

International Journal of Pharmacy and Natural Medicines



Journal Home Page: www.pharmaresearchlibrary.com/ijpnm

RESEARCH ARTICLE

Inhibitor Screening and Quantification of Inhibitor Titres by Bethesda Inhibitor Assay in Patients with Coagulation Disorders

Siva Priya S¹, Uma A N¹*, Lavanya P¹, Kavitha K¹, RakheeKar², Sajini Jacob Elizabeth²

¹Faculty of Allied Health Sciences, Sri Balaji Vidyapeeth, (Deemed to be University), Puducherry-607402. ²Department of Pathology, JIPMER, Puducherry, India.

ABSTRACT

Inhibitor may arise in patients with inherited coagulation disorders or as secondary auto antibodies in previously healthy individuals. This study was conducted over a period of 1 year January 2017 to December 2017 in a large tertiary care centre in Southern India to determine the frequency and the titre of inhibitors among patients with coagulation disorders. A total of 40 patients were enrolled for inhibitor screen of whom 36 were known cases of hemophilia A (33) and B (3), three were new cases of hemophilia and one case of acquired hemophilia. Inhibitor screening was positive in 5 cases of hemophilia A and in the case of acquired hemophilia. In 5 cases the inhibitor was late acting & 1 case the inhibitor positive cases among inherited hemophilias was 12.5%. Bethesda assay was standardized done for quantification of inhibitor titre in all cases where the inhibitor screening was positive the five cases of hemophilia A and the titres ranged from 12BU to >1024BU. By standardizing Bethesda assay exact titres of inhibitor could be given which would be helpful in further management.

Keywords: Bethesda Assay, Hemophilia, Inhibitor Screening.

ARTICLE INFO

*Corresponding Author

Uma A N Faculty of Allied Health Sciences, Sri Balaji Vidyapeeth, (Deemed to be University), Puducherry-607402. MS-ID: IJPNM4120



ARTICLE HISTORY: Received 20 March 2019, Accepted 08 Oct 2019, Available Online 15 December 2019

Copyright© 2019 Siva Priya S, et al. Production and hosting by Pharma Research Library. All rights reserved.

This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution and reproduction in any medium, provided the original work is properly cited.

Citation: Siva Priya S, et al. Inhibitor Screening and Quantification of Inhibitor Titres by Bethesda Inhibitor Assay in Patients with Coagulation Disorders. Int. J. Pharm. Natural Med., 2019, 7(2): 78-83.

CONTENTS

1. Introduction	. 79
2. Materials and Methods.	. 79
3. Results and Discussion.	. 79
4. Conclusion	83
5. References	.83

1. Introduction

The most common coagulation disorders are hemophilias (inheritedor acquired). Hemophiliacs develop bleeding episodes particularly in severe cases and are treated with missing factor replacement therapy i.e. factor VIII or factor IX concentrates¹⁽¹⁾. Inhibitors are acquired de-novo or develop in inherited hemophilias. Acquired inhibitors are associated with an autoimmune disease; it affects mainly elderly people, who present with hemorrhage in the skin. muscle and mucosal membrane. On the other hand inhibitors developing in inherited factor deficiencies occurs early after the beginning of therapy (within <30 exposure days) and most frequent in young hemophiliacs. Patients with inhibitors manifest with excessive bleeding in unusual parts of the body and poor factor therapy recovery⁽²⁾. The inhibitors are IgG antibodies that are against specific deficient factor.

The development of inhibitors is the complication of hemophilia because it occurs shortly after replacement therapy⁽³⁾. Inhibitors are most common in hemophilia A than in hemophilia B but the principles are same for both. Since inhibitors are life threatening cause in hemophiliacs they should be screened for inhibitors in order to prevent serious bleeding and complication⁽³⁾. The inhibitor screening is usually based on a-PTT done on patient plasma mixed with control plasma. The mixing study prolongs the a-PTT in a mixture of patient and normal pooled plasma after incubation at 37°C in presence of inhibitors however FIX and non specific inhibitors are immediate acting, prolonged immediately before incubation whereas FVIII inhibitors aretime dependent, prolonged after incubation at 37°C for 120 minutes⁽⁴⁾. The quantification of inhibitors is done by Bethesda assay. One Bethesda Unit (BU) is defined as the amount of an inhibitor that will neutralize 50% of 1 unit of FVIII: C in normal plasma after 120 minutes incubation at $37^{\circ}C^{(4)}$. It is important that **all people** with hemophilia who use clotting factor concentrates and other bypassing agents are tested for inhibitors at least once a year to prevent from life threatening complication ⁽⁵⁾. Inhibitor titers help to assess whether treatment is working or not working $^{(5)}$.

2. Materials and Methods

Study setting:

The present study was a Cross sectional descriptive study, which was conducted over a period of one year from January 2017 to December 2017 in Department of Pathology (Hematology section), Jawaharlal Institute of Postgraduate Medical Education and Research (JIPMER).

Study participants: Human

Sample size: 40

Initially, 60 samples with prolonged PT &/ APTT were scrutinized. This includes new cases as well as cases on follow up with suspected inhibitors. Patients on anticoagulants and those suspected to have lupus anticoagulant were excluded at the outset. Most of the samples had a $\frac{1}{2}$ patient + $\frac{1}{2}$ control mixing assay. Of these 20 samples were corrected with factor deficient plasma and were having factor deficiency which was excluded. Finally

International Journal of Pharmacy and Natural Medicines

CODEN (USA): IJPNRC | ISSN: 2321-6743

40 samples were put up for inhibitor screening because they were either not corrected or partially corrected on mixing assay or they were clinically suspected to have inhibitors. All cases positive in the inhibitor screen were put up for Bethesda assay.

Bethesda Assay

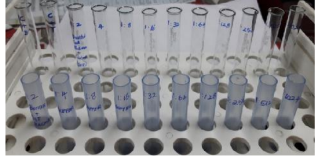
The tubes were taken as follows:

Tube 1: $150\mu l$ control (PNP) + $150\mu l$ of buffer (Owren-koller)

Tube 2: 150µl of control plasma (PNP) + 150µl of test plasma

Tube 3 to 12: 150μ l of control plasma (PNP) in all the tubes + 150μ l of respective diluted test plasma from 1:2 to 1:1024.

Figure 1: From tube 1 to 12



Incubated for 2 hr in water bath at 37°C and performed a Factor VIII Assay on each incubation mixture by using 1/5 dilution

All the tests procedure were done as per the following lab Standard Operative Procedure but we used automated coagulation analyzer –STA Compact (Diagnostic STAGO), here all the test was based on the principle of manual method however instead of water bath a platform with peltier effect is placed, the result is based on viscosity a mechanical detection system enabling immediate delivery of accurate and precise results

Statistical Analysis

The Categorical data like gender, family history, clinical presentation, results of Inhibitor screening were expressed as frequency and percentages. Continuous data like age, values of a PTT and Bethesda Units will be expressed as mean with SD or median with range.

3. Results and Discussion

Study Setting

The study was over a period of 12 months from January 2017 to December 2017, in Haematology section, Department of Pathology at JIPMER, Puducherry.

Sample

Initially, 60 cases with prolonged PT &/ APTT excluding cases of lupus anticoagulant and those under anticoagulants like heparin and warfarin, were screened as per the flow chart given in methods section. Many of these underwent mixing assay and 20 were corrected. These were further mixed with factor deficient plasma and type of factor deficiency was characterized on factor assay. These cases were excluded from further analysis.

Siva Priya S et al, IJPNM, 2019, 7(2): 78-83

A total of 40 samples were put up for Inhibitor screening because they were either not corrected or partially corrected with normal pooled plasma or those cases clinically suspected to have inhibitors. A brief of clinical profile and detailed coagulation work up of these cases is presented.

Distribution of Cases

AGE: The age group of the patients ranged from 1 year to 57 years with a mean of 15 years.

GENDER: Out of 40 cases, 39 were males and 1 was female with acquired hemophilia.

DIAGNOSIS: 40 patients were enrolled for inhibitor screen, 36 were known cases of hemophilia (33 Hemophilia A and 3 Hemophilia B),1 case of acquired hemophilia, 3 were new cases of hemophilia

Clinical Presentation of Patients

The most common clinical presentation in known cases were due to recurrent bleeding, poor factor recovery despite factor replacement, swelling of joints and for routine follow up in cases with prophylactic factor replacement.

Table 1: Clinical presentation of patients

S.no.	Clinical Presentation	No. of cases
1	Follow-up cases	23
	(on prophylaxis)	
2	Poor factor recovery despite	3
	factor replacement	
3	Recurrent bleeding	2
4	Joints swelling	5
5	Hemarthrosis	2
6	Hematoma	3
7	Colicky abdomen pain with	1
	hematuria + clot in urine	

One case diagnosed as acquired hemophilia, on further work-up hadpresented with clinical symptom of bleeds at multiple sites.

Lab Parameters

Inhibitor Screening

All the 40 cases were included for APTT based inhibitor screening. Reading were taken at 0, 1, 2 hrs of fresh mix & incubated mix as shown in table 2.

 Table 2: Reading from inhibitor screening of a positive case.

SAMPLE	'0' Hour	'1' Hour	'2' Hour
Fresh mix	42.1"	44.3°	49.1" 1
Incubated mix	42.1" 🛋	54.5 "	63.5"

As given in the above table, the reading at 1 hr incubated mix is prolonged from 0 hr incubated mix by 12.4 sec and from 1 hr fresh mix by 10.2 sec. The reading at 2 hr incubated mix is prolonged from 1 hr incubated mix by 9 sec and from 2 hr fresh mix by 14.4 sec. As the prolongation seen is more than 8 sec, it is considered positive. Inhibitor screening was positive in 6 cases as International Journal of Pharmacy and Natural Medicines

CODEN (USA): IJPNRC | ISSN: 2321-6743

given in the following table 3. In our cases we found 5 late acting inhibitors which were time and temperature dependent and 1 was showing both late and immediate acting, additionally lupus testing was done and it was negative.

Table 3: Results of inhibitor screening in positive cases

	o states in state of a		1 man come to the set				
	ALSONS CHEARAIRA ALTER peties	150% Distance A 11 constant or when leaves about a 1	AI 1hr from Ohr (incubated mix)	A the fronthe (tresh mx)	At the from the (muthated mix)	At 2hr from 2hr (Loeshonia)	
13		~	12.45 sec	10.25 sec	9 sec	14.4 soc	(at2 hr)
2		~	28.5 აн:	26 sec	12.1 мно	34.1 vec	hate acting (at 2 hr.)
Зs	~		1.9 sec	3.2 sec	16.3 sec	5.8 sec	Both immediat & late acting (at 2 hr)
4		~	21.5 sec	15.9 мгс	12 SHE	26.5 ves	Late acting (at 2 fr.)
h		~	B.4 Net	9.6 vec	4 sec.	5.8 Sec	Late acting (at 1 br)
6*		~	8.3 see:	6.3 vec	4.3 чес	7.3	Late acting (at 1 fr.)

Out of 6 screen positive cases, 5 were hemophilia A and 1 case was acquired hemophilia. The overall prevalence of inhibitors was 15% (6/40). The frequency of inhibitor positive cases among inherited hemophilias was 13.5% (5/40).

Bethesda Assay

Out of 6 positive cases in inhibitor screening test, Bethesda assay was performed in 5 cases according to the procedure mentioned in details in methods section and calculation were done as mentioned below. The 1 case of acquired hemophilia presented on follow up after 2 months when her APTT had normalized and inhibitor screening was negative hence Bethesda assay was not done.

Calculation:

- Choose the dilution of patient's plasma that yields residual factor VIII activity close to 50%.
- The residual factor VIII activity in each is determined using the Factor VIII activity of the control and the respective dilution of the patient's plasma.

The residual factor VIII activity (%)

= <u>Factor VIII activity (patient)</u> × 100 Factor VIII activity (Control)

The residual factor VIII activity is converted to Bethesda unit factor using the following chart (table 1)

Discussion

Inhibitors develops in both inherited and acquired hemophilia, the clinician manages these patients according to the inhibitor titres provided by the coagulation lab⁽¹¹⁾. The development of inhibitors (particularly high titre) is one of the most serious complications of patients under factor replacement therapy leading to poor factor recovery despite factor replacement. Most common complications seen arehemarthrosis, recurrent bleeding, swelling of joints, multiple bleeds, hematoma and hematuria⁽⁶⁾. In our study the inhibitors were found mostly among young patients with a mean age of 15, in other studies it was 13–21 years

Siva Priya S et al, IJPNM, 2019, 7(2): 78-83

in Chicago USA⁽¹²⁾, 11-15 years in Maharashtra India⁽¹³⁾, 18 years in Uttar Pradesh India⁽¹⁴⁾, 17.7 years in India⁽⁷⁾. In our study the prevalence of inhibitor 15%. The frequency of inherited inhibitors were found to be 12.5%, in conjunction with other studies South India (Chennai)-20.99%, Hyderabad-13.33%, Davangere-7.41%, Bangalore- 7.02%⁽⁸⁾. Since hemophilia has X-linked recessive inheritance all cases of inhibitor in hemophilia were males. Only one case of acquired hemophilia was encountered which was a 27 year female⁽¹¹⁾.

Factor inhibitors are time dependent, and the inhibitor will not be detected unless the test is repeated after incubation; hence they are called late acting inhibitor. Nonspecific inhibitors like the lupus anticoagulant usually are not time dependent; the immediate mixture will show prolongation, hence they are immediate acting inhibitors. In this study, 40 cases were enrolled for inhibitor screening, to rule out the immediate and late acting inhibitors. Six out of 40 cases were inhibitor screen positive, 5/6 cases showed late acting inhibitor and in 1/6 casean unexpected finding found to have Inhibitors of both immediate and late acting type. Here

CODEN (USA): IJPNRC | ISSN: 2321-6743

lupus anticoagulant screening was negative. In these cases the common complication was hematoma, and hematuria, hemarthrosis⁽¹¹⁾.

In 1970's, Bethesda assay was developed as the gold standard method for inhibitor determination. The Bethesda assay has become the principal tool for measuring factor inhibitor titres ⁽⁹⁾. So we tried to standardize this Bethesda assay in our lab.Bethesda assay was done in 5/6 cases. It was a one stage assay using normal pooled plasma and buffer. In many studies, patient plasma was diluted with Imidazole bufferand then pooled normal plasmaadded incubated at 37 °C in water bath for 2 hr⁽¹⁰⁾.

In our study we used Owren-koller buffer instead of Imidazole buffer. Bethesda assay was done for the quantification of inhibitor titres. The titre ranges between 12BU to >1024BUand in other studies it was 2.2BU-460.6 BU in Northern part of India⁽⁷⁾. In our study we adopted SOP from an established centre and standardized the Bethesda assay in our setup. The titre value was helpful for the management and treatment of the patients.

Residual VIII %	factor	Residual VIII %	Factor	Residual VIII %	Factor
97	0.05	61	0.7	40	1.35
93	0.1	59	0.75	38	1.4
90	0.15	57	0.8	37	1.45
87	0.2	55	0.85	35	1.5
84	0.25	53	0.9	34	1.55
81	0.3	51	0.95	33	1.6
78	0.35	50	1	32	1.65
75	0.4	48	1.05	30	1.7
73	0.45	46	1.1	29	1.75
70	0.5	45	1.15	28	1.8
68	0.55	43	1.2	27	1.85
66	0.6	42	1.25	26	1.9
64	0.65	41	1.3	25	2

Table 4: Chart to convert the residual factor VIII activity to Bethesda unit

Table 5 – 9 shows the positive cases Bethesda assay calculations Table 5

DILUTION	SEC	FVIII:C	RES. FVIII:C	BU Factor
CONT+ BUFFER	78	64		
CONT+ TEST	108	4	6.25	
1:2	99.9	8	12.5	
1:4	96.8	18	28.1	
1:8	89.2	30	46.8	
1:16	85.6	38	59.3	0.75
1:32	84.3	40	62.5	
1:64	83.6	42	65.6	
1:128	84.5	40	62.5	
1:256	84.5	40	62.5	
1:512				
1:1024				

The residual factor VIII activity (%) Factor VIII activity (patient) × 100 Factor VIII activity (Control)

CALCULATION: 0.75X16(Dil. Factor) = 12BU

Siva Priya S et al, IJPNM, 2019, 7(2): 78-83

CODEN (USA): IJPNRC | ISSN: 2321-6743

Table 6

DILUTION	SEC	FVIII:C	RES. FVIII:C	BU Factor
CONT+ BUFFER	54.2	49		
CONT+ TEST	100.5	1	2.04	
1:2	96.6	1	2.04	
1:4	96.3	1	2.04	
1:8	82.6	3	6.12	
1:16	79.7	5	10.2	
1:32	70.7	10	20.4	
1:64	63.0	20	40.8	
1:128	58.6	30	61.2	0.7
1:256	56.6	38	77.5	
1:512	55.5	42	85.7	
1:1024	55.8	41	83.6	

The residual factor VIII activity (%)

Factor VIII activity (patient) × 100 Factor VIII activity (Control)

CALCULATION:

0.7X128(Dil. Factor) =89.6BU

Table 7

DILUTION	SEC	FVIII:C	RES. FVIII:C	BU Factor
CONT+ BUFFER	52.4	60		
CONT+ TEST	99.7	1	1.66	
1:2	84.6	3	5	
1:4	70.7	10	16.6	
1:8	61.3	23	38.3	
1:16	57.3	35	58.3	0.75
1:32	54.0	50	83.3	
1:64	53.4	54	90	
1:128	52.3	61	101.6	
1:256	53.2	55	91.6	
1:512	52.3	61		
1:1024	51.8	64	106.6	

The residual factor VIII activity (%)

Factor VIII activity (patient) × 100 Factor VIII activity (Control)

CALCULATION:

0.75X16(Dil. Factor) = 12BU

				Table
DILUTION	SEC	FVIII:C	RES. FVIII:C	BU Factor
CONT+ BUFFER	52.8	57		
CONT+ TEST	107.1	1	1.75	
1:2	106.5	1	1.75	
1:4	106.9	1	1.75	
1:8	106.9	1	1.75	
1:16	106.9	1	1.75	
1:32	106.9	1	1.75	
1:64	106.6	1	1.75	
1:128	103.1	1	1.75	
1:256	89.3	2	3.50	
1:512	69.6	11	19.29	
1:1024	60.5	25	43.89	1.2

Table 8

The residual factor VIII activity (%)

Factor VIII activity (patient) × 100 Factor VIII activity (Control)

CALCULATION:

1.2 X 1024(Dil. Factor) = 1228.8 BU

International Journal of Pharmacy and Natural Medicines

DILUTION	SEC	FVIII:C	RES. FVIII:C	BU Factor
CONT+ BUFFER	52.5	59		
CONT+ TEST	106.4	1	1.69	
1:2	107.7	1	1.69	
1:4	106.7	1	1.69	
1:8	107.2	1	1.69	
1:16	107.8	1	1.69	
1:32	108.7	1	1.69	
1:64	106.7	1	1.69	
1:128	106.8	1	1.69	
1:256	101.2	1	1.69	
1:512	91.3	2	3.38	
1:1024	77.8	5	8.47	

Table 9

Factor VIII activity (Control)

The residual factor VIII activity (%) Factor VIII activity (patient) × 100

> RESULT = >1024 BU High inhibitor titres

4. Conclusion

The frequency of inhibitor positive cases among inherited hemophilias was 12.5%. By standardizing Bethesda assay exact titres of inhibitor could be given which was helpful in further management of the hemophilia patients by the clinician.

5. Reference

- [1] Hoot WK, Shapiro AD, Leung LK, et al. Factor VIII and factor IX inhibitors in patients with hemophilia. UpToDate 2011July 11
- [2] Kavaklı K, Aktu lu G, Kemahli S, Ba lar Z, Ertem M, Balkan C, et al. Inhibitor screening for patients with hemophilia in Turkey. Turk J Haematol 2006 Mar 5;23(1):25–32.
- [3] Verbruggen B. Diagnosis and quantification of factor VIII inhibitors. Haemoph Off J World Fed Hemoph. 2010 May;16(102):20–4.
- [4] Phadke S. Hemophilia Care in India: A Review and Experience from a Tertiary Care Centre in Uttar Pradesh. Indian J Hematol Blood Transfus. 2011 Sep;27(3):121–6.
- [5] Miller CH. Laboratory testing for factor VIII and IX inhibitors in haemophilia: A review. Haemoph Off J World Fed Hemoph. 2018;00:1-12.
- [6] Mansouritorghabeh H. Clinical and Laboratory Approaches to Hemophilia A. Iran J Med Sci. 2015 May;40(3):194–205.
- [7] Ghosh K, Shetty S, Kulkarni B, Nair S, Pawar A, Khare A, et al. Development of inhibitors in patients with haemophilia from India. Haemophilia. 2001 May 1;7(3):273–8.
- [8] Lossing TS, Kasper CK, Feinstein DI. Detection of factor VIII inhibitors with the partial thromboplastin time. Blood. 1977, 49(5):793–7.
- [9] Peerschke EIB, Castellone DD, Ledford-Kraemer M, Van Cott EM, Meijer P, NASCOLA Proficiency Testing Committee. Laboratory assessment of factor VIII inhibitor titer: the North American Specialized Coagulation Laboratory

Association experience. Am J Clin Pathol. 2009, 131(4): 552–8.

- [10] Sahud MA. Factor VIII inhibitors. Laboratory diagnosis of inhibitors. Semin Thromb Hemost. 2000, 26(2):195–203.
- [11] Mahmoodi Nesheli H, Hadizadeh A, Bijani A. Evaluation of inhibitor antibody in hemophiliaA population. Casp J Intern Med. 2013;4(3):727–30.
- [12] Leissinger C, Cooper DL, Solem CT, HTRS Investigators. Assessing the impact of age, race, ethnicity and inhibitor status on functional limitations of patients with severe and moderately severe haemophilia A. HaemophOff J World Fed Hemoph. 2011 Nov; 17(6):884–9.
- [13] Kar A, Potnis-Lele M. Descriptive epidemiology of haemophilia in Maharashtra, India. Haemophilia. 2001 Nov 12;7(6):561–7.
- [14] Mishra S, Kumar S, Panwar A, Bhagchandani D, Aneja GK, Verma N, et al. A clinical profile of hemophilia patients and assessment of their quality of life in Western Uttar Pradesh, India: An observational study. Med J DrPatil Vidyapeeth. 2016 May 1;9(3):320.