Absence of in vitro Procoagulant Activity in Immunoglobulin Preparations due to Activated Coagulation Factors

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\textbf{Keywords}
Intravenous immunoglobulin · Subcutaneous immunoglobulin · Procoagulant activities · Factor XIa · Thromboembolic events

\textbf{Summary}
\textbf{Background:} Immunoglobulin (IG) products, including intravenous (IVIG) or subcutaneous (SCIG) immunoglobulins are considered safe and effective for medical therapy; however, a sudden and unexpected increase in thromboembolic events (TE) after administration of certain batches of IVIG products has been attributed to the presence of activated coagulation factors, mainly factor XIa. Our aims were to examine the presence of enduring procoagulant activity during the manufacturing process of IGs, with special focus on monitoring factor XIa, and to evaluate the presence of in vitro procoagulant activity attributed to coagulation factors in different lots of IVIG and SCIG. \textbf{Methods:} Samples of different steps of IG purification, 19 lots of IVIG and 9 of SCIG were analyzed and compared with 1 commercial preparation of IVIG and 2 of SCIG, respectively. Factors II, VII, IX, XI and XIa and non-activated partial thromboplastin time (NAPTT) were assayed. \textbf{Results:} The levels of factors II, VII, IX, X and XI were non-quantifiable once fraction II had been re-dissolved and in all analyzed lots of IVIG and SCIG. The level of factor XIa at that point was under the detection limits of the assay, and NAPTT yielded values greater than the control during the purification process. In SCIG, we detected higher concentrations of factor XIa in the commercial products, which reached values up to 5 times higher than the average amounts found in the 9 batches produced by UNC-Hemoderivados. Factor XIa in commercial IVIG reached levels slightly higher than those of the 19 batches produced by UNC-Hemoderivados. \textbf{Conclusion:} IVIG and SCIG manufactured by UNC-Hemoderivados showed a lack of thrombogenic potential, as demonstrated not only by the laboratory data obtained in this study but also by the absence of any reports of TE registered by the post marketing pharmacovigilance department.

\textbf{Introduction}
Immunoglobulin (IG) products, including intravenous (IVIG) or subcutaneous (SCIG) immunoglobulins, are prepared from pools of human plasma obtained from at least 1,000 individual donors. These products are licensed for treatment of primary and secondary immunodeficiency disorders and some autoimmune and inflammatory diseases [1]. There are substantial differences in the manufacturing process of the IGs; for this reason, these products may vary in their concentration, osmolality, sugar components, sodium content, amino acids and other stabilizing agents as well as in their strategies for viral inactivation [2]. Overall, IG use is considered safe and effective, but common mild to moderate adverse events, including low-grade fever, headache, malaise, nausea, myalgia and urticaria, have been reported [3]. However, with the increasing use of these IGs, more severe side effects (mainly with IVIG) such as acute renal tubular necrosis, aseptic meningitis, and thromboembolic events (TE) have been described [4]. These last manifestations occur at a frequency of 2–3% and are most frequently acute, occurring either during the administration of the
IGs or within the following 24 h. Underlying risk factors of the recipient (advanced age, thrombophilic status) and product factors (dosage, infusion rates, manufacturing process) are the most important matters to be considered as triggering events. In 2010, a sudden and unexpected increase of TE after administration of certain batches of an IVIG product was attributed to the presence of activated coagulation factors, mainly factor XIa (FXIa) [5]. FXI and immunoglobulins co-purify together; thus, additional steps to remove contaminating traces of FXI [6] or an appropriate pasteurization process are required [7]. A recent report from the European Medicines Agency (EMA) informed that the main cause of TE associated with the use of some batches of IVIG is the presence of increased amounts of FXIa in these products [8]. Despite the relatively limited role of FXI in hemostasis, there are increasing evidences that this protease contributes to the development of thromboembolism in humans. Sustained thrombin generation through FIX activation by FXIa may potentiate several pro-thrombotic processes that result in a two-fold increase of the risk of developing TE [9]. Since 1997, the IG products manufactured by UNC-Hemoderivados are obtained from fraction II of Cohn-Onley cold ethanol fractionation, which is produced with material from a pool of plasma donors. This process includes ultrafiltration, pasteurization, disaggregation, and formulation of the final products for intravenous use and pasteurization and formulation of subcutaneous presentations.

These data and the recent recommendations of the European Pharmacopoeia prompted us to conduct this study. Our aims were i) to examine the presence of residual procoagulant activity during the IG manufacturing process, with special focus on FXIa monitoring and ii) to evaluate the presence of ‘in vitro’ procoagulant activity attributed to coagulation factors in different lots of IVIG and SCIG manufactured from 2010 to 2013.

**Material and Methods**

**Materials**

Samples (marked in bold) of the different steps of the IG purification process (fig. 1) were obtained from routine production of seven distinct lots manufactured between 2010 and 2013, and assayed either immediately or kept frozen at –20 °C for no more than 2 months.

19 lots of IVIG and 9 of SCIG, manufactured between 2010 and 2013 in UNC-Hemoderivados, were analyzed and compared with one commercial preparation of 5% IVIG and two of 16% SCIG, respectively. All samples were within their labeled shelf life.

**Quantitative Determination of the Activity of Coagulation Factors**

The activity of FII, FVII, FIX, and FXI was determined with one-stage clotting assays using deficient plasma from Trinity Biotech (Wicklow, Ireland) and an ACL 7000 coagulometer. For each factor, a working reference curve was arranged with a calibrator plasma and dilution buffer. Samples were tested in duplicate, and the results were expressed in IU/ml. The ACL 7000 was used in the mode ‘low calibration curve’ in order to ensure that the values fell within the readable range of the work curve obtained with the reference plasma. Non-quantifiable results were expressed in that manner when the values obtained ranged between 0.02 and 0.015 IU/ml (depending of the factor), which corresponded to the concentration of the lower point of the assay calibration curve.

**Quantitative Determination of FXIa**

FXIa was assayed with a Biophen® chromogenic assay. In this assay, the FXIa present in the tested sample was assessed by the ability to activate FIX to FIXa, which forms an enzymatic complex with FVIIIa, also supplied in excess. This complex activates FX to FXa; the colored product obtained with a specific FXa substrate is directly related to the amount of FXIa to be measured. The released pNA was measured at 405 nm and the results were expressed in ng/ml. The calibrator provided in the kit was used to obtain the calibration curve, yielding values that ranged between 5 and 0.625 ng/ml. Non-quantifiable results were expressed as <0.10 ng/ml, which corresponded to the detection limit calculated by the validation method obtained from 5 assays. Sample blanks were carried out in all assays in order to assure the specificity of the method.

**Determination of Non-Activated Partial Thromboplastin Time (NAPTT)**

NAPTT is a coagulation assay sensitive to the presence of activated coagulation factors, according to the European Pharmacopoeia (Ph. Eur.)(2.6.22). The samples were diluted 1/10 and 1/100 in buffer solution. Human normal plasma was mixed with rabbit brain cephalin and incubated at 37 °C; afterwards, the sample dilutions and CaCl2 were added, and the clotting time was measured. As control, buffer solution was included in the same run of tests. The results were expressed as mean of the duplicates, in seconds, for each dilution assayed. The test was considered valid when the clotting time measured for the control tube (which contained buffer assays instead of sample material) was 200–320 s and the values of the samples yielded results longer than 150 s.

**Statistics**

The results are expressed as mean ± SEM.

**Results**

Samples of 7 batches obtained from 7 different steps of the purification process of IGs were obtained, and the coagulation factors were assayed (table 1).
The results of FXIa and NAPTT are shown in figure 2 and table 2, respectively. Both assays showed a reduction in the thrombogenic potential along with the progress of IG production. In fact, the level of FXIa showed an initial increase of fractions II + III followed by a significant decrease during the subsequent steps of the process. Meanwhile, NAPTT 1/10 and 1/100 behaved in the same way, with the exception that the maximum thrombogenic activity was observed in fractions II + IIIw, compared to the control time (91 ± 7 s vs. 285 ± 19 s).

The levels of the coagulation factors in 19 IVIG and 9 SCIG, including commercial commodities (1 for IVIG and 2 for SCIG) were assayed. The levels of FII, FVII, FIX, FX, and FXI were non-quantifiable in the IVIG presentations of all analyzed products (19 lots and one commercial product) (data not shown). The concentration levels of FXIa (fig. 3a) and NAPTT for IVIG are shown in figure 3b. FXIa in commercial IVIG reached levels slightly higher than the average levels of the 19 batches produced by UNC-Hemoderivados. NAPTT 1/10 and 1/100 yielded similar values in all samples analyzed.

In SCIGs, the levels of FII, FVII, FIX, and FX in the 2 commercial products analyzed; however, an considerable increase of the concentration of FXI (0.95 and 1.98 IU/ml) was detected, which represents an increase of 7 times the average value of this factor in products manufactured by UNC-Hemoderivados (data not shown). Figure 4 shows the values of FXIa (a) and NAPTT (b) in the lots of SCIG. We detected higher concentrations of FXIa in the commercial products, which reached values up to 5 times the average values found in the 9 batches produced by UNC-Hemoderivados. In fact, the values of FXIa ranged from 0.10 to 4 ng/ml but always remained lower than the levels detected in the two commercial products. For NAPTT, all levels were higher than the those recommended by the European Pharmacopeia in all batches produced by UNC-Hemoderivados. The two commercial product presented values lower than 150 s, suggesting a high potential thrombogenic risk.

### Table 1. Levels of the coagulation factors in the different steps of the purification process

<table>
<thead>
<tr>
<th>Samples (n = 7)</th>
<th>IU/ml factor II</th>
<th>IU/ml factor VII</th>
<th>IU/ml factor IX</th>
<th>IU/ml factor X</th>
<th>IU/ml factor XI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plasma pool</td>
<td>0.79 ± 0.30</td>
<td>1.03 ± 0.30</td>
<td>0.72 ± 0.31</td>
<td>1.20 ± 0.40</td>
<td>0.98 ± 0.30</td>
</tr>
<tr>
<td>Fractions II+III</td>
<td>0.87 ± 0.40</td>
<td>3.09 ± 0.40</td>
<td>1.48 ± 0.23</td>
<td>0.51 ± 0.11</td>
<td>1.02 ± 0.40</td>
</tr>
<tr>
<td>Fractions II+IIIw</td>
<td>0.65 ± 0.40</td>
<td>2.95 ± 0.70</td>
<td>1.04 ± 0.54</td>
<td>0.04 ± 0.01</td>
<td>0.87 ± 0.40</td>
</tr>
<tr>
<td>Fraction II</td>
<td>&lt;0.015</td>
<td>0.07 ± 0.02</td>
<td>&lt;0.015</td>
<td>&lt;0.015</td>
<td>0.07 ± 0.02</td>
</tr>
<tr>
<td>Pasteurization</td>
<td>&lt;0.015</td>
<td>&lt;0.015</td>
<td>&lt;0.015</td>
<td>&lt;0.015</td>
<td>&lt;0.015</td>
</tr>
<tr>
<td>Disaggregation*</td>
<td>&lt;0.015</td>
<td>&lt;0.015</td>
<td>&lt;0.015</td>
<td>&lt;0.015</td>
<td>&lt;0.015</td>
</tr>
<tr>
<td>Formulation</td>
<td>&lt;0.015</td>
<td>&lt;0.015</td>
<td>&lt;0.015</td>
<td>&lt;0.015</td>
<td>&lt;0.015</td>
</tr>
</tbody>
</table>

*Only for IVIG products.

### Table 2. NAPTT during the production process of IGs

<table>
<thead>
<tr>
<th>Samples (n = 7)</th>
<th>NAPTT 1/10, s</th>
<th>NAPTT 1/100, s</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plasma pool</td>
<td>116 ± 12</td>
<td>253 ± 46</td>
</tr>
<tr>
<td>Fractions II+III</td>
<td>116 ± 9</td>
<td>245 ± 23</td>
</tr>
<tr>
<td>Fractions II+IIIw</td>
<td>91 ± 7</td>
<td>237 ± 12</td>
</tr>
<tr>
<td>Fractions II</td>
<td>304 ± 40</td>
<td>333 ± 54</td>
</tr>
<tr>
<td>Pasteurization</td>
<td>297 ± 32</td>
<td>311 ± 39</td>
</tr>
<tr>
<td>Disaggregation*</td>
<td>318 ± 44</td>
<td>327 ± 49</td>
</tr>
<tr>
<td>Formulation</td>
<td>319 ± 38</td>
<td>328 ± 40</td>
</tr>
<tr>
<td>Control tube</td>
<td>285 ± 19</td>
<td>289 ± 19</td>
</tr>
</tbody>
</table>

*Only for IVIG products.
Discussion

From a commercial point view, the Cohn-Oncley cold ethanol fractionation [11] is the most frequently used method for plasma fractionation in the large-scale purification of plasma components, such as IG and albumin products. The original process has undergone considerable modifications over time in order to improve the purity and safety of the products or to isolate determined plasma components. Despite these changes, remaining contaminant proteins present in the IG fractions may affect the quality, safety, and efficacy of the products. In fact, the recently revised Ph. Eur. Monograph on human normal immunoglobulin for intravenous ad-
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Administration requires that the production of IG process includes steps that remove thrombosis-generating agents (coagulation factors and/or zymogens) which mainly catalyze activation of FXI to FXIa [12]. The same recommendation is given in the Ph. Eur. Monograph for SCIG production since the raw material and part of the purification process are similar to IVIG manufacturing [13].

In this study, we systematically investigated the presence of coagulation factors with special emphasis on FXIa during the whole IG purification procedure, as well as in the formulated IVIG and SCIG products. Cold ethanol fractionation has been mostly used for IG production by our group, including some modifications in order to improve the purity and viral safety. Our results showed a lack of thrombogenic material from clotting factors in the different steps of the process. In fact, there is a noticeable drop in the concentration of clotting FII, FVII, FIX, FX, and FXI from fraction II, which reached levels below the limits of quantification in those steps that follow IVIG pasteurization, disaggregation, and formulation. These results were predictable since clotting factors are unlikely to be found at this industrial level of production, and only a very low concentration of FVII (0.17 IU/ml) and FXI (0.07 IU/ml) were found on fraction II. Similar results have been observed by José et al. [14] during the purification process of IVIG Flebo-gamma®DIF. The global test NAPTT was used to determine the presence of activated coagulation factors, allowing for detection of an important prolongation of the coagulation time in fraction II and the subsequent steps of the manufacturing process. These results were in accordance with the levels of FXIa since the NAPTT test is highly sensitive to this factor [15]. Indeed, a dramatic drop in the levels of FXIa below the detection limits of the assay was accompanied by an important prolongation of NAPTT. Both assays clearly demonstrated a high procoagulant activity of fractions II+IIIw and undetectable actions in fraction II as well as after pasteurization, disaggregation and the formulation process.

Our results demonstrated very low levels of coagulation FII, FVII, FIX, FX and FXI in the different batches of IVIG in both, commercial (n = 1) and UNC-Hemoderivados products (n = 19). The presence of FXIa in all the 19 IVIG lots analyzed yielded values ≤ 0.2 ng/ml and the commercial product presented an increase levels slightly higher than the average values of the UNC-Hemoderivados products. However, this increase did not represent a potential thrombogenic load since the level of FXIa found in lots responsible for TE yielded results close to 47 ng/ml [16]. On the other hand, when NAPTT rates were analyzed in all batches of our IVIG products and in those of commercial sources, all results were

Fig. 4. Concentration of FXIa (a) and NAPTT (b) in SCIG preparations.

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above the control time, which clearly indicates a lack of thrombo-
genic potential. These results also allowed us to infer that there is a lack of correlation between NAPPT and FXIa when present in very low concentrations. In fact, it has been recently reported that the NAPPT test has satisfactory sensitivity for detection of levels ≥1 ng/ml of FXIa [10].

The results of the SCIG analysis showed a profile similar to IVIG. The levels of FII, FVII, FIX, and FX were very low in both UNC-Hemoderivados (n = 9) and commercial (n = 2) products, but an important difference in the level of FXI was noted. In fact, this factor presented rates that ranged from 1 to 2 IU/ml in the commercial products compared to <0.015 IU/ml observed in the UNC-Hemoderivados batches. This result can be correlated with the levels of FXIa obtained in the same batches of SCIG analyzed. Indeed, an increase of almost 5 times (expressed in ng/ml) was found in the commercial products, compared to those manufactured in UNC-Hemoderivados. FXI is the zymogene of the active enzyme FXIa, and this conversion may increase during the storage of the plasma source used for IG manufacturing at 4 ºC [17]. We observed considerable variations in the levels of FXIa in our SCIG batches, but these were always lower than values of the commercial products. Similarly, the levels of FXIa also demonstrated a correlation with NAPPT. In fact, we observed that one of the commercial lots with a high concentration of FXIa showed a lower value of NAPPT. In contrast, another batch that showed a low concentration of this factor presented NAPPT levels similar or above that of the control tube.

Despite the large number of companies that produce plasma de-

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Disclosure Statement
The authors are salaried employees of UNC-Hemoderivados.