Modification of the Secretion Pattern of Proteases, Inflammatory Mediators, and Extracellular Matrix Proteins by Human Aortic Valve is Key in Severe Aortic Stenosis*

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One of the major challenges in cardiovascular medicine is to identify candidate biomarker proteins. Secretome analysis is particularly relevant in this search as it focuses on a subset of proteins released by a cell or tissue under certain conditions. The sample can be considered as a plasma subproteome and it provides a more direct approximation to the in vivo situation. Degenerative aortic stenosis is the most common worldwide cause of valve replacement. Using a proteomic analysis of the secretome from aortic stenosis valves we could identify candidate markers related to this pathology, which may facilitate early diagnosis and treatment. For this purpose, we have designed a method to validate the origin of secreted proteins, demonstrating their synthesis and release by the tissue and ruling out blood origin. The nLC-MS/MS analysis showed the labeling of 61 proteins, 82% of which incorporated the label in only one group. Western blot and selective reaction monitoring differential analysis, revealed a notable role of the extracellular matrix. Variation in particular proteins such as PEDF, cystatin and clusterin emphasizes the link between aortic stenosis and atherosclerosis. In particular, certain protein levels correlates well, not only with label incorporation trend (only labeled in aortic stenosis group) but, more importantly, with alterations found in plasma from an independent cohort of samples, pointing to specific candidate markers to follow up in diagnosis, prognosis, and therapeutic intervention.

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Degenerative aortic stenosis (AS) is currently the most common cause of valve replacement in Western countries, and a significant increase in its prevalence is expected in the future because of increasing longevity (1). At the same time that our population age increases it can be expected that cardiac valve disease, and AS in particular, will increase in parallel. More and more, clinicians are seeing patients with symptomatic severe AS who are very advanced in age and have severe comorbidities or significant frailty, making operative intervention either impossible or of very high risk in the eyes of the cardiac surgeon. For this reason, the need of new diagnostic and prognostic methods, together with the urgent need of new drugs for therapy, has increased making a correct diagnosis in the early stages of the disease thereby reducing the cost burden to society. Several lines of evidence have demonstrated that degenerative AS is an active process in which inflammation plays a key role. As such, preventative approaches similar to those used in coronary artery disease (CAD) may be also applicable to AS (2, 3, 4, 5, 6). However, recent studies reported no reduction in later states of AS disease using statins, which significantly benefit patients with atherosclerosis (7, 8). Hence, further research is required to elucidate the pathogenic mechanism of this prevalent disease and to identify the similarities and differences in relation to atherosclerosis.

Proteomics has emerged as a particularly suitable platform for the unbiased analysis of proteins involved in the pathogenesis of various diseases, such as AS (9, 10). This type of approach provided the basis for the development of biomarkers to detect patients at risk of developing degenerative AS. Plasma and serum have been the main source used in proteomic studies to identify candidate protein biomarkers, fol-

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1 The abbreviations used are: AS, Aortic Stenosis; ACN, Acetonitrile; FA, Formic acid; HRP, Horseradish peroxidase; HTN, Arterial hypertension; nLC-MS/MS, liquid chromatography mass spectrometry; PBS, Phosphate Buffer Saline; TFA, Trifluoroacetic acid; WB, Western blot.
lowed by tissue and cell samples. However, the use of plasma and serum is hampered by their complex nature and by the large dynamic range of protein concentrations, which may favor the detection of abundant proteins at the expense of those present at lower concentrations (11, 12). As such, the aortic valve secretome has emerged as an attractive target to further understand the AS pathogenic process. Tissue secretomes provide a more accurate model of the in vivo situation and, by minimizing serum contaminants, they facilitate the detection of low abundance proteins secreted into the blood (13).

In the present study, aortic valves from AS patients and nonaffected control subjects were cultured and the secretome analyzed by nLC-MS/MS. The addition of labeled amino acids to the culture media enabled the origin of the secreted proteins to be validated (truly secreted versus serum contaminants) and provided with information related to the dynamics of their synthesis and release from tissue.

EXPERIMENTAL PROCEDURES

Patient and Control Subjects Selection—Heart valves (n = 20) were obtained from AS patients of both sexes (55% male, 45% female), with an average age of 74 (±4) years, presenting at the hospital with one or more of the cardinal AS symptoms (chest pain, dyspnea, syncope) and had undergone aortic valve replacement because of an echocardiography diagnosis of severe degenerative AS. All patients (100%) had hypertension, 50% suffered from hyperlipidemia and 60% from diabetes mellitus. Moreover, 60% of patients suffered coronary artery disease and surgery was indicated according to current practice guidelines. Neither antecedent of rheumatic fever, nor valvular morphologic changes consistent with rheumatic heart disease (e.g. typical mitral valve changes) were present in any patient. Patients with aortic regurgitation were also excluded in the study (Table I A). Control macroscopically normal valves (n = 20, Table I B) were obtained within 4 h from death from autopsies of subjects who had died from causes other than cardiovascular disease and who had no history of coronary artery disease or diabetes mellitus. Before resecting the valve, a cardiac surgeon explored it carefully. Valves with calcifications or cusp restriction were excluded as control cases. If inserted saline in aortic root remained in the closed valve, we ruled out significant regurgitation and we took these aortic valves as a control case. In all the cases, both stenotic and nonstenotic valves, the structure was 3-cuspid.

For a pilot selective reaction monitoring (SRM) validation analysis, peripheral blood samples (n = 14) were collected from AS patients and controls with comparable conditions; both cohorts had an average age of 74 years, were from both sexes (40% male, 60% female) and with analogous cardiovascular risks factors (100% hypertension, 60% diabetes, and 50% hyperlipidemia). AS plasma samples (n = 7) were obtained from patients who underwent aortic valve replacement because of severe degenerative AS. Plasma control samples (n = 7) were obtained from subjects with no apparent cardiovascular illness, and no history of coronary arterial disease. A greater cohort of 80 plasma samples (43 patients and 37 controls) was collected for a subsequent validation analysis using a nonbiased control population. This study was carried out in accordance with the recommendations of the Helsinki Declaration and it was approved by the ethics committee at the Hospital “Virgen de la Salud” (Toledo, Spain). Signed informed consent was obtained from all subjects or relatives in case of necropsies before their inclusion in the study. Clinical information is detailed in Supplementary Material.

Aortic Valve Tissue Culture—The aortic tissue was cultured as described previously (14), with minor modifications. Aortic valves with or without stenosis were transported from the operating room to the laboratory in PBS buffer at room temperature. The tissue was thoroughly washed three times with PBS supplemented with antibiotics and fungizone and washed twice more with 1640 Roswell Park Memorial Institute (RPMI) medium to eliminate serum contaminants. Samples were then transferred to a petri dish (Cell Star®), cut into pieces of about 1 mm³ and incubated separately at 37 °C in l-lysine-arginine free 1640 RPMI medium (Cell Culture Technologies Invitrus) supplemented with 5 mg/ml fungizone, 250 mg/ml amikazin, 2 mg/ml L-lysine 2HCl (U-13C6, 97–99%), and 10 mg/ml L-Arginine HCl (U-13C6, 97–98%: Cambridge Isotope Laboratories Inc., Andover, MA) in a humidified atmosphere of 5% CO2. The culture medium was changed at 4 h and 8 h to eliminate serum contaminants and the tissue was incubated and collected for subsequent analysis at different time points (24 h, 48 h, 72 h, 96 h, and 120 h) to determine the point of maximum protein secretion. Collected media were stored at −80 °C until analysis (Fig. 1).

nLC-MS/MS Identification—Proteins released into the culture media by valve tissue (control or AS) were concentrated by ultrafiltration (Centriplus, 3-kDa cutoff, Millipore) and the protein concentration was determined by the Bradford method (Bradford assay, Bio-Rad, Hercules, CA). Subsequently, 25 μg of total sample protein were loaded onto SDS-PAGE and concentrated in a unique band that concentrated all the proteins in the sample, stained with Coomassie Brilliant Blue G-250 (PageBlue staining solution, Fermentas, Hanover, MD), excised in 15–20 small pieces and stored at 4 °C in ultrapure water for later analysis. Samples were subjected to reduction and alkylation before the in-gel digestion. Samples were in-solution digested by adding modified porcine trypsin (Promega) at a final ratio of 1:50 (trypsin-protein). Digestion proceeded overnight at 37 °C. After digestion, samples were vacuum-dried and finally dissolved in 1% acetic acid for LC-MS analysis. The resulting tryptic peptide mixtures were subjected to nano-liquid chromatography coupled to mass spectrometry for protein identification. Peptides were injected onto a C-18 reversed phase (RP) nano-column (100 μm I.D. and 12 cm, Merck-Millipore) and the protein concentration was determined by the Bradford method (Bradford assay, Bio-Rad, Hercules, CA). An enhanced FT-resolution spectrum (resolution = 60000) followed by the MS/MS spectra from most intense five parent ions were analyzed using a LTQ-Orbitrap mass spectrometer (Thermo Fisher, San José, CA). A Mascot score was set up to search human.fasta (1.0, 199762 entries). Mascot was set up to search a subset of SwissProt (51.6, 236102 entries). Both search engines were run assuming full trypsin digestion. Two missed cleavages were allowed and an error of 15 ppm or 0.8 Da was set for full MS or MS/MS spectra searches, respectively. All identifications were performed by Decoy database search for FDR analysis set at 0.05 by applying corresponding filters (Mascot Significance Threshold, and Score versus Charge state). Oxidation in M, phosphorylation in S, T, or Y, and SILAC+6 in K or R were selected as dynamic modifications.

Antibodies—The primary antibodies used to probe Western blots included: a mouse polyclonal antiserum against tenascin XB; monoclonal antibodies against clusterin, pentraxin 3 and gelsoxin, all from

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Abcam; and a monoclonal pigment epithelium-derived factor (PEDF) antibody (Lifespan Biosciences).

**Western Blotting**—Protein samples obtained from control and AS valve secretomes were resolved on 12% SDS-PAGE gels using a Bio-Rad Miniprotean II electrophoresis unit run at a constant current of 25 mA/gel for 1 h. Proteins were then transferred to a nitrocellulose membrane under constant voltage (15 V for 20 min) and the membranes were stained with Ponceau S to ensure equal protein loading of the secretome samples. The membranes were then blocked with PBS-T containing 7.5% nonfat dry milk for 1 h (15), incubated overnight with the primary antibody in PBS-T containing 5% nonfat dry milk and subsequently incubated with the specific HRP-conjugated secondary antibody in the same solution.

**SRM**—Protein samples were reduced with 100 mM dithiothreitol (Sigma Aldrich) in 50 mM ammonium bicarbonate (99% purity; Scharlau) for 30 min at 37 °C, and alkylated for 20 min at room temperature

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**TABLE IA**

A, Clinical characteristics of the patients with AS. Experiments carried out with every sample are indicated: secretome, western blot (WB), or selected reaction monitoring (SRM). (HTN) arterial hypertension

| Patient num. | Age/Sex | HTN | Diabetes | Dyslipidemia | Discovery phase | WB | SRM |
|--------------|---------|-----|----------|--------------|----------------|----|-----|
| 1            | 74/F    | Yes | No       | Yes          | X              |     |     |
| 2            | 73/M    | Yes | Yes      | No           |                |     | X   |
| 3            | 68/M    | Yes | No       | Yes          |                |     | X   |
| 4            | 81/M    | Yes | No       | Yes          |                |     | X   |
| 5            | 69/M    | Yes | Yes      | Yes          |                |     | X   |
| 6            | 79/F    | Yes | Yes      | No           |                |     | X   |
| 7            | 73/F    | Yes | Yes      | Yes          |                |     | X   |
| 8            | 77/M    | Yes | No       | No           |                |     | X   |
| 9            | 79/F    | Yes | Yes      | No           |                |     | X   |
| 10           | 75/M    | Yes | No       | Yes          |                |     | X   |
| 11           | 74/M    | Yes | Yes      | Yes          |                |     | X   |
| 12           | 74/F    | Yes | Yes      | No           |                |     | X   |
| 13           | 79/F    | Yes | Yes      | No           |                |     | X   |
| 14           | 79/F    | Yes | Yes      | No           |                |     | X   |
| 15           | 72/F    | Yes | No       | No           |                |     | X   |
| 16           | 63/M    | yes | no       | Yes          |                |     | X   |
| 17           | 72/M    | Yes | No       | Yes          |                |     | X   |
| 18           | 75/F    | Yes | Yes      | Yes          |                |     | X   |
| 19           | 78/M    | Yes | Yes      | No           |                |     | X   |
| 20           | 74/M    | Yes | Yes      | No           |                |     | X   |

Mean 74 ± 4/45%F-55%M 100,00% 60,00% 50,00%

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**TABLE IB**

B, Clinical characteristics of control subjects. (COPD) Chronic obstructive pulmonary disease, (WB) Western blot, (SRM) selected reaction monitoring

| Control num. | Age/Sex | Cause of the death | HTN | Discovery phase | WB | SRM |
|--------------|---------|---------------------|-----|----------------|----|-----|
| 1            | 64/M    | Bilateral acute bronchopneumonia | No | X              |     |     |
| 2            | 51/F    | Septic shock        | Yes |                |     |     |
| 3            | 47/H    | Sepsis              | No  | X              |     |     |
| 4            | 47/F    | Bacterial pneumonia | No  |                |     |     |
| 5            | 47/F    | Carcinogenesis      | No  |                |     |     |
| 6            | 34/M    | Acute Pancreatitis  | Yes | X              |     |     |
| 7            | 64/F    | Bacterial Sepsis    | No  |                |     |     |
| 8            | 78/F    | Aspiration bronchopneumonia | No |                |     | X   |
| 9            | 42/F    | Lymphoma            | No  |                |     | X   |
| 10           | 67/M    | Renal insufficiency | No  |                |     | X   |
| 11           | 74/M    | COPD                | No  |                | X   |     |
| 12           | 87/F    | Carcinoma           | No  |                |     | X   |
| 13           | 48/F    | Pulmonary thromboembolism | No |                |     | X   |
| 14           | 55/M    | Atypical pneumonia  | No  |                |     | X   |
| 15           | 87/F    | Cholelithiasis      | No  |                |     | X   |
| 16           | 76/M    | Carcinoma           | No  |                |     | X   |
| 17           | 80/F    | Atrial fibrillation | Yes |                |     | X   |
| 18           | 49/F    | Respiratory insufficiency | No |                |     | X   |
| 19           | 74/M    | Septic shock        | Yes |                |     | X   |
| 20           | 77/F    | COPD                | Yes |                |     | X   |

Mean 69 ± 7/80%F-40%M 25,00%
Peptides were loaded onto a LC/MS/MS, Applied Biosystems). Three replicate injections (2 µg of protein in 4 µL) were performed per sample in the mobile phase A (2% ACN/98% water, 0.1% FA) at a flow rate of 10 µL/min for 5 min. Peptides were loaded onto a µ-Precolumn Cartridge (Acclaim Pep Map 100 C18, 5 µm, 100 Å; 300 µm i.d. X 5 mm, LC Packings) to preconcentrate and desalt samples. Reversed-phase liquid chromatography (RPLC) was performed on a C18 column (Onyx Monolithic C18, 150 × 0.1 mm i.D., Phenomenex) in a gradient of phase A and phase B (98% ACN/2% water, 0.1% FA). The peptides were eluted at a flow rate of 300 nL/min in a continuous acetonitrile gradient: 2–15% B for 2 min, 15–50% B for 38 min, 50 to 90% B for 2 min and 90% B for 3 min. Both the TEMPO nano LC and 4000QTRAP system were controlled by Analyst Software v.1.4.2. The mass spectrometer was set to operate in positive ion mode with ion spray voltage of 2800V and a nanoflow interface heater temperature of 150 °C. The source gas and curtain gas were set to 20 and 10 psi respectively, and nitrogen was applied as both curtain and collision gases.

Theoretical SRM transitions were designed using MRMpilot software v1.1 (ABSciex), with the following settings: Enzyme = trypsin, missed cleavages = 0; modifications in peptide = 3; charge states = +1 from 300 to 600 Da, +2 from 500 to 2000 Da, +3 from 900 to 3000 Da; studied modifications = carboxyamidomethylation; variable modifications = none; min. number of amino acids ≥ 5; max. number of amino acids = 30; ignore multiple modification sites; 3 transitions per peptide (Table II). A pool containing a mixture of all the samples was digested as described previously and analyzed in the 4000QTrap using a MIDAS acquisition method that included the theoretical transitions. Transitions were selected when the three co-eluting peaks (corresponding to the three transitions of the same peptide) had a signal-to-noise ratio over 5 and the MS/MS data matched the theoretical spectrum for that peptide. Collision energy was optimized to obtain the maximum transmission efficiency and sensitivity for each SRM transition. A total of 24 transitions (3 per peptide) were monitored during an individual sample analysis. They were acquired at unit resolution in both Q1 and Q3, with dwell times of 50 ms resulting in cycle times of 1.2 s. The IntelliQuan algorithm included in Analyst 1.4.2 software was used to calculate the peptide abundance on the basis of peak areas after integration.

Pathway Analysis—Proteomics data were uploaded into the Ingenuity Pathways Analysis (IPA) (Ingenuity® Systems, www.ingenuity.com). Providing a list of GI accession numbers, biological networks were generated based on interactions between these selected genes and those stored in the Knowledge Base. Graphically, nodes correspond to genes and edges correspond to the biological relationships between them. IPA analysis computes a score for each network according to the fit of the user’s set of significant genes. The significance value associated with functional analysis for a data set is a measurement of the likelihood that the association between a set of functional analysis molecules in the experiment and a given process or pathway is because of random chance.
Statistical Analysis—Western blot bands were measured using a GS-800 Calibrated Densitometer (Bio-Rad) and the values obtained analyzed using the statistical software packages SPSS 13.0 and GraphPAD Instat (GraphPAD software). A Kolmogorov-Smirnov test was applied to evaluate normal distribution of the population analyzed. When normal distribution was demonstrated, a Levene test for homogeneity of variance was performed and the Student t test was used to compare band intensities and SRM peak areas, selecting adequate p value in view of homogeneity results. For populations with no normal distribution, the nonparametric Mann-Whitney U test was used. Right tailed Fischer Exact Test was applied for IPA analysis. For all tests, statistical significance was accepted when p ≤ 0.05. All LC-MS/MS identifications were performed by Decoy database search for FDR analysis set at 0.05 by applying corresponding filters (Mascot Significance Threshold, and Score versus Charge state).

RESULTS

Secretome Characterization: Metabolic Labeling of Secreted Proteins—The secretome from human control and AS valves from three individuals per group (3 patients, 3 controls) was obtained and the labeled proteins detected were investigated (Fig. 1). The maximum number of labeled proteins from both pathological and healthy valves was obtained at 96 h (Fig. 2A), when 248 proteins were unambiguously identified in total, labeled or not. Among these, 61 proteins were labeled in at least one of the three secretome replicates from either of the two groups (control or patient), 50 in AS and 23 in control aortic valves (Fig. 2B). Twelve proteins of the 61 were secreted by both AS and control valves. All the labeled proteins were analyzed using the SecretomeP software (16) to predict them as being secreted: the presence of a signal peptide indicated secretion by the classical pathway and a NN-score of over 0.5, in the absence of a signal peptide, indicated nonclassical secreted proteins. Of the 61 proteins, 36 contained a signal peptide and 9 proteins were found to be nonclassically secreted. Among the 17 proteins not predicted to be secreted by Secretome P, 5 had been previously detected in serum/plasma (according to the Human Protein Reference Database: www.hprd.org).

The labeled proteins were classified into six different groups based on their molecular function (UniprotKB database): extracellular matrix constituents (ECM)/cytoskeletal, immune response, degradation, transport/metabolic processes, regulatory function, and other (Fig. 2B). Pathway analysis on proteomics data from Table III shows “Cellular Movement, Connective Tissue Disorders, Cardiovascular System Development and Function” as the main associated functional network (see Fig. 3). “Cardiovascular System Development and Function” and “Tissue Development” were the most significant functions related to Physiological System Development, with 18 and 23 proteins from the data set involved, respectively. Additional data can be found in supplemental material and in ftp://PASS00237:BE5743t@ftp.peptideatlas.org/.

Secretion Pattern in Aortic Stenosis—Of the 61 labeled proteins, 50 (82%) incorporated the label in only one of the two groups (control = 11 and AS = 38: Table III). These proteins deserve particular attention as they represent poten-
tial diagnostic and therapeutic targets, given that they reflect the main differences observed between affected and healthy tissue at the protein level. We analyzed additional secretome samples (n = 11005 per group for each analysis by Western blot and n = 10 per group by SRM) to those used in the discovery phase, from both control and AS groups.

**Western Blot Analyses**—A group of five proteins was analyzed in Western blot analysis (Fig. 4), demonstrating that clusterin (Fig. 4A) and tenascin XB (Fig. 4B) were down-regulated in AS valve secretome samples when compared with the control samples (p = 0.0224 and p = 0.0002, respectively). By contrast, the expression of gelsolin (Fig. 4C), pentraxin 3 (PTX3, Fig. 4D) and pigment epithelium-derived factor (Fig. 4E) was up-regulated in AS secretome valves (p = 0.0112, p < 0.0001, p = 0.0002, respectively).

**SRM Analyses**—Secretomes obtained from an independent group of aortic valve samples (n = 10 controls, n = 10 AS patients) were analyzed by SRM. In particular, four proteins that incorporated the label in only one of the two groups were measured. In all cases, the two peptides measured for each protein were found to differ significantly their expression (Fig. 5). This was observed for both angiotensinogen peptides (Fig. 5A, p < 0.0001), for EGF-containing fibulin-like extracellular matrix protein 1 (Fig. 5B, p = 0.0362 peptide 1 and p = 0.0349 peptide 2) and for gelsolin (Fig. 5C, p = 0.0197 peptide 1 and p < 0.0001 peptide 2). The expression of all of these proteins was up-regulated in AS versus control secretomes. In contrast, prostaglandin-D2 synthase (Fig. 5D, p = 0.0084 peptide 1 and p < 0.0001 peptide 2) was down-regulated in AS versus control secretomes (Fig. 5).

**Reflection in Plasma**—In an independent cohort of plasma samples (n = 7 controls, and n = 7 AS patients) matched by sex, age, and AS risk factors (hypertension, diabetes, and dyslipidemia), three proteins were analyzed by SRM: clusterin (p = 0.0053), gelsolin (p = 0.0253), and prostaglandin D2 synthase (p = 0.0236) (see Supplementary Figures). In this first pilot study, differential trends observed by WB or SRM in plasma corroborated previous findings in secretome samples and allowed excluding sex, age, and AS risk factor bias. More importantly, results suggest that some proteins found differentially expressed in the secretome analyses could be validated in patients’ plasma. ROC curves calculated for the three proteins showed the best results for gelsolin (area under the curve, AUC = 0.950) (see Supplementary Figures). Among all proteins tested in secretome samples, gelsolin showed the same trend (increased in AS group) independently of the strategy used for validation and in all the approaches tested (WB and SRM). Interestingly, this protein followed the opposite trend in the secretome from atherosclerotic coronary arteries (13). For all these reasons and considering the limited knowledge about common and specific operating mechanisms in AS and atherosclerosis pathologies, we further validate gelsolin response in a new, wider and independent cohort of samples. To prove the potential of this protein as marker of AS, gelsolin was comparatively analyzed by SRM in a cohort of 80 plasmas (43 patients and 37 controls), finding again that plasma levels increased in patients’ cohort and corroborating previous findings (Fig. 6A). ROC curve (AUC = 0.950) is shown in Fig. 6B. The two peptides measured with three transitions per peptide gave significant variation (p val-
### Table III
Proteins labelled in at least one replicate from either of the two groups. P//C: Number of AS patients (P) or controls (C). Sec, Secretion pathway of the proteins found in plasma, according to the secretome P software and the Human Protein reference database. NS: no secreted, CP: secreted by classical pathway, NCP: secreted by nonclassical pathway, BL: detected in serum or plasma.

| P//C | Sec. | Accession | Gen name | Protein name | Function |
|-----|------|-----------|----------|--------------|----------|
| 3/2 | NS, BL | GI:53791223 | FINC | Fibronectin 1 | Extracellular matrix structural constituent |
| 3/0 | CP | GI:220125 | TIMP | Tissue inhibitor of metalloproteinases | Extracellular matrix structural constituent |
| 2/1 | CP | GI:4502403 | BGN | Biglycan preproprotein | Extracellular matrix structural constituent |
| 3/1 | NS | GI:30851190 | CO6A1 | Collagen, type VI, alpha 1 precursor | Extracellular matrix structural constituent/cell adhesion |
| 3/0 | NS | GI:179716 | CO7 | Complement C7 | Extracellular matrix structural constituent/cell adhesion |
| 1/0 | NS | GI:2065167 | COL14A1 | Collagen, type XIV | Extracellular matrix structural constituent/cell adhesion |
| 1/0 | NS | GI:296439507 | CO1A2 | Prepro-alpha2(i) collagen | Extracellular matrix structural constituent/cell adhesion |
| 3/3 | CP | GI:11342666 | MMP2 | Matrix metalloproteinase 2 (Gelatinase A) | Endopeptidase activity |
| 2/0 | CP | GI:119226208 | TGFDS | Prostaglandin D2 synthase 21kDa | Transporter activity |
| 3/0 | CP | GI:144226251 | CHI3L1 | Chitinase-3-like protein 1 | Chitinase activity |
| 3/0 | CP | GI:40317843 | COMP | Cartilage oligomeric matrix protein | Extracellular matrix structural constituent |
| 1/0 | NS | GI:71773110 | APOB | Apolipoprotein B-100 | Transporter activity |
| 3/1 | NS | GI:311254603 | CHI3L2 | Chitinase 3-like 2 isoform a | Chitinase activity |
| 1/0 | NS | GI:4502693 | CD9 | CD9 antigen (27 kDa diphtheria toxin receptor-associated) | Immune response: platelet activation and aggregation |
| 1/2 | CP | GI:1955047 | LUM | Lumican precursor | Extracellular matrix structural constituent |
| 2/2 | CP | GI:75075054 | CERU | Ceruloplasmin precursor | Chaperone/Ferroxidase enzyme |
| 1/0 | CP | GI:4506141 | HTRA1 | Serine protease HtrA | Serine-type endopeptidase activity |
| 2/0 | CP | GI:11118375 | ANGT | Angiotensinogen (serpin peptidase inhibitor) | Acetyltransferase stimulator activity |
| 1/0 | CP | GI:189580 | PAI1 | Plasminogen activator inhibitor type 1 | Coagulation |
| 1/0 | CP | GI:119592657 | LENG9 | Leukocyte receptor cluster (LRC) member 9 | Receptor activity/Immune response |
| 1/0 | NCP | GI:19923830 | PBP1 | Pre-B-cell leukemia transcription factor interacting protein 1 | transcription corepressor activity |
| 1/0 | NCP | GI:207101417 | CAR82030 | Unnamed protein product Fab | Unknown |
| 1/0 | NCP | GI:7438711 | JEO242 | Ig kappa chain NiG26 precursor | Immune response |
| 2/0 | NCP | GI:229601 | IGH3 | Ig G1 H | Immune response |
| 1/0 | NCP | GI:229585 | IGHA1 | Ig A1 | Immune response |
| 1/0 | NS | GI:189778 | TSP1 | Thrombospondin 1 precursor | Potent inhibitor of angiogenesis |
| 2/0 | CP | GI:181192 | CATB | Preprocathepsin B | Cysteine protease |
| 1/0 | CP | GI:4504165 | GELS | Gelsolin isoform a | Actin binding |
| 1/0 | CP | GI:50659080 | AACT | Alpha-1-antichymotrypsin | Immune response |
| 2/0 | CP | GI:73858750 | IC1 | Plasma protease (C1) inhibitor | Protease inhibitor |
| 1/0 | CP | GI:5901956 | FSTL3 | Follistatin-related protein 1 precursor | BMP signaling pathway |
| 1/0 | NS, BL | GI:119631316 | CO3A1 | Collagen, type III | Extracellular matrix structural constituent |
| 1/0 | NCP | GI:53759113 | B4GT2 | Beta1,4-galactosyltransferase-2 | Metabolic process: N-acetylglactosamine synthase activity |
| 2/2 | NCP | GI:23396609 | IBP7 | Insulin-like growth factor binding protein 7 | Signaling/regulatory function |
| 1/0 | NS | GI:51477704 | SMRC1 | SWI/SNF-related matrix-associated actin-dependent regulator | Transcription regulator |
| 1/0 | NS | GI:11320820 | SPON1 | VSGP/F-spondin | Extracellular matrix structural constituent |
| 1/0 | CP | GI:1709663 | PLTP | Phospholipid transfer protein | Lipid transport, lipid binding |
| 1/0 | CP | GI:15072400 | FBLN3 | EGF-containing fibulin-like extracellular matrix protein 1 | Extracellular matrix structural constituent |
| 1/1 | CP | GI:46981969 | PI3G5 | Proliferation-inducing protein 35 | Negative regulation of angiogenesis |
| 1/0 | CP | GI:77176698 | PTX3 | Pentraxin-related protein PTX3 | Inflammatory response |
| 1/1 | CP | GI:10834984 | IL6 | Interleukin 6 (interferon, beta 2) | Immune defense |
| 1/0 | NCP | GI:119569621 | PCC02 | Procollagen C-endopeptidase enhancer | Extracellular matrix structural constituent |
| 1/0 | CP | GI:4261568 | CATD | Cathepsin D | Involved in degradation: proteolysis |
| 1/1 | CP | GI:41393602 | C1S | Complement subcomponent C1s | Immune response |
| 1/0 | CP | GI:4503107 | CYTC | Cystatin C precursor | Involved in degradation: cysteine-type endopeptidase inhibitor activity |
| 3/0 | NS, BL | GI:62088774 | COL1A1 | Collagen alpha-1 chain precursor | Extracellular matrix structural constituent |
secreted by aortic valve tissue, because incorporation of the label by a protein does not necessarily imply “intentional” secretion. However, once synthesized by the tissue, the release of nonsecreted protein into the media could be attributed to damage-induced tissue leakage, which may be a direct consequence of the pathology under study. This approach is here applied for the first time to human valvular tissue. Hence, valvular tissue showed a dynamic nature as proven by the label incorporation in a significant number of proteins. Based on our previous experiments with other tissues, labeling efficiency depends on the analyzed tissue, in some cases being insignificant (data not shown). At the optimum time point (96 h), 248 unique proteins were detected in the secretome of control and AS aortic valves, with 23 and 50 proteins incorporating the metabolic label in control and AS valves, respectively, most of which (82%) were only labeled in one of the two groups. In this sense, differences in labeling incorporation somehow reflect tissue activity and, thus, these proteins may better discriminate between pathological and healthy conditions, having a significant role in AS development. The functional classification of the selected secreted proteins indicated that a significant number were ECM proteins. This protein type serves as a structural support, maintaining the cell architecture, and these proteins play a pivotal role in both morphogenesis and tissue regeneration through ECM remodeling (17). ECM proteins include collagens, non-collagenous glycoproteins, and proteoglycans. Organization of these components into a fibrillar framework provides the mechanical strength and elasticity required for cell adhesion and migration. The alterations in ECM composition observed may therefore indicate active tissue remodeling with a loss of elasticity in the aortic valve. From this group, differences in labeling incorporation and secretion observed by LC-MS/MS for tenascin XB, EGF-containing fibulin-1, gelsolin, and pigment epithelium derived factor proteins correlated well with protein

| P/C | Sec. | Accession | Gen name | Protein name | Function |
|-----|------|-----------|----------|--------------|---------|
| 1/0 | CP   | GI:15079241 | IBP5     | Insulin-like growth factor binding protein 5 protease | Signaling/regulatory function |
| 1/0 | CP   | GI:29788996 | PON3     | Beta-alanine/gamma-aminobutyrate-H+ symporter | Involved in degradation: arylesterase activity |
| 2/3 | CP   | GI:27373753 | CLUS     | Clustatin precursor | Apoptosis |
| 0/1 | NS, BL | GI:120537932 | TENX    | Tenasin XB | Extracellular matrix structural constituent/Cell adhesion |
| 0/1 | CP   | GI:4504991 | NFkB2    | Leukemia inhibitory factor precursor | Transcription regulator |
| 0/1 | CP   | GI:134218 | SAP      | Saposin precursor | Lipid transport |
| 0/2 | CP   | GI:11528678 | CO3      | Complement C3 precursor | Immune response |
| 0/1 | NCP  | GI:190359980 | PPR3E   | Protein phosphatase 1 regulatory subunit | Glycogen metabolic process |
| 0/1 | NS, BL | GI:6648082 | PYGL    | Glycogen phosphorylase | Energy pathways, phosphorylase activity |
| 0/1 | CP   | GI:47678717 | TIMP3    | Matrix metalloproteinase 3, 3preproprotein | Involved in degradation: metalloendopeptidase inhibitor activity |
| 0/1 | CP   | GI:7949430 | FREM2    | Fras1 related extracellular matrix protein 2 | Extracellular matrix protein |
| 0/2 | CP   | GI:1209010 | TIMP1    | Metalloproteinase tissue inhibitor 1 | Involved in degradation: metalloendopeptidase inhibitor activity |
| 0/2 | CP   | GI:66932947 | A2MG    | Alpha-2-macroglobulin precursor | Proteinase activity |
| 0/1 | NS   | GI:83321247 | KIR3DL3 | KIR antigen | Receptor activity |
abundance validation analyses. Additionally, angiotensinogen, clusterin, and pentraxin 3 trends also correlate. The decrease in tenascin XB in patients could be attributed to a functional similarity with tenascin C, with which it shares a common FNIII domain, several repeats of which are found in tenascin XB. Tenascin C is expressed in the cardiac vasculature, where it is implicated in local vascular remodelling (6). If its secretion pattern is modified in AS compared with a healthy status, the interactions between tenascin-C and both endothelial cells and endothelial precursor cells (EPCs) will also be altered. In this context, and considering the similarity between tenascin C and tenascin XB, this protein and its interactions may represent a novel target to follow disease and, in particular, to act on in order to favor the enhancement of vascular repair mechanisms, known to be down-regulated with age and in pathological conditions (18). Importantly, decreased tenascin XB levels are associated with various problems of mitral and tricuspid valves (16). Furthermore, the decrease in tenascin XB seen in Ehler Danlos syndrome increase the probability of aortic aneurisms (19).

We detected an increase in EGF-containing fibulin-like protein secretion in AS valves. Fibulins are abundant in dynamic connective tissues associated to arteries, lung, skin, and ligaments (20), where they provide elastic recoil and resilience. Fibulins also provide an important adhesion template for cells, regulating growth factor availability (21) and participating in diverse supramolecular structures (22). The increased fibulin secretion observed in AS tissue culture suggests a loss of elasticity in the tissue, which together with other factors and proteins, may promote its calcification. Another fibulin, fibu-
lin-5 has also been implicated in severe hereditary connective tissue diseases such as supravalvular aortic stenosis (8).

Gelsolin is an actin binding protein with multiple regulatory activities that include cytoskeletal remodeling and ion channel regulation (23, 24). Increased gelsolin expression has been reported in failing hearts and it may contribute to the perturbation of thin filament organization (25, 26). Gelsolin has also been implicated in the cardiac remodeling triggered by ischemic heart disease, as described in both animal models of heart failure (15) and human hearts (27). We observed an increase in gelsolin expression in AS valves, which suggests that a similar remodeling process may occur in AS. Indeed, as a consequence of endothelial damage (28), endothelial cells are continuously exposed to mechanical forces generated by blood flow (29), and unidirectional shear stress increases the expression of proteins involved in adaptive processes to reduce flow-induced endothelium damage (18, 30). Moreover, the permanent decline in contractile function may also affect the expression of cytoskeletal proteins (31).

Pigment epithelium derived factor (PEDF), a glycoprotein of the serine protease inhibitor super-family, is a highly effective inhibitor of angiogenesis in cell culture and animal models (32, 33, 34), and it exerts antioxidant (35) and anti-thrombotic (36) effects. As oxidative stress is associated with an increase in cardiovascular risk (37), an increase in PEDF may protect cells from the negative effects of oxidative stress. Recently, an extensive study to analyze coronary plaques using a shotgun proteomic approach described increases in PEDF in atherosclerotic plaques (38). Hence, a link between atherosclerosis and aortic stenosis disease in relation to this concrete pathway may exist, which merits further investigation.

Angiotensinogen, clusterin, and pentraxin 3 differences found in label incorporation and secretion by LC-MS/MS were also correspondent with protein abundance validation analyses. T lymphocytes and macrophages infiltrate the endothelium and release cytokines, which in turn induce cellular proliferation of valve fibroblasts and extracellular matrix remodeling (39). This process results in increased secretion of proteins such as angiotensinogen (AGT), which exert wide-ranging effects on the endothelium and vascular smooth muscle such as vasoconstriction (40), because of its processing to angiotensin II peptide, which contributes to hypertension (21), and activation of endothelial proliferation (41). The increased secretion of AGT in AS valves may explain the thickening of the aortic valve due to its proliferative effect in valve fibroblasts, which precludes valve opening (5). Regarding prostaglandin D2 synthase (PGD2) protein, it inhibits platelet aggregation and induces vasodilatation (42). PGD2 is widely distributed in the peripheral tissues and localized in antigen-presenting cells and mast cells (43), from which it is released as an allergic and inflammatory mediator (44). We detected a decrease in PGD2 secretion in AS valves, implying a more limited vasodilatory effect, which may contribute to the vasoconstriction of aortic stenosis valves provoked at the cellular level.
level by the differentiation of mast cells to macrophages (23, 25). The PTX3 protein, which has recently been implicated in innate immunity, is thought to be a marker of vascular inflammation and damage (45, 46). Given that PTX3 is predominantly released by inflamed tissue in response to proinflammatory cytokines (TNF-α and IL-1β) (47) and, considering the recent proposal that AS may be an inflammatory disease, our findings would be consistent with this hypothesis, because PTX3 secretion was increased in AS valves. Secretion of clusterin protein was much lower in AS valves, in agreement with previous findings in ACS plasma (48), which highlights the occurrence of numerous alterations in protein expression common to both atherosclerosis and AS. Clusterin is a glycoprotein whose function remains unclear, although it has been implicated in complement regulation (49), acute phase response (50), lipid transport (51), apoptosis (52), and membrane protection at the tissue-fluid interface where it is expressed.

These data shows a subset of identified proteins differentially expressed in the secretome of AS human valvular tissue with clinical relevance. A further step in the discovery of novel tools for early diagnostic and prognostic evaluation of AS goes through the possibility to detect physiopathological changes in an easily accessible biological fluid. We evaluated here the variation of the trends of particular proteins in plasma from a wide cohort of patients and healthy subjects, finding gelsolin as a robust potential marker of disease. Gelsolin was found increased both in secretome and plasma AS samples compared with its level in healthy individuals. These data reflect the opposite trend to that found in the secretome from atherosclerotic coronary tissue compared with mammary (control) secretome (13), pointing to clear differences between the mechanisms of atherosclerosis and aortic stenosis. Pathway analysis of labeled proteins (either in control or pathological valves) revealed them as constituents of a network related to "Cellular Movement, Connective Tissue Disorders, Cardiovascular System Development and Function" with maximum significant score. Additionally, cellular movement, assembly and organization, and protein synthesis are functions significantly associated to this set of proteins, pointing to expected mechanisms involved in this pathology.

**Fig. 5. Validation of labeled proteins by SRM analysis.** Graphical representation of the abundance (upper panels) and representative extracted ion chromatograms (lower panels) of the transitions observed for angiotensinogen, EGF containing fibulin, gelsolin, and prostaglandin D2 synthase. Significant increases ($p < 0.05$) in the expression of angiotensinogen, EGF containing fibulin and gelsolin were observed in patients with respect to healthy subjects, whereas prostaglandin D2 synthase expression decreased significantly ($p < 0.01$) in AS patients with respect to healthy subjects. *In chromatograms: Isobaric or nearly isobaric peptides.
In conclusion, our results demonstrate the importance of secretome analysis in aortic stenosis, both to identify new potential biomarkers and to further understand the pathophysiology of this disease. The identification of specific protein secretion patterns altered with disease constitutes valuable information in terms of novel mechanisms of action, and a starting point to further investigate the possibility to act on these proteins through therapy. The use of metabolic labeling unequivocally ensures that all labeled proteins have been secreted or released by aortic valves. Importantly, particular proteins identified are involved in the atherosclerotic process, supporting the emerging hypothesis of a common link between both pathologies. On the other hand, gelsolin has here been demonstrated to be a high sensitivity biomarker of AS, with an opposite trend of secretion to plasma than atherosclerotic arterial tissue (13), which makes it a potential ideal discriminant biomarker of both pathologies. Our approach also exemplifies the development of specific and sensitive assays to validate biomarkers in the secretome. The proteins differentially secreted during aortic stenosis pathogenesis hereby reported, have been validated by two different techniques (WB and SRM) on an independent group of secretomes. In addition, plasma analyses corroborate the observed variations within the blood pointing to specific candidate markers to follow up in diagnosis, prognosis, and therapeutic intervention.

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