-rFVIII, however, have shown discrepancies of 20 - 50%, with the OSA lower than the CSA.²⁸⁻³² EHL recombinant factor concentrates have been shown to pose significant challenges to the laboratory monitoring of post-transfusion samples.³³ These factor replacement products contain various molecular modifications (BDD/truncation, fusion with Fc region of IgG or with albumin, or linkage to polyethylene glycol) intended to extend the half-life. Many of the truncated products show wide variation in measured activity levels between different OSAs as well as between the OSA and CSA.^{9,34} However, not all such products have this issue. For example, glycopegylated Novo Eight (N8-GP), a B-domain truncated rVIII, appears to have acceptable recovery across various OSAs with the exception of one reagent.³⁵ Similarly, N8-GP levels are consistently measured across six distinct CSAs.³⁶ A comprehensive review of assay performance among each EHL factor concentrate is beyond the scope of this paper and can be found elsewhere^{3,34,37} In general OSA and CSA can vary their estimation of factor activity by 20-50% for most products which could be clinically relevant, particularly in individuals undergoing treatment for a major bleed or surgery.

Assay discrepancy for FIX assays in FIX replacement products

Similarly to FVIII, the OSA and CSA both accurately assess FIX activity for pdFIX.³⁷⁻³⁹ Differences between OSA and CSA have been reported with two full-length rFIX, with CSA

results consistently about 70% of OSA results in two rFIX concentrates and in postinfusion samples with one of these products.³⁹ This observation has been confirmed in another report on the evaluation of three CSA FIX kits.⁴⁰ A recent study using samples containing glycopegylated rFIX (N9-GP) showed FIX activity was underestimated by 30 – 70% and overestimated by 500 - 2800%, depending on the reagents used in the OSA.⁴¹ Studies on FIX fused with Fc (rFIX-Fc) indicated accurate measurement of postinfusion plasma rFIXFc was APTT reagent activator-dependency in the OSA with ellagic acid resulting in overestimation, and silica (30%) and kaolin (50%) in underestimation.^{42,43} Another study reported comparable results could be achieved between the CSA and the OSA if selected APTT reagents.⁴⁴

It must be noted the vast majority, if not all, of data that has been published on the performance of different OSA reagents and the CSA kits on recombinant and EHL recombinant factor replacement products was collected using factor concentrates diluted to different concentrations in appropriate factor deficient plasma. This data will need to be confirmed in clinical plasma samples collected postinfusion in patients with hemophilia.

Causes for variability in the factor assays

Review of proficiency testing programs and field surveys demonstrates OSA is currently used in the majority of clinical laboratories for the clinical diagnosis of hemophilia and for monitoring factor replacement therapies.⁴⁵⁻⁴⁷ The OSAs for FVIII and FIX activity use a wide variety of APTT reagents differing in PL source, type, concentration, and activator type. The activator type seems to exert a major influence on activity discrepancy within the OSA. Factor deficient plasma may be from congenital-deficient hemophilia patients or, more frequently, factor-specific immunodepleted plasma. Other variables that have been shown to influence the accurate measurement of factor activity are the analyzer and its methodology, the assay protocol, and the calibration material.⁴⁸⁻⁴⁹

The potency labelling for FVIII and FIX concentrate products, including plasma-derived, recombinant and EHL modified products, is performed by the manufacturers, generally using either the OSA or the CSA. The ISTH/SSC has published recommendations for manufacturer's potency labelling.⁵⁰ These recommendations discuss establishment of valid estimates using OSA and CSA, OSA evaluation performed using different APTT reagents, and potency assessment performed relative to the WHO International Standards (WHO IS). The recommendations also state potency assessment should be conducted using a reference plasma which would not necessarily be used for potency assignment, but could be useful when considering the use of a plasma reference to monitor recovery of new products in the clinical laboratory as well as a plasma reference. These recommendations all directly pertain to improving the accuracy of testing in the clinical laboratory. Potency assessment for many of the FVIII replacement products uses CSA. Indeed, the FVIII and FIX sub-committee of the ISTH/SSC and the European Pharmacopoeia, which sets mandatory standards for medicinal products in Europe, defines the FVIII CSA for potency labeling of FVIII medicinal products; however, the OSA is generally preferred in clinical laboratories in Europe and throughout the world. Presently, there is no recommendation for FIX, presumably because the FIX CSA is only recently available. Some of the available FVIII replacements products have shown no discrepancy in results between OSA and CSA while some products do show discrepancy. Almost all of the EHL-FVIII have shown discrepancy which seems to be reagent-dependent. The clinical laboratory must be well-aware of this information. It should refer to the package insert and the published literature

for guidance as what methodology should be used for monitoring as well as the performance of its assays with a particular factor replacement product, especially the EHL products.

The optimal approach for measurement of factor activity involves the testing against a product reference composed of the same material as that which is infused and the same methodology, that is, a like-versus-like approach for potency assessment. While patient plasma samples at the time of diagnosis and post pd-factor replacement products are assayed versus a like-versus-like reference material traceable to the WHO IS for FVIII or FIX, this concept would be very difficult to implement for the recombinant and EHL factor replacement products. Reference and control material for the recombinant and extended half-life replacement products are not available. To allow appropriate interpretation of laboratory results, it will be necessary for the manufacturer to provide guidance to clinicians whose laboratories use OSA to measure FVIII and FIX activity of products clinically validated and labeled by use of the CSA.⁶ Also, the laboratory must be knowledgeable of the available replacement products as well as the ability of its laboratory assay to accurately measure those products and to communicate such information to the clinician.

Inhibitors

The development of inhibitory alloantibodies following exposure to factor replacement therapy is a major complication of hemophilia. These antibodies are mainly IgG and bind to functional domains on FVIII or FIX, inhibiting its anticoagulant activity.^{51,52} While inhibitors to FIX are uncommon, occurring in approximately 3% of HB patients, anti-human FVIII antibodies develop in approximately 30% of patients with severe HA following exposure to factor replacement. The

presence of an inhibitor results in poor response to standard doses of clotting factor replacement.⁵³ As a consequence, compared to patients without inhibitors, those with inhibitors have more frequent, poorly controlled hemarthroses resulting in more severe joint disease, and are at greater risk for life- or limb-threatening hemorrhage, and have reduced health-related quality of life.⁵⁴⁻⁵⁶

De novo inhibitors to FVIII can also manifest in non-HA patients, in a rare condition called acquired HA (AHA). The majority of AHA cases are idiopathic, while the remaining are associated with autoimmune disease, malignancy, pregnancy and post-partum state.^{57,58} The bleeding phenotype of AHA differs from congenital HA, with soft tissue bleeding being more common than hemarthroses.

Inhibitory antibodies to FVIII can be identified in the laboratory using OSA, most commonly the Bethesda clot-based assay, or enzyme linked immunosorbent assays.⁵⁹ The classic Bethesda assay involves normal pooled plasma as the source of FVIII being incubated in undiluted patient plasma for 2 hrs at 37°C and then assayed for residual FVIII. One inhibitor unit (Bethesda Unit, BU) is defined as the amount of patient plasma that destroys 50% of the FVIII in the mixture, corrected for the deterioration of FVIII in a control consisting of normal plasma incubated with buffer. Positive results may require further dilutions of the patient plasma to be assayed to obtain an accurate result.⁵⁹⁻⁶¹ The Nijmegen modification of this assay has been shown to increase the specificity of low-titer FVIII inhibitors. This modification involves buffering the normal pooled plasma, used in patient and control mixtures, to pH 7.4 with imidazole buffer and using FVIII-deficient plasma in the control mixture for the 2-hour incubation by stabilizing the FVIII in the pooled normal plasma.⁶² Additional modification have been made and evaluated,

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including heating the patient plasma 56-58°C for 60-90 minutes to inactivate any circulating FVIII activity, the appropriated use of FVIII deficient plasma, the use of 4M imidazole solution instead of solid imidazole for buffering, and use of 4% solution of bovine serum albumin in place of FVIII deficient plasma. These as well as the use of CSA to measure the residual FVIII hopefully will continue to improve the laboratory's ability to accurately and reproducibly measure FVIII activity.⁶³⁻⁶⁵

Patients with hemophilia who develop inhibitors (neutralizing anti-drug antibodies) to their factor treatments are particularly difficult to manage. Successful immune tolerance induction (ITI) is the best option and enables use of factor replacement therapy for treating acute or traumatic bleeds.⁵³

Test Costs

A recent article by S. Kitchen et al utilized a computer-based cost analysis model to identify and characterize key cost parameters associated with OSA and CSA for both FVIII and FIX assays.⁶⁴ The authors conclude that the key factors that contribute to costs are factor-deficient plasma and kit reagents for OSA and CSA, respectively. The stability of CSA kit reagents, especially the FIX kit, also limits the cost efficiency compared to OSA. The authors estimate the cost of CSA might be reduced 50 -75% using batch testing, aliquoting and freezing of kit reagents; however, these changes would require validation. A recent publication by Bowyer et al reported that using aliquoting, and freezing and analyzing adequate number of controls in each run, the estimated cost for CSA was 21% more than OSA if 20 samples are analyzed in each batch and, interestingly, 8% cheaper if only one sample is analyzed.⁶⁶ Other findings important to mention

are the perception that CSA are technically complex, the lack of validated protocols by the reagents' manufacturers, and the cost of training.

Conclusion

HA and HB are congenital bleeding disorders, the diagnosis of which hinges are accurate assessment of FVIII and FIX, respectively. The current standard of care is prophylactic infusion of factor replacement products for the severe form (<1% activity) and infusion on demand for the moderate (1 - 5%) activity) and mild (5 - 40%) activity) forms of both HA and HB. Historically, the infusion was achieved first using human donor plasma factor concentrates, and later recombinant forms of factor concentrates. Recently, replacement products have been introduced that contain modifications to the recombinant factors intended to extend their plasma half-life, for the purpose of reducing the frequency of infusion. The OSA assay is the most commonly used test for assessment of FVIII and FIX activity in the clinical laboratory. The CSA is less frequently employed. For the diagnosis of hemophilia reports have shown the OSA and CSA perform very well in detecting severe forms of HA and HB; however, discrepancy is found in the moderate and mild forms, especially the latter, of both HA and HB. Thus, a single method to measure FVIII and FIX may not be sufficient to detect all cases of this disorder. For the assessment of postinfusion factor activity in plasma samples with plasma-derived or recombinant FVIII and FIX replacement products the OSA and CSA perform, with some exceptions, reasonably well. For example, with the introduction of BDD-FVIII some assays became inaccurate using a human FVIII reference plasma. The 'solution' to this problem was the use of a different reference material, requiring the laboratory to perform different FVIII assays for

different purposes. With the new modified EHL-recombinant factor replacement products, discrepancy in results with different OSA reagents as well as between OSA and CSA have been reported. These reports indicate that, dependent on the replacement product monitored and the methodology used, substantial over- or under-estimating of factor activity can occur, resulting in the potential for inappropriate patient management. Thus, the challenge for the laboratory to provide a diagnostic strategy that is sensitive, practical and within budget allowances. To do so will require the clinical laboratory performing factor assays to review the FVIII and FIX assays presently provided in light of the hemophilia factor replacement products used in the medical service to which its service is provided. Good communication between the laboratories and the physician about the product the patient has received is of utmost importance. Looking to the future any clinical laboratory that measures factor activity in hemophilia patients should gain the knowledge about the different factor replacement products, should participate in any dialogue with clinicians and whichever hospital department purchases and dispenses these products, and determine whether their current method for determining factor activity provides an accurate measurement of that particular replacement therapy. For those laboratories that do not have the optimal measuring technique, the treating clinician must be informed about the potential biases and an alternative means of measuring accurate concentration provided.

Bibliography

- Potgieter JJ, Damgaard M, Hillarp A. One-stage vs. chromogenic assays in haemophilia
 A. Eur J Haematol 2015;94 Suppl 77:38-44.
- Barrowcliffe TW. Standardization of FVIII & FIX assays. Haemophilia 2003;9(4):397-402.

- Kitchen S, Kershaw G and Tiefenbacher S. Recombinant to modified factor VIII amd factor IX – chromogenic and one-stage assays issues. Haemophilia 2016;22 Suppl 5: 72-77.
- Kitchen S, Tiefenbacher S, Gosselin R. Factor activity assays for monitoring extended half-life FVIII and factor IX replacement therapies. Semin Thromb Hemost 2017;43(3):331-337.
- Moser KA, Adcock Funk DM. Chromogenic factor VIII activity assay. Am J Hematol 2014;89(7):781-4.
- Peyvandi F, Oldenburg J, Friedman KD. A critical appraisal of one-stage and chromogenic assays of factor VIII activity. J Thromb Haemost 2016;14(2):248-61.
- Barrowcliffe TW, Hubbard AR, Kitchen S. Standards and monitoring treatment. Haemophilia 2012;18 Suppl 4:61-5.
- Viuff D, Barrowcliffe T, Saugstrup T, Ezban M, Lillicrap D. International comparative field study of N8 evaluating factor FVIII assay performance. Haemophilia 2011;17(4):695ba, Pierce GP-702.
- Sommer JM, Moore N, McGuffie-Valentine B, Bardan S, Buyue Y, Kamphaus GD, Konkle. Comparative field study evaluating the activity of recombinant factor VIII Fc fusion protein in plasma samples at clinical haemostasis laboratories. Haemophilia 2014;20(2):294-300.

- Diao L, Li S, Ludden T, Gobburu K. Nestorov I, Jiang H. Population pharmacokinetic modelling of recombinant factor IX Fc fusion protein (rFIXFc) in patients with haemophilia B. Clin pharmacokinet 2014;53(5):467-77.
- Abuchowski A, McCoy JR, Palczuk NC, van Es T, Davis FF. Effect of covalent attachment of polyethylene glycol on immunogenicity and circulating life of bovine liver catalase. J Biol Chem 1977;252(11):3582-6.
- 12. Webster R, Elliott V, Park BK, Walker D, Hankin M, Taupin P, PEG and PEG conjugates toxicity: towards an understanding of the toxicity of PEG and its relevance to PEGylated biologicals. In: Veronese F, editor. PEGylated protein drugs: basic science and clinical applications. Basel, Switzerland: Birkhauser; 2009, p 127-46.
- Harris JM, Chess, RB. Effect of pegylation of pharmaceuticals. Nat Rev Drug Discov 2003;2(3):214-21.
- Kaufman RJ, Powell JS. Molecular approaches for improved clotting factors for hemophilia. Blood 2013;122(22):3568-74.
- 15. Ivens IA, Baumann A, McDonald TA, Humphries TJ, Michaels LA, Mathew P. PEGylated therapeutic proteins for haemophilia treatment: a review for haemophilia caregivers. Haemophilia 2013;19(1):11-20.
- 16. Duncan EM, Duncan BM, Tunbridge LJ, Lloyd JV. Familial discrepancy between the one-stage and two-stage factor VIII methods in a subgroup of patients with haemophiliaA. Br J Haematol 1994;87(4):846-8.

- 17. Cid AR, Calabuig M, Cortina V, Casana P, Haya S, Moret A, Cabrera N, Aznar JA. One-stage and chromogenic FVIII:C assay discrepancy in mild haemophilia A and the relationship with the mutation and bleeding phenotype. Haemophilia 2008;14(5):1049-54.
- Bowyer AE, Van Veen JJ, Goodeve AC, Kitchen S, Makris M. Specific and global coagulation assays in the diagnosis of discrepant mild hemophilia A. Haematologica 2013;98(12):1980-7.
- 19. Pipe SW, Saenko EL, Eickhorst AN, Kemball-Cook G, Kaufman RJ. Hemophilia A mutations associated with 1-stage/2-stage activity discrepancy disrupt protein-protein interactions with the triplicated A domains of thrombin-activated factor VIIIa. Blood 2001;97(3):685-691.
- 20. Trossaert M, Boisseau P, Quemener A, Sigaud M, Fouassier M, Ternisien C, Lefancois-Bettembourg A, Tesson C, Thomas C, Bezieau S. Prevalence, biological phenotype and genotype in moderate/mild hemophilia A with discrepancy between one-stage and chromogenic factor VIII activity. J Thromb Haemost 2011;9(3):524-30.
- 21. Mumford AD, Laffan M, O'Donnell J, McVey JH, Johnson DJ, Manning RA, Kemball-Cook G. A Tyr346→Cys substitution in the interdomain acidic region a1 of FVIII in an individual with FVIII:C assay discrepancy. Br J Haemotol 2002;118(2):589-94.
- 22. Poulsen AL, Pedersen LH, Hvas AM, Poulsen LH, Thykjaer H, Ingerslev J. Assay discrepancy in mild haemophilia A: entire population study in a National Haemophilia Centre. Haemophilia 2009;15(1):285-9.

- Armstrong E, Hillarp A. Assay discrepancy in mild haemophilia A. Eur J Haematol Suppl 2014;76:48-50.
- 24. Oldenburg J, Pavlova A. Discrepancy between one-stage and chromogenic factor VIII activity assay results can lead to misdiagnosis of haemophilia A phenotype. Hamostaseologie 2010;30(4):207-11.
- 25. Kihlberg K, Strandberg K, Rosen S, Ljung R, Astermark J. Discrepancies between onestage clotting assay and the chromogenic assay in haemophilia B. Haemophilia 2017;23(4):620-27.
- 26. Muczynski V, Christophe OD, Denis CV, Lenting PJ. Emerging therapeutic strategies in the treatment of hemophilia A. Semin Thromb Hemost 2017;43(6):581-590.
- 27. Raut S, Sands D, Heath AB, Barrowcliffe TW. Variability in factor VIII concentrate measurement: results from SSC field collaborative studies. J Thromb Haemost 2003;1(9);1927-34.
- 28. Barrowcliffe TW, Raul S, Sands D, Hubbard AR. Coagulation and chromogenic assays of factor VIIII activity: general aspects, standardization and recommendations. Semin Thromb Hemost 2002;28(3):247-56.
- 29. Ingerselv J, Jankowski MA, Weston SB, Charles LA, ReFactor Field Study Participants. Collaborative field study on the utility of a BDD factor VIII concentrate standard in the estimation of BDDr Factor VIII:C activity in hemophilic plasma using one-stage clotting assays. J Thromb Haemost 2004;2(4):623-8.

- Mikaelsson M. Oswaldsson U, Sandberg H. Influence of phospholipids on the assessment of factor VIII activity. Haemophilia 1998;4(4):646-50.
- 31. Pahl S, Pavlova A, Driesen J, Muller J, Potzsch B, Oldenburg J. In vitro characterization of recombinant factor VIII concentrates reveals significant differences in protein content, activity and thrombin activation profile. Haemophilia 2013;19(3):392-8.
- 32. Hubbard AR, Weller LJ, Bevan SA. A survey of one-stage and chromogenic potencies in therapeutic factor VIII concentrates. Br J Haematol 2002;117(1):247-8.
- 33. Kasper CK, Aledort L, Aronson D, Counts R, Edson JR, van Eys J, Fratantoni J, Green D, Hampton J, Hilgartner M, Levine P, Lazerson J, McMillan C, Penner J, Shapiro S, Shulman NR. Proceedings: a more uniform measurement of factor VIII inhibitors. Thromb Diath Haemorrh 1975;34(2):612.
- 34. St. Ledger K, Feussner A, Kalina U. The FVIII plasma activity of rVIII-Single Chain can be measured in both the one-stage and chromogenic substrate assays. (Abstract). J Thromb Haemost 2015:13 Suppl 2:540.
- 35. Hillarp A, Bowyer A, Ezban M, Persson P, Kitchen S. Measuring FVIII activity of glycopegylated recombinant factor VIII, N8-GP, with commercially available one-stage clotting and chromogenic assay kits: a two centre study. Haemophilia 2017;23€:458-65.
- 36. Pickering W, Hansen M, Kjalke M, Ezban M. Factor VIII chromogenic assays can be used for potency labeling and post administration monitoring of N8-GP. J Thromb Haemost 2016;14(8):1579-87.

- 37. Kitchen S, Gray E, Mertens K. Monitoring of modified factor VIII and IX products. Haemophilia 2014;20 Suppl 4:36-42.
- 38. Pouplard C, Trossaert M, Le Querrec A, Delahousse B, Giradeau B. Gruel Y. Influence of source of phospholipids for APTT-based factor IX assays and potential consequences for the diagnosis of mild haemophilia B. Haemophilia 2009;15(1):365-8
- Wilmot HV, Hogwood J, Gray E. Recombinant factor IX: discrepancies between one stage clotting and chromogenic assays. Haemophilia 2014;20(6):891-7.
- 40. Kershaw GW, Dissanayake K, Chen VM, Khoo TL. Evaluation of chromogenic factor IX assays by automated protocols. Haemophilia 2018;24(3):492-501.
- 41. Bowyer AE, Hillarp A, Ezban M, Persson P, Kitchen S. Measuring factor IX acivity of nonacog beta pegol with commercially available one-stage clotting and chromogenic assay kits: a two-center study. J Thromb Haemost 2016;14(7):1428-35.
- 42. Holm PK, Sorensen MH, Hermit MB, Ezban M. The activity of glycoPEGylated recombinant FIX (N9-GP) can be measured in two-stage chromogenic and one-stage clotting assays. (Abstract). J Thromb Haemost 2013;13 Suppl 2:828.
- 43. Bowyer A, Shepherd F, Sampson B, Kitchen S, Makris. The FVIII:C measurement of extended half-life recombinant factor VIII Fc fusion in haemophilia A patients. (Abstract). Research and Practice in Thrombosis and Haemostasis 2018;2 Suppl 1:122.
- 44. Young G, Ezban M, Clausen WHO, Negrier C, Oldenburg J, Shima M. N9-GP activity can be measured using select one-stage clot assays and two-stage chromogenic assays in

clinical samples from adults/adolescents and children with haemophilia B. (Abstract). J Thromb Haemost 2016;14 Suppl 1:57

- 45. Jennings I, Kitchen DP, Woods TAL, Kitchen S, Walker ID, Preston FE. Laboratory performance in the World Federation of Hemophilia EQA programme 2003-2008. Haemophilia 2009;15(2):571-7.
- 46. Favaloro EJ, Meijer P, Jennings I, Sioufi J, Bonar RA, Kichen DP, Kershaw G, Lippi G.
 Problems and solutions in laboratory testing for hemophilia. Semin Thromb Hemost 2013;39(7):816-833 [erratum in: Semin Thromb Hemost 2014;40(1):135].
- 47. Kitchen S, Signer-Romero K, Key NS. Current laboratory practices in the diagnosis and management of haemophilia: a global assessment. Haemophilia 2015;21(4):550-7.
- 48. Butenas S, Parhami-Seren B, Mann KG. The influence of von Willebrand factor on factor VIII activity measurements. J Thromb Haemost 2009;7(1):132-137.
- Verbruggen B, Meijer P, Novakova I, Van Heerde W. Diagnosis of factor VIII deficiency. Haemophilia 2008;14 Suppl 3:76-82.
- 50. Hubbard AR, Dodt J, Lee T, Mertens K, Seitz R, Srivastava A, Weinstein M and on behalf of the factor VIII and factor IX subcommittee of the Scientific and Standardisation Committee of the International Society on Thrombosis and Haemostasis. Recommendations on the potency labelling of factor VIII and factor IX concentrates. J Thromb Haemost 2013;11(5):988-9.

- 51. Hay CR, Brown S, Collins PW, Keeling DM, Liesner R. The diagnosis and management of factor VIII and facor IX inhibitors: a guideline from the United Kingdom Haemophilia Centre Doctor's Organization. Br J Haematol 2016;133(6):591-605.
- 52. Christophe OD, Lenting PJ, Cherel G. Boon-Spijker, Lavergne JM, Boertjes R, de Goede-Bolder A, Goudemand J, Gallard S, d'Orion R, Meyer D, Mertens K. Functional mapping of anti-factor IX inhibitors developed in patients with severe hemophilia B. Blood 2001;98(5):1416-23.
- 53. Leissinger CA. Prevention of bleeds in hemophilia patients with inhibitors: emerging data and clinical direction. Am J Hematol 2004;77(2):187-93.
- 54. Quintana-Molina M, Martinez-Bahamonde F, Gonzalez-Garcia E, Romero-Garrido J, Villar-Camacho A, Jimenez-Yuste V, Fernandez-Bello I, Hernandez-Navarro F. Surgery in haemophilic patients with inhibitor; 20 years of experience. Haemophilia 2004;10 Suppl 2:30-40.
- 55. Rodriguez-Merchan EC. Surgery in haemophilic patients with inhibitors. Haemophilia 2004;10 Suppl 2:1-2.
- 56. Scalone L, Mantovani L, Mannucci P, Gringeri A, SOCIS Study Investigators. Quality of life is associated to the orthopaedic status in haemophilic patients with inhibitors. Haemophilia 2006;12(2):154-62.
- 57. Knoebl P, Marco P, Baudo F, Collins P, Huth-Kuhne A, Nemes L, Pellegrini F, Tengborn L, Levesque H, EACH2 Registry Contributors. Demographic and clinical data in

acquired hemophilia A: results from the European Acquired Haemophilia Registry (EACH2). J Thromb Haemost 2012;10(4):622-31.

- 58. Collins P, Hirsch S, Baglin T, Dolan G, Hanley J, Makris M, Keeling DM, Liesner R, Brown SA, Hay CR, UK Haemophilia Centre Doctors' Organisation. Acquired hemophilia A in the United Kingdom: a 2-year national surveillance study by the United Kingdom Haemophilia Centre Doctors' Organisation. Blood 2007;109(5):1870-7.
- 59. Peerschke EI, Castellone DD, Ledford-Kraemer M, Van Cott EM, Meijer P. Laboratory assessment of factor VIII inhibitor titer: the North American Specialized Coagulation Laboratory Association experience. Am J Clin Pathol 2009;131(4):552-8.
- Kenshaw G, Favaloro EJ. Laboratory identification of factor inhibitors: an update. Pathology 2012;44(4):293-30.2.
- 61. Miller CH, Rice AS, Boylan B, Shapiro AD, Lentz SR, Wicklund BM, Wicklund BM, Kelly FM, Soucie JM, Hemophilia Inhibitor Research Study Investigation. Comparison of clot-based, chromogenic and fluorescence assays for measurement of factor VIII inhibitors in the US Hemophilia Inhibitor Research Study. J Thromb Haemost 2013;11(7):1300-9.
- 62. Verbruggen B, Novakova I, Wessela H, Boezeman J, van den Berg M, Mauser-Bunschoten E. The Nijmegen modification of the Bethesda assay for factor VIII:C inhibitors: improved specificity and reliability. Thromb Haemost 1995;73(2):247-51.
- Favaloro EJ, Verbruggen B and Miller CH. Laboratory testing for factor inhibitors. Haemophilia 2014;20 Suppl 4:94-98.

- 64. Verbruggen B, van Heerde W, Novakova I, Lillicrap D and Giles A. A 4% solution of bovine serum albumin may be used in place of factor VIII:C deficient plasma in the control sample in the Nijmegen modification of the Bethesda factor VIII:C inhibitor assay. Thromb Haemost 2002;88:362-4.
- 65. Kitchen S, Blakemore J, Friedman KD, Hart DP, Ko RH, Perry D, Platton S, Tan-Castillo D, Young G, Luddington RJ. A computer-based model to assess costs associated with the use of factor VIII and factor IX one-stage and chromogenic activity assays. J Thromb Haemost 2016;14(4):757-64.
- 66. Bowyer AE, Duncan EM, Antovic JP. Role of chromogenic assays in haemophilia A and B diagnosis. Haemophilia 2018;00:1-6 https//doi.org/10.1111/hae.13520.

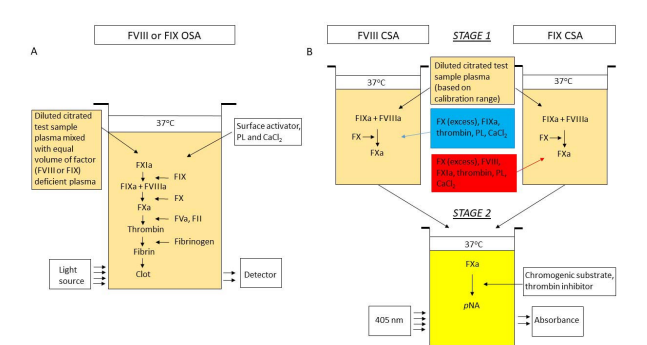
Legends:

Fig. 1. Schematics of FVIII and FIX activity assays. (A) One-stage clotting assay. (B) Chromogenic substrate assay. CaCl₂, calcium chloride; CSA, chromogenic substrate assay; FII, prothrombin (factor II); FIX, factor IX; FIXa, activated factor IX; FVIII, factor VIII; FVIIIa, activated factor VIII; FX, factor X; FXa, activated factor X; FXIa, activated factor XI; PL, phospholipid; *p*NA, *para*-nitroanaline.

Table 1. List of available Chromogenic Substrate kits. (FDA, U.S. Food and Drug Administration; RUO, research use only; N.R., not reported; ^aHigh range and low range calibration curves suggested; ^bSingle calibration curve; ^cValue obtained from reagent package insert; ^dValue for low range calibration curve listed)

Table 2. List of approved EHL products. (EHL, extended half-life; FDA, U.S. Food and Drug Administration; FIX, factor IX; FVIII, factor VIII; PEG, polyethylene glycol;)





Factor VIII Kits				
Name	Manufacturer	FDA status	Detection Limit ^{c,e}	
Biophen FVIII:C ^a	Hyphen-Biomed	None (RUO)	2%	
Coamatic Factor VIII ^a	Chromogenix	Approved	0.5%	
Coatest SP FVIII ^b	Chromogenix	Approved	1%	
Coatest SP4 FVIII ^b	Chromogenix	Approved	1%	
FVIII Chromogenic Assay ^b	Siemens Healthcare	Approved	NR	
Hemosil Electrachrome FVIII ^a	Insturmentation Laboratory	None (RUO)	0.5%	
Technoclone FVIII:C ^a b	Technoclone	None (RUO)	0%	
Factor IX Kits				
Name	Manufacturer	FDA status	Detection Limit	
Biophen Factor IX ^a	Hyphen-Biomed	None (RUO)	0.5%	
ROX Factor IX ^a	Rossix	None (RUO)	0.5%	

Table 1. Chromogenic Substrate Assay Kits

\mathbf{D} VIII V:

Table 2. FDA approved EHL products				
Variant Name	Manufacturer	Year approved	Half-Life	
EHL-FVIII products				
Adynovate (Amino-group PEGylation of full-length FVIII)	Shire	2015	14.7 hours	
Eloctate (Fc-fusion with BDD FVIII)	Bioverativ	2014	19.0 hours	
EHL-FIX products				
Alprolix (Fc-FIX fusion)	Bioverativ	2014	82.1 hours	
Idelvion (Albumin-FIX fusion)	CSL Behring	2015	90.0 hours	
N9-GP, Rebinyn, Refixia, (GlycoPEGylated-FIX)	Novo-Nordisk	2017	92.7 hours	

