Improvement of 2D-PAGE resolution of human, porcine and canine follicular fluid: comparison of two immunodepletion columns
Somayyeh Fahiminiya, S. Roche, Nadine Gérard

To cite this version:

HAL Id: hal-01129661
https://hal.archives-ouvertes.fr/hal-01129661
Submitted on 18 Dec 2017

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
Short Communication

Improvement of 2D-PAGE Resolution of Human, Porcine and Canine Follicular Fluid: Comparison of Two Immunodepletion Columns

S Fahiminiya1, S Roche2,3 and N Gérard1

1Physiologie de la Reproduction et des Comportements, UMR 6175, INRA, Nouzilly, France; 2Institut de Génétique humaine du CNRS, UPR1142, Montpellier, France; 3Plateforme de Protéomique Clinique, Hôpital Saint-Eloi, Montpellier, France

Contents

Follicular fluid provides the microenvironment within which somatic cells proliferate and differentiate, and the oocyte matures. It contains a number of soluble factors implicated in various stages of follicular development, most of them being functionally unknown. The presence of several high-abundance proteins, mainly originating from the blood circulation, is a major challenge of follicular fluid proteomic analysis, as these proteins can mask or decrease the visualization of follicle-specific proteins. In this study, we evaluated the efficiency of two immunodepletion columns (ProteomeLab™ IgY-HSA and MARS-6) on follicular fluids of human, porcine and canine prior to 2D-PAGE. Our results showed that both columns were suitable to remove some of the high-abundance proteins present in human and canine follicular fluid. In conclusion, we demonstrated that the immunodepletion strategy enables the detection of new protein spots, increases resolution and highly improves the intensity of low-abundance proteins by 2D-PAGE.

Introduction

Follicular fluid (FF) accumulates in the antrum of growing follicles. It provides the microenvironment within which somatic cells proliferate and differentiate, and the oocyte matures. Previous proteomic studies on follicular fluid of various species (Cabrera et al. 1985; Angelucci et al. 2006; Fahiminiya et al. 2010) demonstrated that it contains several high-abundance proteins (albumin, immunoglobulins, alpha-1-antitrypsin, transferrin and haptoglobin), which constitute approximately 85–90% of the total protein mass (Jacobs et al. 2005). The presence of these high-abundance proteins in follicular fluid can preclude the identification, or even the detection, of follicle-specific proteins (low-abundance proteins) which locally produced and secreted by ovarian cells into the follicular fluid (Fortune et al. 2004). Detection and identification of these low-abundance proteins like growth or differentiation factors in follicular fluid would help to better understand ovarian follicular growth and oocyte development in mammals. Therefore, an efficient approach to overcome this challenge might be to remove some high-abundance plasma proteins from follicular fluid prior to its proteomic analysis. Recently, the application of immunodepletion columns has been raised to remove up to 20 high-abundance proteins from human biological fluids. To our knowledge, no pre-fractionation method was developed specially for follicular fluid. In this study, we evaluated the efficiency of two immunodepletion columns (ProteomeLab™ IgY-HSA and MARS-6) to remove high-abundance proteins from human (HFF), porcine (PFF) and canine (CFF) follicular fluids prior to 2D-PAGE analysis. These two columns are based on either immunoglobulin G (IgG) in mammals (Martosel-lä et al. 2005) or immunoglobulin yolk (IgY) in chickens (Zhang 2003; Huang et al. 2005; Linke et al. 2007). Indeed, the immunodepletion of follicular fluid allowed us to visualize several peptides and proteins on 2D gels, previously masked by high-abundance proteins.

Material and Methods

Porcine ovaries were collected at the slaughterhouse, and the follicular fluids were punctured from ovaries with a 20-G needle. Canine follicular fluids were aspirated from ovaries collected by ovariotomy (Fahiminiya et al. 2010). Human follicular fluids were collected according to Guerif et al. (2003) method. In each species, the follicular fluid was collected from three different ovaries. After collection, follicular fluid samples were centrifuged for 10 min at 1500 × g and stored at −80°C till immunodepletion step.

Two immunodepletion spin columns were used to remove high-abundance plasma proteins from the follicular fluids of human, porcine and canine species. As far as, we know these two immunodepletion methods has no side effect on the protein structure. All steps were performed at room temperature as recommended by the manufacturers. The depleted fractions were concentrated by 10-kDa molecular mass cut-off centrifugal concentrators (Microcon™; Millipore, Bedford, MA, USA), and the pH of fractions was neutralized by 50 mM Tris pH 8.8. Names and properties of the two immunodepletion columns are presented in Table 1.

All the chemicals and materials used for proteomic analysis were purchased from Bio-Rad (Marnes-la-Coquette, France), unless otherwise indicated. The protein assay and 2D-PAGE analysis were performed as previously described (Fahiminiya et al. 2010). Briefly, the total protein content of crude and depleted follicular fluids was determined using DC (detergent compatible) assay with bovine serum albumin as standard (Pierce, Rockford, IL, USA) and statistically analysed by the non-parametric Kruskal–Wallis test. For analytical separations, 2D-PAGE was performed on three individual samples in each species: 100 µg of proteins was loaded upon ReadyStrip™ IPG (11-cm IPG strips, pH
Results and Discussion

The major challenge of follicular fluid proteomic analysis is its high similarity with plasma, in which a few proteins (high abundance) constitute more than 80% of the total protein content. One efficient methodology prior to its proteomic analysis could be the depletion of high-abundance proteins, to visualize, and consequently identify some lower-abundance ones. The latter may be most probably produced locally by follicle cells. To our knowledge, no method has been developed to deplete follicular fluid before its proteomic analysis. Here, we compared the efficiency of two immunodepletion columns in three species, with the aim to determine whether the removal of high-abundant proteins could improve the resolution of 2D protein patterns of follicular fluid.

Depletion efficiency and reproducibility of ProteomeLab™ IgY-HSA and MARS-6 on HFF, PFF and CFF were determined after the measurement of the total protein content (Table 2) and 2D-PAGE analysis.

Table 2. Total protein concentration and the percentage of depletion efficiency in crude and depleted FF samples

<table>
<thead>
<tr>
<th>Total protein concentration (mg/ml; mean ± SEM)</th>
<th>ProteomeLab™ IgY-HSA</th>
<th>Multiple affinity removal spin cartridge – human 6 (MARS-6)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HFF</td>
<td>81.85 ± 16.45</td>
<td>87.27 ± 2.56</td>
</tr>
<tr>
<td>CFF</td>
<td>85.00 ± 9.00</td>
<td>83.13 ± 1.74</td>
</tr>
<tr>
<td>PFF</td>
<td>91.40 ± 2.10</td>
<td>84.98 ± 1.02</td>
</tr>
</tbody>
</table>

CFF, canine follicular fluid; HFF, human follicular fluid; PFF, porcine follicular fluid; IgY, immunoglobulin yolk.

All values are based on three independent experiments.

Differences are not significant.

The depletion efficiency was calculated with following formula: \( \frac{(\text{Protein amount (pg)} \times 100)}{\text{protein amount (pg) in crude FF)}} \)
were completely depleted in all three species. This result was in accordance with the depletion efficiencies (HFF and CFF > PFF). Figure 1C clearly showed that HFF depleted with MARS-6 exhibit many more protein (+254) spots in entire pH interval as compared with crude samples. Again, this can be explained by the specificity of this immunodepletion column against human proteins. Concerning CFF, some proteins showed a higher intensity after depletion by MARS-6, compared with crude follicular fluid, but no new proteins were revealed. Finally, the effective removal of immunoglobulins from PFF did not exhibit any new proteins after depletion by MARS-6.

Taken together, our data suggest that immunodepletion of the six most highly abundant plasma proteins using the MARS-6 column is more effective at reducing the dynamic range of follicular fluid proteins than the ProtocomeLab™ IgY-HSA column, and results in the visualization of more medium- and low-abundance follicular fluid proteins by 2D-PAGE.

Conclusion
This study provides the first comprehensive investigation of commercially available immunodepletion columns, to improve the protein profiling of follicular fluid derived from human, porcine and canine. In this study, we showed that both immunodepletion columns were more effective to remove high-abundance proteins especially albumin from HFF and CFF, compared with PFF. In addition, new proteins were revealed in HFF 2D-PAGE after depletion by MARS-6. Based on our results, we conclude that both columns are efficient and reproducible methods that can be used in combination.
with 2D-PAGE and/or mass spectrometry before proteomic analysis of human and CFF.

Acknowledgements
The authors thank Dr Dominique Royère, Dr Joëlle Dupont and Dr Karine Reynaud for providing human and canine follicular fluids, respectively. Pascal Papillier is acknowledged for collecting porcine ovaries from slaughterhouse. SF and NG thank Dr Sylvain Lehmann for accepting them in his laboratory during the test of immunodepletion columns. This research programme was financially supported by INRA, France. Somayyeh Fahiminiya was PhD student supported by a fellowship from INRA.

Conflict of interests
None of the authors have any conflict of interest to declare.

Author contributions
SF carried out immunodepletion, 2D-PAGE analysis of HFF, PFF and CFF, computerized image analysis and drafted the manuscript. SR helped for immunodepletion of follicular fluid samples. NG designed and supervised the study, participated to the immunodepletion of follicular fluid samples and revised the manuscript. All authors read and approved the final manuscript.

References
Submitted: 3 Jun 2011; Accepted: 13 Oct 2011
Author’s address (for correspondence): N Gérard, UMR 6175, INRA, 37380 Nougazilly, France. E-mail: nadine.gerard@tours.inra.fr