Technical Brief: A novel strategy for enrichment of trabecular meshwork proteome proteome

Renata Picciani, Anna K. Junk, Sanjoy K. Bhattacharya

Bascom Palmer Eye Institute, University of Miami, Miami, FL

We present a novel and simple enrichment strategy to capture trabecular meshwork (TM) proteome proteome. The method relies on fractionation of TM tissue into cytosolic and nuclear extracts and subsequent affinity enrichment of proteases on peptide inhibitors. A large repertoire of available protease substrate analog peptides enables an improved capture of TM protease proteome compared to SDS–PAGE fractionation alone. Peptide analog inhibitors of protease substrates are immobilized on a protein A or G column using 254 nm intense ultraviolet (UV) light. The TM cytosolic protein extract incubated on the column is eluted with salt or a buffer with a low hydrogen ion concentration. The resultant protein solution is precipitated with acetone, fractionated on SDS–PAGE, in situ trypsin digested, and subjected to mass spectrometry. This relatively simple protocol enables improved capture of cytosolic proteases. We identified 20 previously reported TM proteins from a single donor tissue using affinity enrichment. The majority of identified proteins were either intracellular proteases or known protease inhibitors. Both serine and cysteine proteases were captured using this strategy with improved coverage compared to our previous identification without affinity enrichment.

However, intracellular proteases also have the potential to provoke ECM changes by processing expressed matrix gene products [13,14] and modifying different forms of receptors/transporters that are important for entry and exit of key cellular homeostatic components such as the endothelin B receptor [15]. However, comprehensive studies in changes of TM intracellular proteases have not been performed. Recent investigations suggest that important alterations including impairment of intracellular proteases due to posttranslational modifications occur in the glaucomatous TM [6,14]. The changes in the intracellular protease proteome can be identified using high-throughput proteomic mass spectrometric approaches. One of the challenges in proteomic studies lies in reducing the complexity in capturing, identifying, and eventually quantifying the proteome changes [16]. Here, we present a simple and efficient method that allows better capturing of TM intracellular proteases. This method will enhance our understanding of TM intracellular protease changes that occur as part of glaucoma pathophysiology.

METHODS

Tissue procurement: Normal eyes (Table 1) were procured from the National Disease Research Institute (Philadelphia, PA) and the Lions Eye Bank (Miami, FL). The eyes were enucleated within 10 h of death, placed in a moisture chamber at 4 °C, and transported. These eyes were dissected within 48 h, and the TM was carefully excised for study. Fresh porcine TM tissue was isolated from freshly enucleated eyes from euthanized pigs procured from the University of Miami Department of Surgery following IACUC approved protocols.
Preparation of cytosolic extract: TM was carefully dissected from normal cadaver eyes. TM cytosolic and nuclear protein extracts were obtained using the NE-PER Nuclear and Cytoplasmic Extraction Reagents kit (Cat number 78833; Pierce Biotechnology, Rockford, IL) and following the protocols recommended by the manufacturer. The recovered proteins were subjected to spectrophotometric quantification using the Bradford assay and subsequently aliquoted for use or stored at −80 °C for future analysis. All protein aliquots were either freshly used or subjected to only one freeze–thaw cycle.

Western blot analyses: Approximately 10 µg of total cytosolic and nuclear protein extracts were fractionated on a 4%–20% Tris-glycine gradient gel (Invitrogen Corporation, Carlsbad, CA), transferred onto a polyvinylidene fluoride (PVDF) membrane, and incubated overnight at 4 °C with the antibodies (−5 µg/ml) detailed below. For these analyses, rabbit monoclonal antibody against histone H3 (Cat number 05–928; Upstate®, Billerica, MA) and rabbit polyclonal antibody against GAPDH (Cat number sc-25778; Santa Cruz Biotechnology Inc., Santa Cruz, CA) were used. Subsequently, a 2 h incubation at 4 °C with the appropriate horseradish peroxidase secondary antibodies was performed. All incubations occurred in 5% milk. Detection was performed using electrochemiluminescence (ECL; catalog number 32106; Pierce Biotechnology, Rockford, IL).

Preparation of enrichment column and capture of proteases: The enrichment column was prepared by ultraviolet (UV) cross-linking of peptides present in commercial protease inhibitor cocktails. Two different protease inhibitors (100 µl each; Protease inhibitor cocktail for use with mammalian cell and tissue extracts and Sigma Fast Protease inhibitor tablets for general use, Cat number P8340 and S8820, respectively; Sigma-Aldrich Inc., St. Louis, MO) were incubated with 200 µl (2.5 mg) of Protein A Sepharose beads (Cat number 32106; Pierce Biotechnology, Rockford, IL) was equilibrated with 1 ml 1X PBS. The samples were passed through the column three times and then washed with 250 µl 1X PBS twice. Subsequently, 100 µl of 1 M NaCl plus 100 mM KCl were used for elution. Finally, the elution products were subjected to acetone precipitation with the addition of four volumes of cold acetone and to incubation at room temperature for 15 min with or without carrier tRNA (Yeast tRNA; Catalog number M2271Lkit; AMRESCO Inc., Solon, Ohio). Approximately, 10 µg of the TM protein extracts (total cytosolic and nuclear extract) was also loaded on the 4%–20% Tris-glycine gel. On 4%–15% PHAST gel, 4 µg of cytosolic extract was also loaded in addition to eluents.

For mass spectrometric identification of proteins, gel slices were excised and digested in situ with sequencing grade porcine trypsin (Promega Biosciences Inc., San Luis Obispo, CA). The products of in situ digestion were loaded onto precolumns 360 mm od x 100 mm id fused silica (Polymeric Technologies, Phoenix, AZ) packed with 3 cm of irregular 5–15 µm non-spherical C18 (YMC Inc., Wilmington, NC). The gel was washed with 0.1 M acetic acid (AcOH) for 5 min before switching in-line with the resolving column (7-cm spherical C18, 360 od x 100 id). Once the columns were in-line, the peptides were gradient eluted with a 0%–100% B in 30 min where A was 0.1 M AcOH in nanopure water and B was 0.1 M AcOH in 80% acetonitrile. All samples were analyzed using a Thermo Electron Finnigan LTQ (San Jose, CA). Electrospray was accomplished using an Advion Triversa Nanomate (Advion Biosystems, Ithaca, NY) with a voltage of 1.7 kV and 200 nA. The mass spectrometer was operated in data dependent mode with the top five most abundant ions in each spectrum being selected for sequential tandem mass spectrometry (MS/MS) experiments. The exclusion list was used (1 repeat, 180 s return time) to increase the dynamic range. All MS/MS spectra were searched with Sequest version 2.7 (ThermoFinnigan, San Jose, CA) and Mascot using NCBI.

Table 1. Donor details.

| Age | Race | Gender | Time of death to enucleation (h) |
|-----|------|--------|---------------------------------|
| 55  | W    | M      | 8                               |
| 53  | W    | M      | 7                               |
| 62  | W    | F      | 10                              |
| 76  | W    | F      | 9                               |
| 55  | W    | M      | 9                               |

The Caucasian race is indicated by W. Letters M and F denote male and female, respectively.
non-redundant, Ensemble, and SwissProt databases. The search of database entries were restricted to Human and Metazoa (animals) and allowed for a maximum of two missed cleavages. The protonated molecule ions “MH+” and “Monoisotopic” were defined for the peak mass data input. The SEQUEST Sp and Xcorr cutoff scores of the protonated “MH+” peptides were 500 and 1.8, respectively. For Mascot searches, the score was greater than 78. All spectra were visually inspected for determination of correct database assignment. The potential chemical modifications of a peptide such as alkylation of a cysteine (carbamidomethyl [C]), oxidation of a methionine residue (M), and acetylation of a lysine (K) were also considered in the search. No restrictions were imposed for either the protein isoelectric point or the molecular weight during the search.

RESULTS

Enrichment of proteins in immobilized protease inhibitor peptide columns: Human TM samples were solubilized using buffers provided in the NE-PER Nuclear and Cytoplasmic Extraction Reagents kit (Pierce Biotechnology Inc.) and subjected to fractionation into cytosolic and nuclear fractions as described in Methods. The cytosolic fraction was subjected to affinity enrichment for proteases over a P8340 or a S8820 peptide inhibitor cocktail column. The column affinity enrichment resulted in the recovery of very small amounts of proteins. SDS–PAGE fractionation of proteins (4%–20% gradient mini gels; Invitrogen Inc.) did not allow detection of protein bands upon Coomassie Blue (Figure 1A) or silver staining (Figure 1B). The same amount of eluted proteins in a final volume of 4 µl was loaded onto a six-well PHAST gel (GE Healthcare) and subjected to fractionation and staining. Both Coomassie blue and silver staining allowed detection of the eluted bands (Figure 1C,D). To ensure the fractionation of cytosolic and nuclear proteins, the total protein, cytosolic, and nuclear fractions were subjected to western blot analyses (Figure 1E,F). When the blot was probed with antibody against GAPDH, only total and cytosolic extracts showed presence of GAPDH immunoreactivity (Figure 1E). On the other hand, probing the blot with histone H3 antibody revealed reactivity in the total and nuclear extracts and not in the cytosolic extract (Figure 1F), suggesting fractionation of total proteins into cytosolic and nuclear extracts.

Quantification of peptide column recovered proteins: The method presented here is to enable the capture of the protease proteome profile from single tissue samples. This method is aimed to eventually be applied to surgical TM tissue samples. While it is possible to quantify the proteins recovered from immobilized peptide columns with established biochemical methods for large initial protein loads (milligram quantities), the limited initial protein load in actual tissue samples prohibits quantification with the current biochemical methods (i.e., Bradford, Folin-Ciocalteau, Lowry, and Amido Black). For this purpose, we have performed experiments with porcine TM tissue to reduce or limit the use of precious human tissue for method development alone. Our studies indicate that reproducible relative quantification is possible for fractionated proteins on the PHAST gel system using silver staining and densitometry, but absolute quantification is not. Using porcine TM extract and densitometric scans of silver staining, the impact of the initial material (within a range of 50–500 µg) and time of incubation (1–16 h) was found to have an effect on the relative recovery of proteins (Figure 2). The densitometric scan of the recovered products from 550 µg of initial TM extract was incubated with immobilized peptide beads for 3 h, eluted with 1 M NaCl plus 100 mM KCl followed by acetone precipitation; 10 µg of carrier yeast tRNA was used as a reference (considered as 100 percent). This reference was used for calculation of the relative recovery ratios for all other elutions. To determine the effect of incubation time on relative recovery of eluted products, an initial load of 500 µg of porcine TM and was used, and elution was performed in an identical fashion as noted above.

Enhanced recovery as a function of carrier molecules and/or electrolyte addition. The recovery of proteins within a 50–250 µg range of initial material was found to be enhanced with the use of carrier molecules (yeast t-RNA). The electrolytes, a

![Figure 1. Analyses of SDS–PAGE fractionated proteins in enrichment steps.](Image)
combination of sodium and potassium, have an effect on the recovery of silver stained proteins. The addition of potassium chloride increased the recovery of proteins (data not shown). The addition of potassium chloride and carrier tRNA also reduced batch to batch variation in recovery especially with low initial loads (50–250 µg).

**Mass spectrometric identification of recovered trabecular meshwork proteases:** In normal TM, a total of 20 proteins were identified (Table 2). The majority of the identified proteins are proteases encompassing both serine and cysteine proteases (Table 2). Several protease inhibitors such as antichymotrypsin, anti-trypsin, and anti-thrombin were also identified (Table 2). Some non-protease proteins were also identified (Table 2). At this point, it is not well understood why these proteins/peptides showed strong binding. Proteases can bind to peptides, which are not their known high affinity substrate analog. Consequently, the TM proteases identified by affinity enrichment may or may not have a direct peptide inhibitor/analog. We identified all proteases from one single TM tissue in our current analyses that were identified from combined multiple numbers of TMs by proteomic analysis in a previous study [17]. The enhancement of identification in the current study is due to affinity enrichment of proteases. Although the total number of intracellular proteases identified are the same in our previous and in our current analyses (Table 2), they are different in two respects. First, all identified proteases are from a single TM tissue sample rather than from multiple TM samples, and second, all identified protease peptide matches (coverage) are greater than our previous analysis. In the previous analyses, fewer proteases were identified from a single TM tissue. All the proteases enlisted in Table 2 were identified only when six different TM samples were combined [17]. There are some non-protease proteins (for example, protease inhibitor anti-trypsin) for which fewer peptides were identified in the current analysis than in our previous analysis. However, overall a better capture of protease peptides was obtained with affinity enrichment as described above.

**DISCUSSION**

Glaucoma is thought to be due to increased resistance at the TM, which impedes aqueous humor outflow. Changes in the ECM are responsible for the altered resistance. Increased production of ECM proteins or, alternatively, processed secreted protein products may result in changes in the matrix composition and in ECM remodeling. Several factors such as elevated pressure or mechanical stretching have also been shown to influence ECM gene expression of the TM in vivo and in cultured TM cells [13, 18]. While the role of matrix metalloproteases has been well studied in TM remodeling [4, 10], changes in the TM intracellular proteases have been the subject of scant studies. Cellular proteasome have been found to undergo oxidative modification, resulting in the impairment of their function in the glaucomatous TM [6, 19, 20]. A comprehensive and quantitative investigation of the TM proteasomal activities also remains to be performed.

The proteomic analysis of clinically relevant tissues requires access to fresh or frozen specimens with adequate morphological consistency and sufficient quantity as well as proper and detailed clinical records [21]. The availability of such properly preserved tissues is often scarce especially ocular tissues. This fact limits meaningful outcome of proteomic analyses. Obtaining sufficient quantity of different ocular tissues is important for proteomic and genomic analyses, one providing complementary information to the other. Enrichment of proteins is a necessity due to the tiny nature of tissue regions in the case of ocular tissues. The TM is one such region where the availability of the amount of tissue material from a single donor is often limited (due to the tiny nature of the region), and without enrichment, successful identification of protease proteome will be severely limited.

Identification of proteins by mass spectrometry can be limited by several factors even for very sensitive mass
The proteins that mostly proteases identified by LC MS/MS of protein extracts were derived from the TM of a 55-year-old male Caucasian donor after enrichment on an immobilized immunocolumn and SDS–PAGE fractionation as described in Methods. Swiss-Prot database entries are shown; peptide matches in current mass spectrometric analyses; peptide matches and number of donors from our previous TM proteomic analyses [17] without affinity enrichment; identified protease inhibitors; identified proteins that are not proteases; cathepsin D has been identified as bovine and human protein in our previous and current TM proteomic analysis, here human protein accession number has been provided; not all complement components possess proteolytic activity; however, they act as a complex which results in proteolytic activity, here we have placed them in the protease category.

The immobilized inhibitors or inhibitory peptides have been in use for affinity purification. Immobilized peptide arrays have been used for epitope mapping as well as protein–protein interaction, enzyme-substrate interaction, and enzyme-inhibitor interaction studies [23]. Inhibitor affinity chromatography, using immobilized substrate analogs or other inhibitors, has been employed to isolate enzyme proteomes and is often referred to as the chemiproteomic approach [24]. Affinity enrichment of many different enzymes such as kinases, phosphatases, and matrix metalloproteases (MMPs) have benefited from the use of this approach [24–28]. However, such approaches are yet to be applied to the comprehensive enrichment of intracellular proteases, particularly in the ocular tissues [28,29]. Immobilized inhibitors have not only been used for affinity enrichment but also for several other novel uses [30–32]. Surface-immobilized, oriented peptide aptamers have been used for the detection of specific target proteins in complex biologic solutions [27]. Aprotinin-linked fibrinogen in a novel approach has been used for controlling degradation of fibrin gels [32]. The precursor state of factor Xa heparin-binding exosite was revealed by Xolaris binding to factor X in a novel fashion [31]. Label-free, high-throughput functional lytic assays based on immobilized inhibitors or substrates have made the development of refractive index-sensitive resonant waveguide grating biosensors possible [30]. Affinity enrichment of MMPs is a routine procedure for non-ocular tissues [28,29]. The approach outlined here can be used in combination with immobilized MMP target peptides and immobilized metal and gelatin QG columns to enable the capture of MMPs and their tissue inhibitors. Sequential capture of matrix and intracellular proteases from the same tissue is possible by combination of these techniques, maximizing the utility of tissue toward comprehensively discovering changes in the protease proteome composition in glaucoma (or any other disease) in comparison to the normal state. Additives or carrier molecules, which are usually proteins, electrolytes, and tRNAs, enhance recovery of biomacromolecules including proteins and DNA [33,34]. The carrier molecules usually act as chaperones and help entrap low amounts of protein following elution. Electrolytes typically promote charge-charge interaction and help in coagulation thus facilitating enhanced recovery. We have used two different inhibitor peptide cocktails to determine whether there are vast differences in their efficiency to capture target proteins. tRNA was added to enhance the recovery of proteins while it serves as a carrier molecule during acetone precipitation.

Compared to our previous analyses [17], current analyses captured a greater number of protease peptides from a single TM sample, demonstrating the utility and efficacy of the
fractionation strategy presented here. Several serine and cysteine proteases have thus been identified (Table 2). In particular, several complement components have also been identified. It is important to note that complement components have been found to impair synapses and have been implicated in neurodegeneration of retinal ganglion cells, a process that is thought to occur in glaucoma [35]. Our analyses also revealed the release and capture of mitochondrial neurolysin and tripeptidyl peptidase II (Table 2). Tripeptidyl peptidase II is a component of the proteolytic cascade acting downstream of the 26S proteasome in the ubiquitin-proteasome pathway [36,37]. Tripeptidyl peptidase II is thought to be able to complement the 26S proteasome function under conditions in which the latter is inhibited. Therefore, the expression of tripeptidyl peptidase II in normal TM suggests that perhaps in at least some eyes, the proteasomal function is impaired long before the pathology is apparent. Our efforts to capture protease proteome in the DBA/2J mouse model using combinatorial affinity enrichment methods outlined above is under progress to further our understanding of these proteases and their alterations.

ACKNOWLEDGMENTS

This work was supported by NIH grant EY15266 and a RPB career award (SKB). We thank Dr. Catherine Birt and Dr. Rajesh Dalvi for their helpful comments.

REFERENCES

1. Quigley HA. Number of people with glaucoma worldwide. Br J Ophthalmol 1996; 80:389-93. [PMID: 8695555]
2. Morrison JC, Acott TS. Anatomy and physiology of aqueous humor outflow. In: Morrison JC, Pollack IP, editors. Glaucoma Science and Practice. New York: Thieme Medical Publishers Inc.; 2003. p. 34-41.
3. Desmettre T, Rouland JF. Hypothesis on the role of nutritional factors in ocular hypertension and glaucoma. J Fr Ophthalmo 2005; 28:312-6. [PMID: 15883498]
4. Acott TS, Kelley MJ. Extracellular matrix in the trabecular meshwork. Exp Eye Res 2008; 86:543-61. [PMID: 18130351]
5. Buller C, Johnson D. Segmental variability of the trabecular meshwork in normal and glaucomatous eyes. Invest Ophthalmol Vis Sci 1994; 35:3841-51. [PMID: 7928181]
6. Govindarajan B, Laird J, Salomon RG, Bhattacharya SK. Isoleucylglutamin-modified proteins, including elevated levels of inactive calpain-1, accumulate in glaucomatous trabecular meshwork. Biochemistry 2008; 47:817-25. [PMID: 18085799]
7. Kaufman PL. Enhancing trabecular outflow by disrupting the actin cytoskeleton, increasing uveoscleral outflow with prostaglandins, and understanding the pathophysiology of presbyopia interrogating Mother Nature: asking why, asking how, recognizing the signs, following the trail. Exp Eye Res 2008; 86:3-17. [PMID: 18053966]
8. Johnson DH. Histologic findings after argon laser trabecuoplasty in glaucomatous eyes. Exp Eye Res 2007; 85:557-62. [PMID: 17822700]
9. Friedl P, Brocker EB. The biology of cell locomotion within three-dimensional extracellular matrix. Cell Mol Life Sci 2000; 57:41-64. [PMID: 10949580]
10. Sanka K, Maddala R, Epstein DL, Rao PV. Influence of actin cytoskeletal integrity on matrix metalloproteinase-2 activation in cultured human trabecular meshwork cells. Invest Ophthalmol Vis Sci 2007; 48:2105-14. [PMID: 17460268]
11. Alexander JP, Samples JR, Van Buskirk EM, Acott TS. Expression of matrix metalloproteinases and inhibitor by human trabecular meshwork. Invest Ophthalmol Vis Sci 1991; 32:172-80. [PMID: 1846130]
12. Ronkonk S, Rekoven P, Kaarniranta K, Puustjarvi T, Terasvirta M, Uusitalo H. Matrix metalloproteinases and their inhibitors in the chamber angle of normal eyes and patients with primary open-angle glaucoma and exfoliation glaucoma. Graefes Arch Clin Exp Ophthalmol 2007; 245:697-704. [PMID: 17028863]
13. Keller KE, Kelley MJ, Acott TS. Extracellular matrix gene alternative splicing by trabecular meshwork cells in response to mechanical stretching. Invest Ophthalmol Vis Sci 2007; 48:1164-72. [PMID: 17325160]
14. Sanchez-Sanchez F, Martinez-Redondo F, Aroca-Aguilar JD, Coca-Prados M, Escribano J. Characterization of the intracellular proteolytic cleavage of myocilin and identification of calpain II as a myocilin-processing protease. J Biol Chem 2007; 282:27810-24. [PMID: 17650508]
15. Rosenthal R, Choritz L, Zorn R, Munzer G, Fromm M, Pfeiffer N, Thieme H. Endothelin receptor B in trabecular meshwork. Exp Eye Res 2007; 85:482-91. [PMID: 17669399]
16. Righetti PG, Castagna A, Antonioli P, Boschetti E. Prefractionation techniques in proteome analysis: the mining tools of the third millennium. Electrophoresis 2005; 26:297-319. [PMID: 15657944]
17. Bhattacharyya SK, Rockwood EJ, Smith SD, Bonilha VL, Crabb JS, Kuchey RW, Robertson NG, Peachey NS, Morton CC, Crabb JW. Proteomics reveals cochlin deposits associated with glaucomatous trabecular meshwork. J Biol Chem 2005; 280:6080-4. [PMID: 15579465]
18. Gonzalez P, Epstein DL, Borras T. Genes upregulated in the human trabecular meshwork in response to elevated intraocular pressure. Invest Ophthalmol Vis Sci 2000; 41:352-61. [PMID: 10670462]
19. Caballero M, Liton PB, Challa P, Epstein DL, Gonzalez P. Effects of donor age on proteasome activity and senescence in trabecular meshwork cells. Biochem Biophys Res Commun 2004; 323:1048-54. [PMID: 15381105]
20. Caballero M, Liton PB, Epstein DL, Gonzalez P. Proteasome inhibition by chronic oxidative stress in human trabecular meshwork cells. Biochem Biophys Res Commun 2003; 308:346-52. [PMID: 12901875]
21. Paik YK, Kim H, Lee EY, Kwon MS, Cho SY. Overview and introduction to clinical proteomics. Methods Mol Biol 2007; 428:1-32. [PMID: 18287765]
22. Kim H, Eliuk S, Deshane J, Meleth S, Sanderson T, Pinner A, Robinson G, Wilson L, Kirk M, Barnes S. 2D gel proteomics: an approach to study age-related differences in protein abundance or isoform complexity in biological samples. Methods Mol Biol 2007; 371:349-91. [PMID: 17634592]
23. Reimer U, Reineke U, Schneider-Mergener J. Peptide arrays: from macro to micro. Curr Opin Biotechnol 2002; 13:315-20. [PMID: 12323352]

24. Tian R, Jiang X, Li X, Jiang X, Feng S, Xu S, Han G, Ye M, Zou H. Biological fingerprinting analysis of the interactome of a kinase inhibitor in human plasma by a chemproteomic approach. J Chromatogr A 2006; 1134:134-42. [PMID: 16982065]

25. Daub H, Godl K, Brehmer D, Klebl B, Muller G. Evaluation of kinase inhibitor selectivity by chemical proteomics. Assay Drug Dev Technol 2004; 2:215-24. [PMID: 15165517]

26. Lolli G, Thaler F, Valsasina B, Roletto F, Knapp S, Uggeri M, Bachi A. Inhibitor affinity chromatography: profiling the specific reactivity of the proteome with immobilized molecules. Proteomics 2003; 3:1287-98. [PMID: 12872229]

27. Johnson S, Evans D, Laurensen S, Paul D, Davies AG, Ko Ferrigno P, Walti C. Surface-immobilized Peptide aptamers as probe molecules for protein detection. Anal Chem 2008; 80:978-83. [PMID: 18186614]

28. Freije R, Klein T, Ooms B, Kauffman HF, Bischoff R. An integrated high-performance liquid chromatography-mass spectrometry system for the activity-dependent analysis of matrix metalloproteases. J Chromatogr A 2008; 1189:417-25.

29. Freije JR, Klein T, Ooms JA, Franke JP, Bischoff R. Activity-based matrix metalloprotease enrichment using automated, inhibitor affinity extractions. J Proteome Res 2006; 5:1186-94. [PMID: 16674108]

30. O'Malley SM, Xie X, Frutos AG. Label-free high-throughput functional lytic assays. J Biomol Screen 2007; 12:117-25. [PMID: 17175523]

31. Monteiro RQ, Rezaie AR, Bae JS, Calvo E, Andersen JF, Francischetti IM. Ixolaris binding to factor X reveals a precursor state of factor Xa heparin-binding exosite. Protein Sci 2008; 17:146-53. [PMID: 18042685]

32. Smith JD, Chen A, Ernst LA, Waggoner AS, Campbell PG. Immobilization of aprotinin to fibrinogen as a novel method for controlling degradation of fibrin gels. Bioconjung Chem 2007; 18:695-701. [PMID: 17432824]

33. Ambudkar SV, Maloney PC. Bacterial anion exchange. Use of osmolytes during solubilization and reconstitution of phosphate-linked antiport from Streptococcus lactis. J Biol Chem 1986; 261:10079-86. [PMID: 3090028]

34. Kishore R, Reef Hardy W, Anderson VJ, Sanchez NA, Buoncristiani MR. Optimization of DNA extraction from low-yield and degraded samples using the BioRobot EZ1 and BioRobot M48. J Forensic Sci 2006; 51:1055-61. [PMID: 17018081]

35. Stevens B, Allen NJ, Vazquez LE, Howell GR, Christopherson KS, Nouri N, Micheva KD, Mehalow AK, Huberman AD, Stafford B, Sher A, Litke AM, Lambris JD, Smith SJ, John SW, Barres BA. The classical complement cascade mediates CNS synapse elimination. Cell 2007; 131:1164-78. [PMID: 18083105]

36. Denis NJ, Vasilescu J, Libert JP, Smith JC, Figgey D. Tryptic digestion of ubiquitin standards reveals an improved strategy for identifying ubiquitinated proteins by mass spectrometry. Proteomics 2007; 7:868-74. [PMID: 17370265]

37. Tomkinson B, Wernstedt C, Hellman U, Zetterqvist O. Active site of tripeptidyl peptidase II from human erythrocytes is of the subtilisin type. Proc Natl Acad Sci USA 1987; 84:7508-12. [PMID: 3313395]