Heparin Cryoprecipitation Reduces Plasma Levels of Non-Traditional Risk Factors for Atherosclerosis in vitro

E. Meilin, S. Sela, B. Kristal

Aims: To show that heparin cryoprecipitation (HCP), an in vitro method of plasma purification, reduces the levels of in vivo modified proteins and non-traditional risk factors from plasma of atherosclerotic hemodialysis (HD) patients. Methods: HCP was applied to plasma obtained from HD patients and controls, forming a precipitate – cryogel. Levels of fibrinogen, albumin, CRP, TNF-α, IL-6, advanced oxidation protein products, carbonylated fibrinogen and carbonylated albumin were determined in plasma before and after applying HCP and in the cryogel. Results: Treatment of HD plasma with HCP, beyond the significant reduction of the increased levels of all the above-mentioned molecules, reduced fibrinogen, TNF-α, carbonylated fibrinogen and carbonylated albumin to control levels which were simultaneously found in the cryogel. Conclusions: HCP applied to plasma enables the simultaneous precipitation of modified molecules and circulating non-traditional risk factors for atherosclerosis. This study may serve as a base for the future development of a clinical purification technique.
proteins to aggregate, to be less soluble and to precipitate [5, 7, 8]. In addition, oxidation of albumin, for example, will reduce drug-binding properties and antioxidant properties and may induce tissue factor activity in culture of human umbilical vein endothelial cells [9]. Oxidized albumin is able to trigger the oxidative burst of human neutrophils [10, 11] while oxidized fibrinogen loses the ability to form a solid clot [5, 12], may induce platelet aggregation in the circulation, and enhance IL-8 secretion from human umbilical vein endothelial cells [13].

This study aimed to specifically reduce the levels of modified molecules, especially those that could serve as non-traditional risk factors for atherosclerosis. This was achieved by using the newly developed heparin cryoprecipitation (HCP) method, which is based on an interaction between heparin and proteins, combining freezing temperatures and centrifugation. This precipitation of modified molecules and risk factors forms a pellet, called cryogel. The interaction between heparin and fibrinogen in cooling temperatures has previously been described by Smith et al. [14].

The different effects of temperature, centrifugation and heparin, comprising the HCP method, were evaluated on in vitro modified/untreated fibrinogen and albumin. The HCP method was also applied to commercial TNF-α. Plasma before and after HCP, and cryogels were obtained from atherosclerotic HD patients and the levels of various inflammatory mediators were compared to healthy controls.

HCP preferentially reduces the levels of oxidized proteins, and almost no native proteins, from plasma of atherosclerotic patients. Simultaneously, precipitation of non-modified circulating risk factors was observed from atherosclerotic plasma.

Material and Methods

Blood Samples

Blood was drawn from 25 patients (age range 48–73) with end-stage renal disease on chronic HD treatment, three times a week for 4 h. All the included patients were clinically diagnosed as suffering from accelerated atherosclerosis. 20 healthy controls (HC), age- and gender-matched, served as the control group. Blood from HD patients was always drawn before the start of a dialysis session. The inclusion of HC subjects in the study was based upon clinical examination with laboratory confirmation. Informed consent was obtained from all the patients and subjects participating in this study according to the protocol approved by the institutional committee in accordance with the Helsinki Declaration.

In vitro Oxidation of Fibrinogen and Albumin

Fibrinogen and albumin (Sigma-Aldrich, St. Louis, Mo., USA) were oxidized in vitro using the metal-catalyzed oxidation system comprised of iron and ascorbate to yield a highly oxidized (carbonylated) protein, according to Michellis et al. [15]. Briefly, fibrinogen oxidation (5.5–6 mg/ml) was carried out for 5 h at 37°C in PBS containing 5 mM ascorbate (Sigma, St. Louis, Mo., USA) and 100 µM FeCl₃ (Merck, Germany), while albumin oxidation (28–42 mg/ml) was carried out for 52 h at 37°C in PBS containing 25 mM ascorbate and 100 µM FeCl₃. The reactions were stopped at 4°C by the addition of ethylenediaminetetraacetic acid (EDTA) at pH 8.0 (Sigma) to a final concentration of 1 mM. The oxidizing reagents were removed by overnight dialysis, at room temperature, against PBS.

Determination of Carbonyls Content on Proteins

Levels of carbonylated fibrinogen and albumin were determined, according to Michellis et al. [15]. Briefly, samples were derivatized with 2,4-dinitrophenylhydrazine and separated on sodium dodecyl sulfate-polyacrylamide gel electrophoresis gels (SDS-PAGE): 6% for fibrinogen and 10% for albumin. A fixed amount of the in vitro oxidized fibrinogen/oxidized albumin served as a standard [15]. For each protein studied, two gels were run in parallel, one gel was stained for proteins by Coomassie blue, and the other was transferred on to nitrocellulose membrane in transfer buffer (25 mmol/l Tris, 192 mmol/l glycine) for Western blot analysis. The detection of carbonyls was performed with rabbit polyclonal anti-dinitrophenyl serum (Sigma) followed by secondary goat anti-rabbit IgG peroxidase conjugate (Sigma). In order to confirm that the carbonyls detected were on fibrinogen/albumin, several nitrocellulose membranes were stripped in water for 30 min and used again for Western blot analysis of fibrinogen/albumin and to untreated proteins (ta-

Heparin Cryoprecipitation Method

Evaluation of Temperature, Centrifugation and Heparin Effect on Protein Precipitation

The separate effects of the heparin, temperature and centrifugal force, comprising the HCP method, were applied to in vitro oxidized fibrinogen and albumin and to untreated proteins (tables 1, 2).

Solutions of fibrinogen/albumin (Sigma-Aldrich) and their oxidized forms were divided into four tubes after measuring the protein concentration by absorbance at 280 nm. Two tubes contained heparin (14 IU/ml) and two without heparin served as control for each protein (evaluation of the effect of heparin). The concentrations of in vitro fibrinogen/albumin and in vitro oxidized fibrinogen/albumin were measured and compared to their initial concentrations after: a) freezing (–20°C for 24 h) and thawing...
Application of HCP to Plasma

Blood was drawn into tubes containing 14 IU/ml of sodium heparin (Vacuette®, Greiner Bio-One GmbH), centrifuged at 800 g to obtain cell-free plasma (Plasma before). This plasma was frozen for 24 h, then thawed at 4°C and centrifuged again (800 g) at 4°C, followed by the immediate removal of the supernatant (Plasma after). This procedure resulted in a pellet referred by us as cryogel. The cryogel was insoluble in cold temperatures but mostly soluble at 37°C.

Effect of HCP on Commercial TNF-α

Purified TNF-α (Sigma-Aldrich) was dissolved in PBS and used for in vitro studies. The levels of purified TNF-α were determined by enzyme-linked immunosorbent assay (Bender MedSystems GmbH, Vienna, Austria) before and after application of the HCP method. Commercial TNF-α was prepared in tubes containing 14 IU/ml of sodium heparin (Vacuette®, Greiner Bio-One GmbH, Kremsmünster, Austria) (Before HCP), frozen for 24 h (–20°C), then thawed to 4°C and centrifuged again (800 g) at 4°C, followed by the immediate removal of the supernatant (After HCP).

Application of HCP to Plasma

Blood was drawn into tubes containing 14 IU/ml of sodium heparin (Vacuette®, Greiner Bio-One GmbH), centrifuged at 800 g to obtain cell-free plasma (Plasma before). This plasma was frozen for 24 h, then thawed at 4°C and centrifuged again (800 g) at 4°C, followed by the immediate removal of the supernatant (Plasma after). This procedure resulted in a pellet referred by us as cryogel. The cryogel was insoluble in cold temperatures but mostly soluble at 37°C.

Determination of Biochemical and Inflammatory Parameters

Fibrinogen and albumin levels in plasma and cryogel of HD patients and HC subjects were determined using the Cobas Mira analyzer (Roche Diagnostics, Basel, Switzerland) using the K-assy kit (Kamiya Biomedical Co., Seattle, Wash., USA) for fibrinogen and using a colorimetric assay for albumin (Sentinal Diagnostics, Milan, Italy). Plasma and cryogel levels of IL-6, CRP and TNF-α in HD patients and HC were determined using enzyme-linked immunosorbent assay by Bender MedSystems GmbH.

Determination of Advanced Oxidation Protein Products

Advanced oxidation protein products (AOPP) serve as general markers for the evaluation of protein oxidation. AOPP levels in plasma and cryogel of HD patients and HC were evaluated. Determination of AOPP is based on spectrophotometric detection [16]. Plasma samples were diluted 1:5 in PBS; pure acetic acid (99%; Sigma) was added to all tubes and the absorbance at 340 nm was measured immediately. A standard calibration curve was performed using Chloramine-T solution (Sigma, The Netherlands), 0–100 µmol/l, followed by the addition of 1.16 mol/l potassium iodide (Sigma). PBS served as a blank.

Protein Phenotyping of Cryogel

Analysis of the protein contents of the cryogel by sequencing and identifications was carried out by mass spectrometry, in the
Statistical Analysis

Data are expressed as means ± SD. Values were compared by Wilcoxon signed ranks test. A p value < 0.05 was considered statistically significant.

Results

Effect of Temperature, Centrifugation and Heparin on in vitro Protein Precipitation

The separate effects of heparin, temperature and centrifugation on the precipitation of commercial oxidized fibrinogen/albumin vs. control, untreated fibrinogen/albumin were evaluated. The results are shown in table 1 for fibrinogen and table 2 for albumin.

Untreated fibrinogen (UT-FIB), with or without heparin, followed by freezing and thawing (22°C), resulted in the precipitation of a small percentage of fibrinogen (4%), indicating that heparin, freezing and thawing had very little effect on the precipitation of native UT-FIB. Centrifugation of these tubes (4°C) resulted in additional precipitation of the UT-FIB, but to a much greater extent in the presence of heparin (11 vs. 1%).

Freezing and thawing of the in vitro oxidized fibrinogen (FIB-CO), with and without heparin, resulted in about 20% precipitation of the protein. The addition of centrifugation resulted in further precipitation of the protein, which was augmented by the addition of heparin (49 vs. 40%). Both FIB-CO and UT-FIB precipitated more in the presence of heparin, although modified fibrinogen precipitated more than the native protein (table 1, n = 4).

The conditions (heparin 14 IU, freezing –20°C, thawing 4°C, centrifugation 800 g, 4°C) where we succeeded in precipitating maximum FIB-CO were used in all the following experiments and are referred to hereafter as HCP. The effect of HCP on the precipitation of commercial fibrinogen is also shown in figure 1a.

The in vitro FIB-CO was precipitated from an average of 484 ± 98 mg/dl before application of HCP to 247 ± 32 mg/dl after the application of HCP (49% decrease), while UT-FIB was precipitated from 512 ± 127 to 439 ± 93 mg/dl (14% decrease) (fig. 1a).

The levels of carbonyls in the UT-FIB and in vitro FIB-CO are depicted in figure 1b. Higher levels of carbonyls per fibrinogen molecule were expressed on in vitro FIB-CO compared to UT-FIB, 3.21 ± 1.45 vs. 0.54 ± 0.47 nmol carbonyl/mg fibrinogen, respectively. HCP significantly reduced the levels of the highly carbonylated fibrinogen molecules, from 3.21 ± 1.45 to 2.30 ± 1.26 nmol carbonyl/mg (28% reduction) while in UT-FIB the lower carbonyl levels were non-significantly reduced from carbonyl levels of 0.54 ± 0.47 to 0.47 ± 0.28 nmol carbonyl/mg; the higher the carbonylation, the greater the precipitation.

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Fig. 1. Effect of HCP on fibrinogen and on in vitro oxidized fibrinogen. a Purified fibrinogen was prepared and divided into two samples: one served as control (untreated; UT-FIB) and the other was oxidized in vitro (FIB-CO). The fibrinogen levels in the two samples were measured before and after the application of HCP method by the Cobas Mira analyzer. The dashed line represents the average results (n = 4). * p < 0.05 before vs. after for both FIB-CO and UT-FIB. b The effect of HCP on precipitation of carbonylated fibrinogen. The levels of carbonyls on untreated purified fibrinogen and on in vitro oxidized fibrinogen were evaluated by SDS-PAGE before and after application of the HCP method. The dashed line represents the average results (n = 4). * p < 0.05 FIB-CO before vs. UT-FIB before; ** p < 0.05 before vs. after.
Commercial untreated albumin (UT-ALB), with or without heparin, followed by freezing and thawing (22°C), resulted in the precipitation of a small percentage of albumin (about 5–7%), indicating that heparin, freezing and thawing had very little effect on the precipitation of native UT-ALB. Cold temperatures and centrifugation of these samples did not add to the precipitation of native albumin (table 2, n = 3).

Freezing and thawing of the in vitro oxidized albumin (ALB-CO), with and without heparin, resulted in about 11–14% precipitation of the protein. The addition of cold temperatures and centrifugation resulted in an additional precipitation of the protein, which was augmented by the presence of heparin (23 vs. 17%). ALB-CO precipitated to a greater extent in the presence of heparin (table 2).

The particular effect of HCP on the precipitation of commercial albumin is also shown in figure 2a.

ALB-CO was precipitated from 3.41 ± 0.7 to 2.58 ± 0.3 g/dl (23% decrease) while UT-ALB was precipitated

### Table 3. Effect of HCP on levels of inflammatory mediators and modified molecules in HD and HC plasma

<table>
<thead>
<tr>
<th></th>
<th>HD plasma before</th>
<th>HD plasma after</th>
<th>Δ</th>
<th>HC plasma before</th>
<th>HC plasma after</th>
<th>Δ</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fibrinogen, mg/dl</td>
<td>664 ± 162*</td>
<td>388 ± 140**</td>
<td>-42%</td>
<td>444 ± 78</td>
<td>283 ± 142**</td>
<td>-36%</td>
</tr>
<tr>
<td>Albumin, g/dl</td>
<td>4.02 ± 0.35*</td>
<td>3.82 ± 0.29**</td>
<td>-5%</td>
<td>4.64 ± 0.26</td>
<td>4.61 ± 0.28</td>
<td>-8%</td>
</tr>
<tr>
<td>CRP, mg/l</td>
<td>28.1 ± 2.1*</td>
<td>23.8 ± 2.0**</td>
<td>-16%</td>
<td>1.3 ± 0.12</td>
<td>1.2 ± 0.10</td>
<td>-8%</td>
</tr>
<tr>
<td>TNF-α, pg/ml</td>
<td>2.70 ± 1.29*</td>
<td>0.58 ± 0.84**</td>
<td>-78%</td>
<td>0.44 ± 0.90</td>
<td>0.34 ± 0.67</td>
<td>-24%</td>
</tr>
<tr>
<td>IL-6, pg/ml</td>
<td>5.8 ± 3.05*</td>
<td>5.13 ± 2.94**</td>
<td>-12%</td>
<td>0.88 ± 0.38</td>
<td>0.87 ± 0.37</td>
<td>-8%</td>
</tr>
<tr>
<td>AOPP, μM</td>
<td>186 ± 63*</td>
<td>157 ± 62**</td>
<td>-16%</td>
<td>69 ± 6.3</td>
<td>63 ± 4.8</td>
<td>-12%</td>
</tr>
<tr>
<td>FIB-CO, nmol Carb/mg protein</td>
<td>1.22 ± 0.58*</td>
<td>0.68 ± 0.46**</td>
<td>-44%</td>
<td>0.35 ± 0.15</td>
<td>0.36 ± 0.21</td>
<td>-14%</td>
</tr>
<tr>
<td>ALB-CO, nmol Carb/mg protein</td>
<td>8.28 ± 3.01*</td>
<td>6.85 ± 2.77**</td>
<td>-17%</td>
<td>3.38 ± 1.66</td>
<td>2.91 ± 1.42</td>
<td>-14%</td>
</tr>
</tbody>
</table>

Results are mean ± SD of the following sample size: fibrinogen, n = 25 for HD, n = 20 for HC; albumin and AOPP, n = 20 for HD, n = 10 for HC; CRP, TNF-α, and IL-6, n = 8 for HD; CRP, TNF-α, and IL-6, n = 4 for HC; FIB-CO and ALB-CO, n = 10 for HD, n = 5 for HC. Before and after relate to the application of the HCP procedure. FIB-CO = Carbonylated fibrinogen; ALB-CO = carbonylated albumin.

* p < 0.05 HD before vs. HC before; ** p < 0.05 before vs. after.
from 3.66 ± 0.5 to 3.46 ± 0.3 g/dl (7% decrease) (fig. 2a).

The levels of carbonyls on untreated and oxidized albumin are depicted in figure 2b. ALB-CO expressed higher levels of carbonyls per albumin molecule compared to UT-ALB and had a higher carbonyl content of 8.82 ± 5.05 vs. 1.65 ± 0.46 nmol carbonyl/mg albumin, respectively. HCP reduced significantly highly carbonylated albumin molecules from 8.82 ± 5.05 to 7.40 ± 3.54 nmol carbonyl/mg (16% decrease) albumin while in UT-ALB the carbonyl levels were not reduced (1.65 ± 0.46 to 1.57 ± 0.62 nmol carbonyl/mg albumin) following HCP; the higher the carbonylation, the greater precipitation.

The difference of the carbonyl content on in vitro FIBCO vs. ALB-CO (8.82 ± 5.05 nmol carbonyl/mg for albumin vs. 3.21 ± 1.45 for fibrinogen) is a result of the different oxidation protocol. Albumin was oxidized for a much longer period and with higher ascorbate contents than fibrinogen (see Material and Methods).

**Effect of HCP on HD and HC Plasma**

Plasma from atherosclerotic patients on chronic HD treatment and from healthy subjects was evaluated before and after the application of HCP. The effect of HCP on inflammatory parameters and modified molecules in HD and HC plasma is stated in table 3. Significantly elevated levels of fibrinogen, CRP, IL-6 and oxidized molecules such as advanced oxidative protein products, carbonylated fibrinogen and carbonylated albumin and significantly lower levels of albumin characterized the HD plasma. Following treatment by HCP, the levels of fibrinogen, albumin, CRP, TNF-α, IL-6, AOPP, carbonylated fibrinogen, and carbonylated albumin significantly decreased in HD plasma. It must be emphasized that the pathologically high levels of fibrinogen and TNF-α in HD plasma were corrected to the normal healthy plasma level. In contrast, the only parameter that decreased significantly in HC treated by HCP was fibrinogen, which did not precipitate below the normal range (200–400 mg/dl). In both groups, the levels of non-carbonylated albumin were hardly affected.

**Effect of HCP on the in vitro Precipitation of TNF-α**

TNF-α is a main inflammatory cytokine involved in inflammatory states. Since high levels of TNF-α were found in HD plasma (table 3), we studied the effect of HCP on the non-traditional risk factor TNF-α precipitation in vitro. TNF-α concentrations before and after application of the HCP method were measured. HCP significantly precipitated commercial TNF-α levels from an average of 2.79 ± 0.87 to 2.03 ± 0.45 pg/ml (–27%).

**Cryogel Composition**

**SDS-PAGE Analysis of Cryogel.** A representative picture of three different concentrations of HD cryogel as analyzed by SDS-PAGE is shown in figure 3. Cryogel was precipitated from HD plasma, separated on SDS-PAGE (8%), stained by Coomassie blue and analyzed. The HD cryogel contained fibrinogen (340 kDa), immunoglobulins (150 kDa), transferrin (76 kDa), and albumin (66 kDa), in addition to other unidentified proteins (fig. 3).

**Mass Spectrometry Analysis of Cryogel.** The main proteins identified in the HD cryogel after analysis by mass spectrometry are depicted in table 4. It can be seen that the cryogel contains fibrinogen, haptoglobin, immunoglobulins, transferrin, and albumin. This analysis is semiquantitative, and the detected proteins shown in table 4 are in a decreasing order as to their relative amounts in the cryogel. This analysis supported the results obtained by the SDS-PAGE (fig. 3).

**Characterization of Fibrinogen/Carbonylated Fibrinogen in Plasma and Cryogel.** Plasma and cryogel of HD and HC were separated on SDS-PAGE (6%) and analyzed by Western blot analysis for fibrinogen and carbonylated fibrinogen (fig. 4a, b). HD plasma contained higher levels of fibrinogen, its degradation products, aggregates and carbonylated forms than form HC plasma. The HCP method reduced all forms of fibrinogen, which are abundant in the cryogel (fig. 4a, b). When comparing cryogel
from HC and HD it is evident that the amount of fibrinogen, its degradation products, aggregates and its carbonylated forms are precipitated to a greater extent in HD than in HC plasma (fig. 4a, b). The quantified fibrinogen and carbonylated fibrinogen levels in plasma are shown in table 3 and in cryogel in table 5. It has to be noted that the oxidation causes higher molecular weight aggregates in HD than in HC (fig. 4b).

Characterization of Albumin/Carbonylated Albumin in Plasma and Cryogel. Plasma and cryogel of HD and HC were separated on SDS-PAGE (10%) and analyzed by Western blot analysis for albumin and carbonylated albumin (fig. 5a, b). HD plasma contained higher levels of carbonylated albumin than HC plasma, which was reduced by the HCP method (fig. 5b). Cryogel from HD showed significantly higher amounts of carbonylated albumin than HC cryogel (fig. 5b). The quantified levels of fibrinogen and its degradation products, aggregates and its carbonylated forms are precipitated to a greater extent in HD than in HC plasma (fig. 4a, b). The quantified fibrinogen and carbonylated fibrinogen levels in plasma are shown in table 3 and in cryogel in table 5. It has to be noted that the oxidation causes higher molecular weight aggregates in HD than in HC (fig. 4b).
albumin and carbonylated albumin are shown in plasma (table 3) and in cryogel (table 5).

A summary of cryogel contents is depicted in table 5. HD cryogel contained significantly higher levels of fibrinogen, CRP, TNF-α, IL-6, AOPP, carbonylated fibrinogen and carbonylated albumin, while albumin levels were similar in both HD and HC cryogels.

Table 5. Cryogel contents: acute phase proteins, cytokines and modified molecules

<table>
<thead>
<tr>
<th></th>
<th>HD cryogel</th>
<th>HC cryogel</th>
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<tbody>
<tr>
<td>Fibrinogen, mg/dl</td>
<td>239 ± 114*</td>
<td>153 ± 66</td>
</tr>
<tr>
<td>Albumin, g/dl</td>
<td>0.38 ± 0.14</td>
<td>0.4 ± 0.12</td>
</tr>
<tr>
<td>CRP, pg/ml</td>
<td>1.0 ± 1.0*</td>
<td>0.1 ± 0.1</td>
</tr>
<tr>
<td>TNF-α, pg/ml</td>
<td>0.84 ± 0.58</td>
<td>0.78 ± 0.33</td>
</tr>
<tr>
<td>IL-6, pg/ml</td>
<td>0.96 ± 0.67*</td>
<td>0.12 ± 0.12</td>
</tr>
<tr>
<td>AOPP, μM</td>
<td>26.6 ± 10*</td>
<td>7.7 ± 4.8</td>
</tr>
<tr>
<td>FIB-CO, nmol Carb/mg protein</td>
<td>7.19 ± 4.6*</td>
<td>&lt;1.5</td>
</tr>
<tr>
<td>ALB-CO, nmol Carb/mg protein</td>
<td>6.8 ± 2.45*</td>
<td>2.37 ± 1.23</td>
</tr>
</tbody>
</table>

Results are mean ± SD of the following sample size: fibrinogen and albumin, n = 15 for HD, n = 10 for HC; CRP, TNF-α, and IL-6, n = 8 for HD, n = 4 for HC; AOPP, FIB-CO and ALB-CO, n = 10 for HD, n = 5 for HC. FIB-CO = Carbonylated fibrinogen; ALB-CO = carbonylated albumin.

* p < 0.05 vs. HC cryogel.

Discussion

HCP, a unique molecular precipitating method, was developed in this study. This method combines the use of heparin, freezing temperatures and centrifugation to preferentially precipitate a cluster of risk factors from atherosclerotic patients’ blood (in vitro).

During the development of the HCP method, we tried to find the optimal conditions to precipitate mainly modified molecules and cardiovascular risk factors while trying to avoid precipitation of native molecules. Preliminary studies evaluating the effect of various concentrations of heparin on precipitation level were carried out. These studies indicated that 14 IU/ml of sodium heparin is the lowest level of heparin required for application of the HCP method (data not shown). Heparin, a highly charged polyanion, is capable of forming soluble and insoluble complexes with proteins, which upon freezing and the application of centrifugal force will accelerate their precipitation [17, 18]. The precipitate or the cryogel is insoluble in cold temperatures but mostly soluble at 37°C, enabling its removal by centrifugation at lower temperatures. We show that heparin complexes of native proteins hardly precipitate, while complexes with these in vitro oxidized proteins (oxidized albumin and oxidized fibrinogen) precipitate better by freezing, thawing and centrifugation.

Plasma of atherosclerotic patients on chronic HD contains high levels of modified molecules and high levels of circulating non-traditional risk factors for atherosclerosis such as AOPP, carbonylated fibrinogen, carbonylated albumin, fibrinogen, CRP, IL-6 and TNF-α. All these parameters, together with the low levels of albumin, reflect a state of systemic oxidation and inflammation in these patients [1–3, 11, 13, 15, 19, 20].

When the HCP method was applied to plasma of HD patients, acute phase proteins and cytokines such as fibrinogen, CRP, IL-6, TNF-α and modified molecules such as AOPP, carbonylated fibrinogen and carbonylated albumin precipitated, while hardly any precipitation was observed in HC plasma. HD plasma contained significantly higher levels of AOPP, CRP, IL-6 and TNF-α levels than HC, which were significantly reduced by the HCP method.

It is interesting to note that in our healthy subjects, fibrinogen levels that were in the upper quartile of the normal range precipitated by the HCP to the lower normal levels. The risk of precipitating native molecules by HCP is one of the hazards of this method. Nevertheless, the described HCP method precipitated threefold more of the oxidized proteins compared to the native ones, with a minimal effect on the native ones. As other studies have shown that fibrinogen and its carbonylated form are in-
flammatory mediators, we imply that significant reduction in their levels from HD plasma may be beneficial in attenuation of the atherosclerotic process and cardiovascular morbidities in these patients [3, 5, 13, 21, 22].

Hypoalbuminemia, frequently observed in HD patients, is a measure of malnutrition, acute phase inflammation, accelerated atherosclerosis (MIA syndrome), and a strong predictor of cardiovascular morbidity and mortality [23–26]. Himmelfarb and McMonagle [19] and Mera et al. [11] have reported that HD patients on chronic HD have lower albumin and higher carbonylated albumin levels than HC. We show that the HCP method is efficient in precipitating particularly the carbonylated albumin, without affecting the levels of the unmodified molecules both in HD patients and HC subjects. This is especially important in view of accumulating data showing that decreased albumin levels predict mortality in all populations, and are a powerful predictor of CVD in chronic HD patients [27]. Oxidized albumin, in addition to being a reliable marker for oxidative stress in HD patients [11], triggers an oxidative burst of human neutrophils and as such serves as a prooxidant too [9–11]. Thereby, the reduction in oxidized albumin levels from HD blood by HCP will contribute to the attenuation of the systemic oxidative stress in these atherosclerotic patients. A lack of interaction between the anionic albumin with the polyanionic heparin may explain the minor effect of heparin on the precipitation of the native albumin, nevertheless the HCP method precipitates better the oxidized form of albumin.

Our findings support the study by Himmelfarb and McMonagle [19] which demonstrated that albumin is the major plasma target of excess oxidation in HD patients. Albumin can accumulate oxidative modifications possibly due to its long turnover [28] in plasma (15–19 days); this, together with the high albumin concentration in plasma, compose albumin as the major plasma antioxidant protein.

AOPP is a surrogate marker of oxidative stress and serves as an indicator for protein oxidation [29]. AOPP also mediates inflammation and triggers oxidative reactions of neutrophils, monocytes and T lymphocytes, which contribute to inflammation and to the impairment of the immune system found in atherosclerosis [30]. The reduction in AOPP levels by HCP from HD plasma, beyond confirming our hypothesis that HCP is an effective method in the reduction of modified molecules, may also be beneficial in reducing the inflammatory state in HD patients.

The reduction of CRP by HCP is also of biological importance. Beyond being a reliable marker for inflammation and a strong predictor of future cardiovascular events [31], CRP is suggested as a mediator of the atherosclerotic process [32]. CRP production, which is predominantly under the control of IL-6, is involved in the induction of adhesion molecule expression in human endothelial cells, is able to activate the complement system [32], and may play a direct role in promoting the inflammatory component of atherosclerosis [31–33].

IL-6 and TNF-α are central cytokines involved in the proinflammatory process. IL-6 is suggested as a predictor of future heart disease and is the only cytokine that can stimulate the synthesis of all acute phase proteins involved in the inflammatory response [21].

TNF-α is regarded as the master regulator of the cytokine cascade. TNF-α induces thrombotic and inflammatory reactions in endothelial cells, and also induces production of IL-6 by these cells [34]. We suggest that the highly significant reduction in TNF-α from HD plasma may result from its binding to heparin [35], forming a complex that is then precipitated by the HCP method.

By reducing the levels of all the mentioned cytokines and inflammatory mediators, a deceleration and retardation of the vicious cycle of oxidative-stress-inducing inflammation will be achieved, finally attenuating the atherosclerotic process.

As a mirror image, the HD cryogel contained significantly higher levels of fibrinogen, CRP, IL-6, AOPP, carbonylated fibrinogen, and carbonylated albumin compared to HC cryogel, confirming the enhanced precipitation of these inflammatory mediators and modified molecules from HD plasma.

Treatment of atherosclerosis is principally by the traditional pharmacological approach which deals mainly with one risk factor at a time, and is only partially successful in the reduction of CVD morbidity/mortality.

The heparin-mediated extracorporeal low-density lipoprotein precipitation procedure (HELP apheresis) is the main non-traditional technique used for selective removal of plasma low density lipoproteins in the indication of familial hypercholesterolemia unresponsive to medication. This procedure is based on application of high levels of heparin (100 IU/ml) together with the lowering of plasma pH values [36].

HELP apheresis manages to reduce levels of proatherogenic lipoproteins and of several circulating proinflammatory markers as well [36]. Although HCP uses lower levels of heparin and does not require the lowering of the pH values, our results are in agreement with results presented by Wang et al. [37], showing reduction in fibrinogen and CRP levels. It has to be emphasized that
HCP did not reduce fibrinogen levels below normal levels as in the data presented by Wang et al. [37].

Many atherosclerotic patients have normal levels of atherogenetic lipoproteins [38], suggesting that they are not always the major risk factor of atherosclerosis. These patients do not qualify for treatment by HELP apheresis.

In order to effectively attenuate the atherosclerotic process, there is a genuine need for a new therapeutic approach, such as HCP, which can simultaneously reduce multiple risk factors, independent of lipoprotein levels.

This study evaluated the effect of HCP on the in vitro precipitation of modified molecules and circulating non-traditional risk factors for atherosclerosis in plasma removed from HD patients. We believe that the data presented in this study may serve as a basis for future studies focusing on the development of an applicable in vivo clinical purification technique. Additionally, evaluation of the effect of HCP in vitro on plasma removed from other groups of atherosclerotic patients (such as patients with dyslipidemia/hypercholesterolemia) has to be carried out. The HCP method may be developed to complement the pharmacological treatments designed to reduce levels of circulating non-traditional risk factors for atherosclerosis and CVDs, in achieving attenuation in the propagation of atherosclerosis.

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References


