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# Comparative Study of Factor Xa Fluorogenic Substrates and Their Influence on the Quantification of LMWHs

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43	Abstract	<p>Low molecular weight heparins (LMWHs) are recognised as the preferred anticoagulants in the prevention and treatment of venous thromboembolism. Anti-Factor Xa (anti-FXa) levels are used to monitor the anticoagulant effect of LMWHs and such assays are routinely employed in hospital diagnostic laboratories. In this study, a fluorogenic anti-FXa assay was developed using a commercially available fluorogenic substrate with an attached 6-amino-1-naphthalene-sulfonamide (ANSN) fluorophore and was used for the determination of two LMWHs, enoxaparin and tinzaparin and the heparinoid, danaparoid. The assay was based on the complexation of heparinised plasma with 100 nM exogenous FXa and 25 <math>\mu</math>M of the fluorogenic substrate Mes-D-LGR-ANSN (C<sub>2</sub>H<sub>5</sub>)<sub>2</sub> (SN-7). The assay was tested with pooled plasma samples spiked with anticoagulant concentrations in the range 0–1.6 U mL<sup>-1</sup>. The statistically sensitive assay range was 0–0.4 U mL<sup>-1</sup> for enoxaparin and tinzaparin and 0–0.2 U mL<sup>-1</sup> for danaparoid, with assay variation typically below 10.5%. This assay was then compared with a previously published fluorogenic anti-FXa assay developed with the peptide substrate, methylsulfonyl-<math>\epsilon</math>-cyclohexylalanyl-glycyl-arginine-7-amino-4-methylcoumarin acetate (Pefafluor FXa). Both assays were compared in terms of fluorescence intensity, lag times and sensitivity to anticoagulants.</p>	
44	Keywords separated by ' - '	FXa - Fluorogenic substrate - AMC - ANSN - Low molecular weight heparins - Danaparoid	
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## Comparative study of Factor Xa fluorogenic substrates and their influence on the quantification of LMWHs

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4-methylcoumarin acetate (Pefafleur FXa). Both assays were compared in terms of fluorescence intensity, lag times and sensitivity to anticoagulants.

**Keywords** FXa · Fluorogenic substrate · AMC · ANSN · Low molecular weight heparins · Danaparoid

### Introduction

Low molecular weight heparins (LMWHs) are efficacious anticoagulants administered for both the prophylaxis and treatment of venous and arterial thromboembolic disorders and acute coronary syndromes [1–3]. LMWHs require less laboratory monitoring than unfractionated heparin (UFH) as they exhibit a more predictable pharmacokinetic response with reduced bleeding risk [4]. However, clinical situations do arise where the anticoagulant effect of LMWHs is unpredictable. As a result, laboratory monitoring of LMWHs is necessary in special patient cohorts including pregnant women, the elderly, children, patients with renal insufficiency, and patients at the extremes of body weight [2, 5].

The recommended test for monitoring LMWHs therapy is the anti-Factor Xa (anti-FXa) assay due to the strong inhibition of FXa by LMWHs [5]. The first anti-FXa assay was a clotting time test based on the heparin-accelerated inhibition of FXa [6]. Anti-FXa assays currently employed in the central laboratory setting are chromogenic and use a synthetic FXa peptide substrate coupled to a cleavable chromophore [7] and [2]. The application of synthetic peptide substrates to the analysis of coagulation proteins confers many advantages over traditional clot-based assays, in terms of greater sensitivity, specificity, accuracy, and simplicity [8].

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64 The endopeptidases involved in coagulation and fibrinolysis are trypsin-like serine endopeptidases [9]. Research into the use of synthetic substrates for the evaluation of coagulation endopeptidases began in the 1950s and 1960s. Initially, chromogenic substrates based on the chromophore *p*-nitroaniline were developed for thrombin, plasmin, and trypsin analysis [8, 10], followed by the development of assays using fluorogenic substrates for thrombin [11, 12]. Once the specific peptide cleavage sequence was known, these developments were rapidly extended to other coagulation proteins in the 70s and 80s, resulting in a new strategy for the biochemical study of the coagulation system [10].

65 Factor Xa occupies a critical position in the coagulation cascade; hence its joint popularity with thrombin as a therapeutic target for anticoagulant therapy. The first photometric anti-FXa assay was introduced in 1976 [13] followed by the rapid development of more specific chromogenic substrates and eventually a fluorogenic substrate specific for FXa which was introduced in 1977 [14]. Both chromogenic and fluorogenic assays are well suited to the measurement of heparin, as there is a direct relationship between the functional activity of heparin and its effect on antithrombin allowing for greater precision in its determination [8]. Fluorescence-based measurements offer further advantages over colorimetric assays, such as superior sensitivity and specificity [14, 15], as well as measurement in a broader range of sample types such as platelet poor plasma, platelet rich plasma, and whole blood [16, 13].

66 Two fluorophores are principally used in commercial substrates for the analysis of coagulation proteins, namely 7-amino-4-methylcoumarin (AMC) and 6-amino-1-naphthalene-sulfonamide (ANSN) [17, 18, 19, 20]. The majority of studies have focussed on the reactivity of substrates containing 7-amino-4-methylcoumarin, as it is the most common fluorogenic leaving group used in bioassays [21]. Coumarins can be described as a broad class of fluorophores resulting in UV or near-UV excitation wavelengths with a high quantum yield and low extinction coefficient [22]. The naphthalenesulfonamides were reported in 1992 as a new fluorescent group for substrates of amidases which are also excited in the UV range. Their high quantum yield and the presence of a sulfonyl moiety allows for a range of chemical modifications which can enhance enzymatic substrate specificity [9]. However, given the availability of these substrates, few publications have evaluated their usefulness in the development of anti-FXa assays.

67 A novel fluorogenic anti-FXa assay was developed in this study using a peptide substrate for FXa with an attached ANSN fluorophore and was compared to a previously developed assay based on an AMC fluorescent

68 leaving group [16]. A comparison of assay responses to pharmacological concentrations of LMWHs and danaparoid in commercial pooled plasma is reported.

69 **Materials and methods**

70 **Reagents**

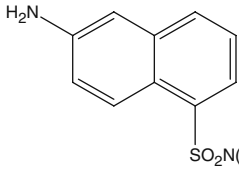
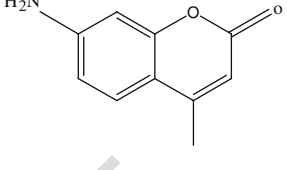
71 Water (ACS reagent) and HEPES (minimum 99.5% titration) were purchased from Sigma-Aldrich (Dublin, Ireland). Filtered HEPES was prepared at a concentration of 10 mM (pH 7.4). A 100 mM filtered stock solution of CaCl<sub>2</sub> from Fluka BioChemika (Buchs, Switzerland) was prepared from a 1 M CaCl<sub>2</sub> solution. The fluorogenic substrate methylsulfonyl-D-cyclohexylalanyl-glycyl-arginine-7-amino-4-methylcoumarin acetate (Pefafluor FXa) was purchased from Pentapharm (Basel, Switzerland). It was reconstituted in 1 mL of water having a final concentration of 10 mM, aliquoted and stored at -20 °C. Dilutions from 10 mM stock solutions down to 10 μM were freshly prepared with water when needed. Subsequent dilutions were prepared in 10 mM HEPES. Tubes were covered with aluminium foil to protect from exposure to light. The fluorogenic substrate Mes-D-LGR-ANSN (C<sub>2</sub>H<sub>5</sub>)<sub>2</sub> (SN-7), containing the fluorescent reporter group ANSN, was acquired from Haematologic Technologies Inc. (Vermont, USA). Stock solutions of 10 mM in DMSO were stored at -20 °C and also protected from light with aluminium foil. Dilutions of stock solutions were performed with 10 mM HEPES. Table 1 summarises the kinetic constants and physico-chemical properties of both the Pefafluor FXa and SN-7 fluorogenic substrates. These parameters were obtained from the suppliers. Kinetic constants refer to the cleavage of both fluorogenic substrates by the endopeptidase FXa. Purified human FXa (serine endopeptidase; code number: EC 3.4.21.6) was obtained from HYPHEN BioMed (Neuville-Sur-Oise, France). Tinzaparin (Innohep®) was obtained from LEO Pharma (Ballerup, Denmark). Enoxaparin (Clexane®) and danaparoid (Orgaran®) were from Sanofi-Aventis (Paris, France) and Schering-Plough (New Jersey, USA), respectively. Human pooled plasma was purchased from Helena Biosciences Europe (Tyne and Wear, UK). Lyophilised plasma was reconstituted in 1 mL of water and left to stabilise for at least 20 min at room temperature prior to use.

72 **Apparatus and software**

73 Fluorescence intensities were measured on an Infinite M200 microplate reader from Tecan Group Ltd. (Männedorf, Switzerland) equipped with a UV Xenon flashlamp. Flat, black-bottom 96-well polystyrol FluorNunc™ micro-



t1.1 **Table 1** Kinetic constants and physico-chemical properties of both fluorogenic substrates

Fluorogenic substrate	SN-7	Pefafleur FXa
<b>Formula</b>	Mes-D-LGR-ANSN(C <sub>2</sub> H <sub>5</sub> ) <sub>2</sub>	CH <sub>3</sub> SO <sub>2</sub> -D-CHA-Gly-Arg-AMC.AcOH
<b>Physical form</b>	Liquid	Lyophilised powder
<b>Fluorophore chemical structure</b>		
<b>M<sub>w</sub> (g mol<sup>-1</sup>)</b>	682.8	679.8
<b>K<sub>m</sub> (μM)</b>	125	220
<b>k<sub>cat</sub> (s<sup>-1</sup>)</b>	36	162
<b>k<sub>cat</sub>/K<sub>m</sub> (M<sup>-1</sup> s<sup>-1</sup>)</b>	290,000	~740,000
<b>Wavelength maxima (nm)</b>	352 (λ <sub>ex</sub> ), 470 (λ <sub>em</sub> )	342 (λ <sub>ex</sub> ), 440 (λ <sub>em</sub> )

164 plates from Thermo Fisher Scientific (Roskilde, Denmark)  
165 were used.

166 Fluorogenic anti-FXa assay

167 Measurements were carried out in reconstituted citrated  
168 human pooled plasma. FXa and the ANSN-based fluoro-  
169 genic substrate were titrated within the range of 0.1–  
170 100 nM and 8.3–33.3 μM (K<sub>m</sub>=125 μM), respectively.  
171 Samples consisting of 6.25 μL of 100 mM CaCl<sub>2</sub>, 43.75 μL  
172 of pooled plasma, and 50 μL of FXa (0.1–100 nM) were  
173 incubated at 37 °C for 3 min and shaken for the first 150 s.  
174 The reaction was started by adding 50 μL of ANSN-based  
175 fluorogenic substrate (8.3–33.3 μM). Samples within wells  
176 were mixed with the aid of orbital shaking at 37 °C for  
177 30 s. Finally, immediately after shaking, fluorescence  
178 measurements were recorded at 37 °C for 60 min with  
179 integration time of 20 μs. Fluorescence excitation was at  
180 352 nm and emission was monitored at 470 nm,  
181 corresponding to the excitation/emission wavelengths of  
182 the ANSN fluorophore. All measurements were carried out  
183 in triplicate. Following optimization of assay conditions,  
184 pooled commercial plasma samples were spiked with  
185 pharmacologically relevant concentrations (0–1.6 U mL<sup>-1</sup>)  
186 of therapeutic anticoagulants including enoxaparin, tinza-  
187 parin, and danaparoid. The reaction rate (slope), which is  
188 defined as the change in fluorescence divided by the change  
189 in time (i.e., dF/dt), was measured as the linear portion of  
190 the fluorescence response profile and plotted versus  
191 anticoagulant concentration. The assay using the Pefafleur  
192 FXa fluorogenic substrate was optimised as previously

described by Harris et al. [16]. The fluorescence excitation  
and emission wavelengths were 342 nm and 440 nm,  
respectively, corresponding to the excitation/emission  
wavelengths of the AMC fluorophore. The only difference  
between these assays was the insertion of an incubation  
step in the ANSN-based assay to improve reproducibility.

Statistical analysis

All graphs were plotted using SigmaPlot 8.0. and SPSS  
17.0 was used for statistical analysis. Intra-assay differences  
within the anticoagulant concentration range were com-  
pared using one-way analysis of variance (ANOVA), with  
subsequent post-hoc analysis performed (Scheffe's test) if  
significance was observed. A result of p<0.05 was  
considered statistically significant. Inter-assay differences  
between the two fluorogenic substrates employed were  
statistically analysed using the paired Student's t test at a  
significance level of 5%.

**Results**

Assay optimisation

In this study, two fluorogenic substrates with two different  
fluorophore leaving groups were compared using the anti-  
FXa assay principle, whereby exogenous FXa added to  
heparinised plasma was inhibited by the heparin-AT  
complex and the resulting FXa activity was measured using  
fluorescence. A new assay is reported that measures the rate

218 of ANSN fluorophore release as a result of FXa substrate  
 219 cleavage and is compared to the rate of AMC fluorophore  
 220 release which has been published previously [16]. Opti-  
 221 mised assay concentrations for this new fluorogenic assay  
 222 were determined by performing titrations of FXa from 0.1  
 223 to 100 nM and fluorogenic substrate concentrations from  
 224 8.3 to 33.3  $\mu\text{M}$ . To allow for differentiation in the assay  
 225 reaction rates, the concentration range of the ANSN-based  
 226 substrate was selected so that it was lower than the  $K_m$   
 227 value. The final working assay concentrations of 25  $\mu\text{M}$   
 228 ANSN-based fluorogenic substrate and 100 nM FXa,  
 229 conform to the optimisation criteria which included short  
 230 lag times, fast reaction rates, and a broad fluorescence  
 231 dynamic range for the differentiation of anticoagulant  
 232 concentrations. The same criteria were also selected for  
 233 the optimization of the AMC-based fluorogenic anti-FXa  
 234 assay.

235 Evaluation of the ANSN-based anti-FXa fluorogenic assay

236 Three anticoagulant drugs were tested using the fluorogenic  
 237 anti-FXa assay with the ANSN-based substrate in commer-  
 238 cial pooled plasma.

239 The fluorescence intensity profiles over time for tinza-  
 240 parin plasma samples in the fluorogenic anti-FXa assay can  
 241 be seen in Fig. 1 and the inset shows the dose–response  
 242 profile versus anticoagulant concentration. As the concen-  
 243 tration of tinzaparin increased, the fluorescence profiles  
 244 reached lower fluorescence intensity values after 1 h. In the  
 245 absence of tinzaparin, the maximum intensity value was  
 246 56,000 AU; with 0.2  $\text{U mL}^{-1}$  the profile reached  
 247 40,000 AU and 27,000 AU for 0.4  $\text{U mL}^{-1}$ . From 0.4 to  
 248 1.6  $\text{U mL}^{-1}$ , the maximum fluorescence intensity value was  
 249 between 22,000 and 27,000 AU.

250 Lag times were calculated by extrapolation of the  
 251 linear portion of the progression curve to its intersection  
 252 with the x-axis [23]. In the presence of higher anticoag-  
 253 ulant concentrations, longer lag time values were ob-  
 254 served. At low concentrations of 0 and 0.2  $\text{U mL}^{-1}$   
 255 tinzaparin, the reaction was very quick and no lag  
 256 time was observed. Lag times increased to 310 and  
 257 630 s for 0.4  $\text{U mL}^{-1}$  and 0.6  $\text{U mL}^{-1}$  respectively. At  
 258 concentrations of 0.8–1.6  $\text{U mL}^{-1}$  lag time values were  
 259 1,010 s.

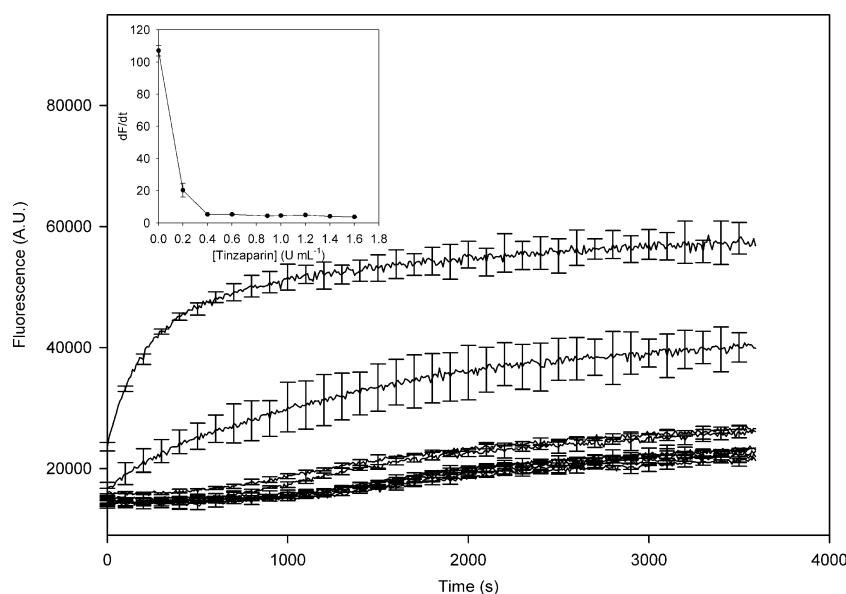
260 The reaction rates of the fluorescence profiles were  
 261 calculated to generate dose–response profiles and a similar  
 262 trend was observed for all drugs analysed, in that as  
 263 anticoagulant concentration increased, the rate of the  
 264 reaction decreased. The slope values were statistically  
 265 different up to 0.4  $\text{U mL}^{-1}$  ( $p < 0.003$ ) with significantly  
 266 reduced sensitivity in the upper tinzaparin range (0.6–  
 267 1.6  $\text{U mL}^{-1}$ ).

268 The anti-FXa assay was also tested with enoxaparin,  
 269 another member of the low molecular weight heparin  
 270 family. The fluorescence profiles are very similar to those  
 271 of tinzaparin (Fig. 2). The fluorescence profiles for  
 272 0  $\text{U mL}^{-1}$  reached 52,000 AU after 1 h, for 0.2  $\text{U mL}^{-1}$   
 273 the intensity value was 38,000 AU and with 0.4  $\text{U mL}^{-1}$  the  
 274 profile reached 28,000 AU. At higher enoxaparin concen-  
 275 trations, the profiles reached maximum intensity values of  
 276 21,000 and 28,000 AU.

277 At low concentrations of enoxaparin lag times were  
 278 absent but were calculated as 110 s for 0.4 and 0.6  $\text{U mL}^{-1}$   
 279 enoxaparin. At higher concentrations, lag times increased to  
 280 590 s for 0.8 and 1  $\text{U mL}^{-1}$ , 730 s for 1.2  $\text{U mL}^{-1}$ , and  
 281 910 s for 1.4 and 1.6  $\text{U mL}^{-1}$ .

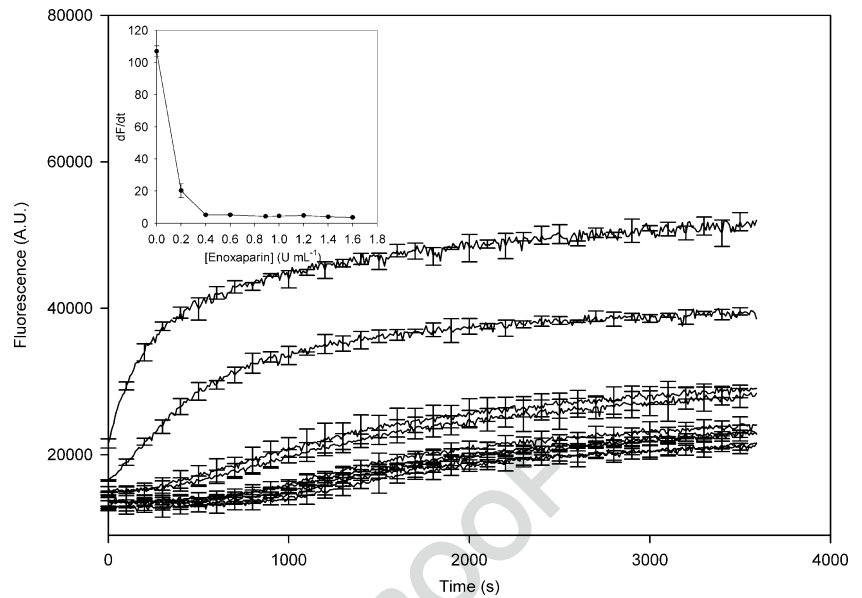
282 The dose–response curve calculated from the linear  
 283 slopes of the fluorescence profiles of enoxaparin can be

**Fig. 1** Fluorescence intensity vs. time for the anti-FXa activity with tinzaparin ( $n=3$ ). Tinzaparin concentration increases from top to bottom. Inset Dose–response calibration curve of tinzaparin in pooled plasma in the ANSN-based fluorogenic anti-FXa assay ( $n=3$ ), calculated from the linear slopes ( $dF/dt$ ) of the fluorescence response curves





**Fig. 2** Fluorescence intensity vs. time for the anti-FXa activity with enoxaparin ( $n=3$ ). Enoxaparin concentration increases from top to bottom. *Inset* Dose-response calibration curve of enoxaparin in pooled plasma in the ANSN-based fluorogenic anti-FXa assay ( $n=3$ ), calculated from the linear slopes ( $dF/dt$ ) of the fluorescence response curves



284 seen in Fig. 2 and good sensitivity was achieved with  
285 concentrations up to  $0.4 \text{ U mL}^{-1}$  ( $p < 0.003$ ).

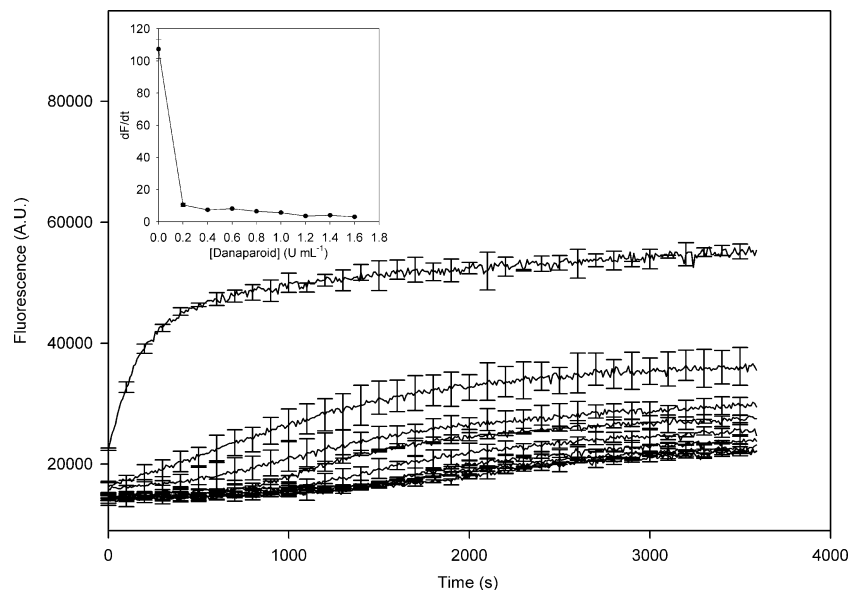
286 Figure 3 shows the fluorescence profiles of the anti-Xa  
287 assay in the presence of danaparoid. The fluorescence  
288 profile for  $0 \text{ U mL}^{-1}$  reached 55,000 AU after 1 h, at  
289  $0.2 \text{ U mL}^{-1}$  the maximum fluorescence intensity value was  
290 35,000 AU and from 0.4 to  $1.6 \text{ U mL}^{-1}$  danaparoid,  
291 profiles reached fluorescence intensities between 22,000  
292 and 29,000 AU.

293 Similar to the LMWHs, lag times were not observed at  
294 low concentrations of danaparoid. Lag times increased  
295 from 200 s at  $0.4 \text{ U mL}^{-1}$  to 780 s at  $0.6 \text{ U mL}^{-1}$ , 1,050 s  
296 at  $0.8 \text{ U mL}^{-1}$  and 1180 s at  $1 \text{ U mL}^{-1}$ . At  $1.2 \text{ U mL}^{-1}$  and  
297  $1.6 \text{ U mL}^{-1}$  danaparoid lag times were 1,320 s and at  
298  $1.4 \text{ U mL}^{-1}$  the lag time was calculated as 1,480 s.

Inset of Fig. 3 shows the dose-response calibration curve  
of danaparoid in the fluorogenic anti-FXa assay. Statistical  
analysis returned significant differences in the variances  
with the Levene's test ( $p=0.011$ ), so one-way ANOVA  
could not be applied. When equal variances were not  
assumed, the Tamhane test returned significant differences  
between 0 and  $0.2 \text{ U mL}^{-1}$  danaparoid ( $p=0.031$ ).

The reaction rates were compared for all drugs tested  
using the normalised data (ratios of the dose-responses  
against the averaged dose-response value at  $0 \text{ U mL}^{-1}$ ). The  
reaction rates for enoxaparin were higher than tinzaparin  
rates at all concentrations from 0 to  $1.6 \text{ U mL}^{-1}$ . Both low  
molecular weight heparins returned higher reaction rates than  
danaparoid in the statistically sensitive range of 0–  
Enoxaparin reaction rates were the highest of

**Fig. 3** Fluorescence intensity vs. time for the anti-FXa activity with danaparoid ( $n=3$ ). Danaparoid concentration increases from top to bottom. *Inset* Dose-response calibration curve of danaparoid in pooled plasma in the ANSN-based fluorogenic anti-FXa assay ( $n=3$ ), calculated from the linear slopes ( $dF/dt$ ) of the fluorescence response curves



314	all drugs tested and only at concentrations of 0.6, 0.8, and	returned higher slopes than the ANSN fluorogenic assay	359
315	1 U mL <sup>-1</sup> , were tinzaparin reaction rates lower than	in all instances. The inter-assay variability between the two	360
316	danaparoid, but these fall into the statistically insensitive	fluorogenic substrates for each anticoagulant drug was	361
317	range.	statistically evaluated using the paired Student's <i>t</i> test.	362
318	The analytical errors (standard deviation and percentage	Results indicate that there is a significant mean difference	363
319	coefficient of variation) related to each of the slope	( <i>p</i> <0.05) between the two fluorogenic anti-FXa assays for	364
320	measurements for all three anticoagulants were calculated.	each anticoagulant at all concentrations, except for	365
321	All CV values were <10.5% with the exception of	0 U mL <sup>-1</sup> as it corresponds to the unit value of normal-	366
322	0.2 U mL <sup>-1</sup> tinzaparin which returned a CV value of 21%.	isation in all cases.	367
323	Comparative studies of AMC versus ANSN-based	<i>Assay sensitivity</i>	368
324	fluorogenic assays		
325	The two synthetic peptide substrates containing two	The intra-assay variability between different concentrations	369
326	different fluorogenic leaving groups were compared in	of anticoagulants was assessed for both fluorogenic	370
327	terms of their fluorescence intensity, lag times, and slope	substrates. Table 3 outlines the statistically sensitive range	371
328	values.	for both fluorogenic anti-FXa assays each containing a	372
329	<i>Fluorescence intensity</i>	different leaving group in the presence of tinzaparin,	373
330	All three of the AMC-based fluorogenic assay reaction	enoxaparin, and danaparoid. When the ANSN-based fluo-	374
331	progress curves for enoxaparin, tinzaparin, and danaparoid,	rogenic anti-FXa assay was performed in the presence of	375
332	reported by Harris et al. [16], approached a similar	tinzaparin, the slopes of the enzymatic reaction were	376
333	maximum plateau value of product formation at approx-	statistically different up to 0.4 U mL <sup>-1</sup> ( <i>p</i> <0.05) at intervals	377
334	imately the same level (i.e., 48,000–53,000 AU) independ-	of 0.2 U mL <sup>-1</sup> . In the case of the Pefaf fluor FXa substrate	378
335	ent of anticoagulant type and concentration. This is an	the assay proved to be sensitive up to 0.6 U mL <sup>-1</sup>	379
336	indication of substrate depletion and also that the enzyme	tinzaparin ( <i>p</i> <0.001). This reduction in assay sensitivity	380
337	remains stable under the conditions tested. In the ANSN-	for the ANSN-based fluorogenic anti-FXa assay was also	381
338	based study, the maximum value of product formed for all	observed for enoxaparin. The fluorogenic anti-FXa assay	382
339	three anticoagulant drugs reached different fluorescence	using the ANSN-based substrate resulted in a narrower	383
340	intensity values depending on anticoagulant concentration.	statistically sensitive range from 0 to 0.4 U mL <sup>-1</sup> ( <i>p</i> <0.05)	384
341	As the concentration of anticoagulant increased from 0 to	when compared with the AMC-based substrate assay range	385
342	1.6 U mL <sup>-1</sup> , the fluorescence intensity values decreased	of 0–0.8 U mL <sup>-1</sup> ( <i>p</i> <0.001) for enoxaparin. The effect of	386
343	from 56,000 to 22,000 for tinzaparin, 52,000 to 21,000 for	danaparoid was also evaluated in both fluorogenic assays	387
344	enoxaparin and finally, from 55,000 to 22,000 for	and a larger difference in assay sensitivity was observed.	388
345	danaparoid.	Response slopes to danaparoid concentrations at	389
346	<i>Lag time values</i>	0.2 U mL <sup>-1</sup> intervals were statistically different up to	390
347	The increase in lag time with increasing anticoagulant	0.2 U mL <sup>-1</sup> and 1 U mL <sup>-1</sup> for the ANSN and AMC-based	391
348	concentration was common to both fluorogenic assays	substrate assays, respectively. In all cases, the AMC-based	392
349	as shown in Table 2. Furthermore, the ANSN-based	substrate seems to offer a wider assay sensitive range	393
350	anti-FXa fluorogenic assay returned smaller lag time	compared with the ANSN-based fluorogenic substrate	394
351	values than the AMC-based assay for all anticoagulant	when quantifying both LMWHs and danaparoid.	395
352	concentrations.	<i>Reproducibility (%CV values)</i>	396
353	<i>Slope values</i>		
354	Analysis of slope values at 0 U mL <sup>-1</sup> returned 25%	Reproducibility of the two developed assays was deter-	397
355	variation in all assays. As a result, all slope values were	mined using the percentage coefficient of variation (%CV).	398
356	normalised with respect to those at 0 U mL <sup>-1</sup> for each	Comparing CV values between the two fluorogenic assays,	399
357	anticoagulant. As can be seen in Fig. 4, the fluorogenic	results in lower variation for both low molecular weight	400
358	anti-FXa assay using the AMC fluorogenic substrate	heparins and danaparoid with the AMC fluorophore.	401
		Studies of precision for the fluorogenic anti-FXa assay	402
		using the AMC-based substrate showed a coefficient of	403
		variation ranging between 0.5% and 7% [16]. In the case of	404
		the ANSN-based assay, CV values were below 10.5%	405
		except for 0.2 U mL <sup>-1</sup> tinzaparin which returned a CV of	406
		21%.	407

Comparative study of Factor Xa fluorogenic substrates

t2.1 **Table 2** Lag times of averaged progress curves for both anti-FXa fluorogenic assays in the presence and absence of tinzaparin, enoxaparin, and danaparoid

Conc (U mL <sup>-1</sup> )	Enoxaparin <sup>a</sup> (s)		Tinzaparin <sup>a</sup> (s)		Danaparoid <sup>a</sup> (s)		
	AMC	ANSN	AMC	ANSN	AMC	ANSN	
0	200	0	160	0	220	0	t2.2
0.2	280	0	240	0	560	0	t2.3
0.4	420	110	380	310	800	200	t2.4
0.6	580	110	670	630	1,420	780	t2.5
0.8	850	590	900	1,010	2,010	1,050	t2.6
1	940	590	1,040	1,010	2,070	1,180	t2.7
1.2	1,020	730	1,440	1,010	2,440	1,320	t2.8
1.4	1,260	910	1,800	1,010	>3,600	1,480	t2.9
1.6	1,310	910	2,000	1,010	>3,600	1,320	t2.10

<sup>a</sup> Averaged of triplicate measurements (n=3)

408 **Discussion**

409 FXa is a serine endopeptidase which occupies a central role  
 410 in the coagulation cascade. LMWHs are anticoagulant  
 411 drugs used in the treatment of thrombosis and cardiovas-  
 412 cular diseases and can effectively neutralise FXa to prevent  
 413 bleeding [24]. At present, commercially available anti-FXa  
 414 tests capable of measuring LMWHs within the therapeutic  
 415 range are all chromogenic assays. While chromogenic anti-  
 416 FXa assays have some advantages over traditional clot-  
 417 based assays, they do still suffer from some drawbacks as  
 418 highlighted by Harris et al. [16]. To counteract these  
 419 limitations, they presented a new fluorogenic anti-FXa  
 420 assay based on the AMC fluorogenic substrate which can  
 421 differentiate and quantify UFH, enoxaparin, tinzaparin, and  
 422 danaparoid within the therapeutic range and with CV values  
 423 below 7%.

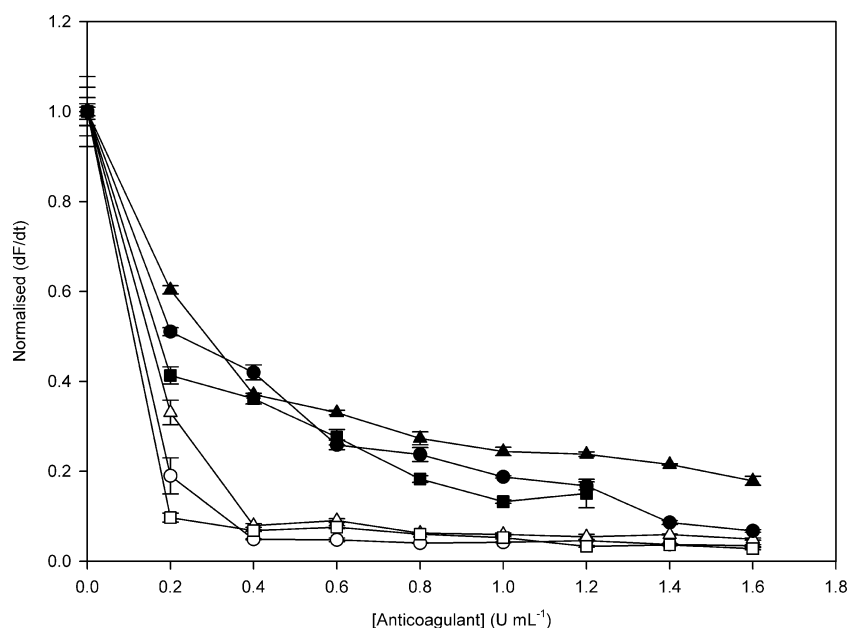
424 A number of studies have been devoted to the  
 425 identification of specific FXa fluorogenic substrates with

desirable properties, such as the photo and chemical  
 stability of the generated fluorescent group, the Stokes  
 shift, the solubility of substrate and fluorescent product in  
 aqueous buffer, and the fluorescence quantum yield, to  
 mention just a few [9]. To date, several synthetic fluoro-  
 genic substrates targeting FXa have been developed with  
 different peptidic groups, attached to just two fluorescent  
 detecting groups namely ANSN [9] and AMC [14].

This paper sought to develop a new fluorogenic anti-FXa  
 assay using the commercially available ANSN-based fluo-  
 rogenic substrate for FXa and compare it with the AMC-  
 based fluorogenic anti-FXa assay previously investigated by  
 Harris et al. [16] in the presence and absence of two  
 LMWHs (i.e., enoxaparin and tinzaparin) and danaparoid.

The principle of the assay is to measure the rate of  
 fluorescent tag release which results from FXa proteolysis  
 of the two synthetic peptide substrates, in the presence and  
 absence of three anticoagulants. From the results obtained,  
 different observations can be made: firstly, the anti-FXa

**Fig. 4** Dose–response curves (normalised data) of the fluorogenic: ANSN-based anti-FXa assay to tinzaparin (empty circle), enoxaparin (empty upright triangle), and danaparoid (empty square); AMC-based anti-FXa assay to tinzaparin (filled circle), enoxaparin (filled upright triangle), and danaparoid (filled square)



**Table 3** Comparison of the statistically sensitive range for each anticoagulant tested in two fluorogenic anti-FXa assays based on different fluorophore leaving groups

Anticoagulant drug	Statistically sensitive range (U mL <sup>-1</sup> )	
	ANSN	AMC
Tinzaparin	0–0.4	0–0.6
Enoxaparin	0–0.4	0–0.8
Danaparoid	0–0.2	0–1

assay presented in this paper was statistically differentiated at intervals of 0.2 up to 0.4 U mL<sup>-1</sup> for enoxaparin and tinzaparin. It was also found to be statistically sensitive for danaparoid up to 0.2 U mL<sup>-1</sup>. CV values were <10.5% for all anticoagulant drugs except for tinzaparin at 0.2 U mL<sup>-1</sup>, which returned 21%. The sensitivity and dynamic range of the fluorogenic anti-FXa assay developed in this study are smaller than the values shown by the commercially available chromogenic anti-FXa assays. Most of the chromogenic anti-FXa assays are able to measure concentrations of LMWHs up to 1 U/ml with %CV values between 5% and 10% [16].

Secondly, the reaction progress curves of the ANSN-based assay show a very different time-course of product formation compared with those of the AMC-based assay. In this study, we observed a decrease in fluorophore formation with increasing anticoagulant concentration. In other words, the reaction progress curves reach lower fluorescence intensities as the concentration of anticoagulant increases. This could be indicative of inhibition by the accumulating products, instability of the enzyme or some other component of the assay system, the presence of enzyme-activated irreversible inhibitors [25], i.e., antithrombin (ATIII) in human plasma or the anticoagulants added to the assay. In comparison, the AMC-based fluorogenic assay showed similar maximum plateau values at all concentrations of FXa inhibitors, indicating enzyme stability and fluorogenic substrate depletion. Moreover, the ANSN substrate did not yield as good a signal-to-noise ratio as the AMC-based substrate; however, the former was not consumed as rapidly.

Thirdly, lag time values for the ANSN-based assay were shorter compared with the AMC-based assay for all anticoagulants and at all concentrations. Considering that the  $K_m$  value for ANSN is 125 and 220 μM for AMC, it is expected that the ANSN substrate would have greater binding affinity for FXa (i.e., shorter lag times) than the AMC substrate. It has been suggested by Butenas et al. [9] that this is due to the limited possibility for modification of the AMC group. The lack of an adequate functional group does not allow for optimisation of substrate interaction compared to the presence of a modifiable sulfonyl moiety

in the ANSN group which can enhance substrate specificity. Additionally, lag times were found to increase with increasing anticoagulant concentration. This shows the ability of FXa inhibitors to form a complex with ATIII and FXa. Any FXa not neutralised is free to react with the fluorogenic substrates; therefore lag times can be due to the time taken for the concentrations of the intermediate enzyme–substrate and enzyme–product complexes to rise to their steady-state levels [25].

And finally, the rates of the enzymatic reaction (i.e., rate of fluorophore formation) were higher for the AMC-based fluorogenic anti-FXa assay in all cases. In the presence of enoxaparin the AMC-based fluorogenic assay resulted in reaction rates that were between 3.6- and 4.8-fold higher than the ANSN assay except for 0.2 U mL<sup>-1</sup> which was 1.8-fold higher. Tinzaparin reaction rates were between 2.3- and 8.4-fold higher than ANSN rates, and finally, danaparoid resulted in rates of reaction that were 2.6- to 5.1-fold higher than those in the ANSN assay. This behaviour is related to the catalytic constant ( $k_{cat}$ ) values of FXa with respect to the two synthetic peptide substrates. The  $k_{cat}$  value for the ANSN substrate is 36 and 162 s<sup>-1</sup> for the AMC substrate. Thus, the higher the  $k_{cat}$  value, the faster the enzyme–substrate complex conversion to fluorophore. Hence, these values indicate that FXa should convert the AMC-based fluorogenic substrate into its fluorophore 4.5-fold quicker than the ANSN-based fluorogenic substrate. This explanation is consistent with our findings.

In terms of assay sensitivity and reproducibility, the AMC-based substrate offers a wider assay sensitive range for enoxaparin, tinzaparin, and danaparoid compared with the ANSN detecting group in the fluorogenic anti-FXa assay. The ANSN-based (AMC-based) assay resulted in detection ranges of 0–0.4 U mL<sup>-1</sup> (0–0.8 U mL<sup>-1</sup>) for enoxaparin, 0–0.4 U mL<sup>-1</sup> (0–0.6 U mL<sup>-1</sup>) for tinzaparin, and 0–0.2 U mL<sup>-1</sup> (0–1 U mL<sup>-1</sup>) for danaparoid. In addition, assay reproducibility for both LMWHs and danaparoid when using the ANSN-based substrate was <10.5%, while the AMC-based substrate returned CVs of <7%.

Assay sensitivity as well as assay precision can also be analysed in terms of the  $k_{cat}/K_m$  (“specificity constant”) values of FXa with respect to the two fluorogenic substrates. FXa has a  $k_{cat}/K_m$  of 290,000 M<sup>-1</sup> s<sup>-1</sup> for the ANSN fluorogenic substrate and ~740,000 M<sup>-1</sup> s<sup>-1</sup> for the AMC substrate. The importance of  $k_{cat}/K_m$  is that it determines the specificity of an enzyme for competing substrates. Therefore taking into account these two values, FXa shows a 2.5-fold greater specificity for AMC than for ANSN which translates into a wider assay sensitive range and better assay reproducibility. Therefore, what appears to be a poor substrate in terms of affinity for FXa (i.e. Peflafluor FXa has a relatively high  $K_m$  of 220 μM



540 compared with 125  $\mu\text{M}$  for SN-7), it actually shows an  
 541 overall greater effectiveness for substrate hydrolysis.  
 542 However, it has been suggested by Bromfield et al. [18]  
 543 who performed competitive inhibition assays for FXa with  
 544 the fluorogenic coumarin substrate Boc-IEGR-AMC, that  
 545 substrates with low  $K_m$  values rather than high  $k_{\text{cat}}/K_m$   
 546 values may be better indicators of inhibitor potential for a  
 547 peptidic sequence. Nevertheless, it is a complex area of  
 548 research due to the open and flat architecture of the FXa  
 549 active site [26, 27] which results in its low selectivity for  
 550 peptide substrates [28].

551 Overall, fluorogenic substrates incorporating the AMC  
 552 leaving group are commonly used probes in proteolytic  
 553 assays [29], fluorogenic thrombin generation assays [30,  
 554 31, 32, 33, 34] and more specifically in fluorogenic FXa  
 555 assays [35, 36, 37, 38, 39]. The use of ANSN-based  
 556 fluorogenic assays has also been described, but to a lesser  
 557 extent for activated protein C [19], human factor VIIa and  
 558 factor VIIa-tissue factor [40], and lately, for the *Mytilus*  
 559 *edulis* anticoagulant peptide [41].

560 In summary, a new fluorogenic anti-FXa assay has been  
 561 investigated in human pooled plasma based on the  
 562 fluorescent reporter group ANSN. It was capable of  
 563 statistically differentiating enoxaparin up to 0.4 U  $\text{mL}^{-1}$ ,  
 564 tinzaparin up to 0.4 U  $\text{mL}^{-1}$ , and danaparoid up to  
 565 0.2 U  $\text{mL}^{-1}$  with CVs of <10.5%. Moreover, it was  
 566 compared with the 7-amino-4-methylcoumarin-based fluo-  
 567 rogenic anti-FXa assay previously developed by Harris et  
 568 al. [16]. Results indicate that the AMC-based fluorogenic  
 569 anti-FXa assay is quicker, it has a wider sensitive range and  
 570 it is more precise than the ANSN-based fluorogenic anti-  
 571 FXa assay examined in this study.

572  
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