

Proteomics, glycomics, and glycoproteomics of matrisome molecules

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Abstract

The most straightforward applications of proteomics database searching involve intracellular proteins. While intracellular gene products number in the thousands, their relatively well-defined post-translational modifications (PTMs) makes database searching practical. By contrast, cell surface and extracellular matrisome proteins pass through the secretory pathway where many become glycosylated, modulating their physicochemical properties, adhesive interactions, and diversifying their functions. While matrisome proteins number only a few hundred, their high degree of complex glycosylation multiplies the number of theoretical proteoforms by orders of magnitude. Given that extracellular networks that mediate cell-cell and cell-pathogen interactions in physiology depend on glycosylation, it is important to characterize the proteomes, glycomes and glycoproteomes of matrisome molecules that exist in a given biological context. In this review, we summarize proteomics approaches for characterizing matrisome molecules, with an emphasis on applications to brain diseases. We demonstrate the availability of methods that should greatly increase the availability of information on matrisome molecular structure associated with health and disease.

Introduction

Viewed in evolutionary terms, the diversification of genes for intracellular versus extracellular proteins has separate drivers (1-3). Evolution of intracellular biology focuses on the regulation of signaling events, transcription and translation, through phosphorylation and other PTMs that influence allosteric enzyme regulation and signaling cascades through activation/deactivation of recognition domains, for example, SH2, SH3, bromo-, chromo- and tudor domains (4, 5). Significantly, many of the PTMs that occur inside the cell produce well-defined molecular additions (phosphorylation, acetylation, acylation and methylation) that are compatible with the established database searching workflows (5-7). Ubiquitination leaves a recognizable peptide tag after tryptic digestion that is amenable to proteomics approaches (8-10). Evolution of such PTMs arose through the need for complex control of gene expression through regulated signaling networks. By contrast, complex glycosylation reflects the evolutionary response to pathogen pressure and the evolving need for multicellular complexity (2, 3). Organisms need a multicellular organization with the ability to distinguish self from non-self and orchestrated responses to infection and regulated tissue plasticity. Therefore, much of cellular biology responds to signals received from the extracellular environment. Complex glycosylation is heterogeneous as a rule at each protein site, multiplying the number of molecular forms and requiring specialized proteomics methods.

As summarized in recent reviews (11-13), the mechanisms of tissue homeostasis and most diseases include interactions with the extracellular microenvironment. The matrisome constitutes the non-cellular components that control biochemical and biomechanical cues, growth factor and morphogen gradients, and physical scaffolds that define tissue phenotypes including morphogenesis, differentiation, and homeostasis (12). While each tissue has a unique extracellular environment, the number of gene products that code matrisome proteins in the entire body are limited (14-18). Cell surface receptors regulate adhesion and cytoskeletal connections to the matrisome. The structure and organization of the matrisome require maintenance as it

adapts to tissue growth needs. Matrisome proteins become glycosylated in the secretory pathway, may be proteolytically processed and cross-linked. The resulting physical and biochemical characteristics reflect the organized networks that depend on numerous molecular interactions that arise from PTMs.

To date, the ability to apply established proteomics methods that depend on database searching to such highly modified and heterogeneous proteins remains far from adequate (19). While we can detect many matrisome proteins using proteomics, the low sequence coverage leaves many structural elements of functional interest unidentified. In addition, the heterogeneity of glycosylation at each of the many glycosylation sites of matrisome proteins results in astronomically large numbers of possible proteoforms, if taken as multiples of the variants at each site. While the existence of such large numbers of proteoforms seems unlikely, the number of functional proteoforms that exist in a given biological context remains largely undefined.

In this review, we summarize methods for proteomics, glycomics and glyoproteomics of matrisome molecules. The goal is to characterize matrisome molecular structure in the greatest detail possible using wide-angle omics experiments. The high extent of matrisome protein glycosylation and other post-translational modifications requires special consideration of sample workup and proteomics database searching. We summarize matrisome physiology with emphasis on brain diseases. We summarize experimental approaches for matrisome workup and mass spectrometric analysis.

Extracellular matrix physiology and pathophysiology

Dysregulation of the cellular microenvironment occurs in cancers (20-22), neurodevelopmental and neuropsychiatric diseases (23, 24). Known as the matrisome, networks of extracellular matrix and cell surface molecules control the availability of growth factors to cellular receptors and the

mechanical-physical properties of the cell microenvironment. Currently, the limited understanding of the regulation of matrisome glycosylation hinders understanding of the roles of glycosylation-dependent matrisome networks in the basic mechanisms necessary for the targeted intervention of many diseases.

The extracellular environment (14-16) consists of glycoproteins, proteoglycans, collagens, and their interacting partners. Matrisome protein functions are elaborated by biosynthetic enzymes of the secretory pathway that generate mature molecules with spatially- and temporally-regulated glycosylation. Thus, glycoproteins have context-specific structures and biological functions that remain largely undefined due to the lack of effective methods for quantifying changes to site-specific protein glycosylation. This means that it is necessary to achieve complete matrisome protein coverage in order to determine the changes to these molecules that occur during disease mechanisms.

Progress in developing treatments that target interactions among cells and their extracellular microenvironments, including dysregulated cell growth, morphogenesis, and host-pathogen interactions, is limited by the ability to quantify the extent to which changes in matrisome networks resulting from altered glycoprotein glycosylation determine disease mechanisms. Many matrisome proteins contain lectin domains that recognize particular glycan epitopes. Thus, the glycosylation of matrisome molecules determines their binding interactions with other matrisome molecules and with soluble growth factors. The result is organized assemblies of matrisome molecules that compose the spatially and temporally regulated microenvironments through which cells receive signals in physiology and pathophysiology.

The core matrisome consists of 195 glycoproteins, 44 collagens and 35 proteoglycans (16, 18, 25), many of which are glycosylated. Extensive proteomics studies have cataloged the abundances of matrisome molecules in different tissues (18); however, such studies have not defined the glycosylation states of matrisome molecules necessary to define networks of

interactions with lectin-containing binding partners. Many matrisome molecules have several points of glycosylation, each with microheterogeneity, each representing a functional epitope that needs to be defined in order to characterize biological functions (26).

Glycan branching regulates glycoprotein dynamics and the residency of cell surface cytokine receptors (27). Interactions among cell surface and extracellular glycoproteins with lectins including galectins, C-type lectins, and siglecs, drive the clustering of cell surface molecules into networks that define tissue microenvironments. These organized assemblies of extracellular molecules (the matrisome) adapt cellular microenvironments to phenotypic needs. While glycosylated proteins represent potential therapeutic targets, their macro- and micro-heterogeneities pose a significant challenge to exploitation. Because their functions depend on glycosylation and other PTMs, it is necessary to produce detailed proteolytic maps of matrisome proteins and follow the changes that occur during aging and disease development.

The dynamic brain matrisome

The brain extracellular space has been referred to as the final frontier in neuroscience (28). Many of the matrisome molecules common to systemic organs are not found in brain (29). Matrisome functions depends on networks of interaction among glycosylated proteins and glycan-binding lectins. As illustrated in Figure 1 for the brain, networks of cell surface and extracellular glycoproteins and proteoglycans bind many families of growth factors and growth factor receptors (30-32). They modulate receptor tyrosine kinase signaling pathways at the heart of mechanisms including tissue stiffness and growth factor transport.

In the brain, the extracellular space occupies ~20% of brain volume (28, 33). As shown in Figure 1, matrisome in the central nervous system includes the interstitial matrix, basement membranes, and perineuronal nets (PNNs), the fine structures of which vary spatially and temporally (34, 35).

The matrisome provides the environment necessary for cell homeostasis, repair, regeneration, and neural plasticity in a brain region-specific manner (36-38).

The basement membranes that line cerebral blood vessels consist of collagen IV, laminins and heparan sulfate proteoglycans (HSPGs) perlecan and agrin (39, 40). In the brain, neural interstitial matrix separates cells and consists of networks of chondroitin sulfate proteoglycans (CSPGs), tenascins, hyaluronan, and link proteins. The PNN consist of many of the same CSPGs, tenascins, hyaluronan, and link proteins condensed that surround some neuronal cell bodies and dendrites.

Dysregulation of matrisome molecular networks characterize pathophysiologies of cancer (41, 42), autoimmune (43-45), cardiovascular (46-48), and neuropsychiatric diseases (23, 24, 34). In the brain, region-specific regulation of matrisome molecule glycosylation controls the neuronal microenvironment and becomes dysregulated in neuropsychiatric diseases (23, 24, 34, 49). For example, in neurodegeneration, proteoglycans bind to and play roles in the aggregation of proteins including A β , tau, prion protein, and α -synuclein (50, 51). In Alzheimer's disease, proteoglycans participate in amyloid plaque formation, leading to disease pathology from altered proteolytic processing and accumulation of toxic aggregates (52, 53).

Perineuronal nets are lattices of matrisome molecules that surround the cell body and dendrites of neurons. They are thought to serve as a reservoir for cations and provide the connective architecture that controls synaptic plasticity. Deficits in PNN structure appear to contribute to dysfunction in cortical circuitry in schizophrenia (54). The number of PNN in visual cortex increases during postnatal development, paralleling the critical period for synaptic plasticity and playing an important role in critical period closure. Significantly, the adult inability to repair spinal cord injury can be restored by the treatment of the injury site with chondroitinase enzymes (55, 56) with the implication that CSPGs maintain the extracellular environment in adult neural tissue that limits neural plasticity (37, 57-59). PNN structure differs spatially and temporally in the brain

in association with injury, repair, development, aging, learning, memory, neuropsychiatric diseases, neurodegeneration, and in response to drug abuse (24, 60, 61).

Variation in matrisome molecular structure among brain regions, with development, aging, and pathologies. Traditional antibody-based techniques including immunohistochemistry show spatial and temporal regulation of matrisome molecule expression in the brain (62-64). While antibody binding indicates the levels of individual epitopes, the antibody specificity and underlying structure are assumed. Matrisome molecule glycosylation can also be stained using lectins, leaving the underlying matrisome site-specific glycosylation structure undefined (24, 54, 65).

Because antibodies reflect the presence of a particular structural epitope on highly complex matrisome molecules, the changes in structure unrelated to such epitopes are not defined. This is illustrated for the aggrecan proteoglycan in Figure 2. Aggrecan contains three globular domains and an extended region modified with more than 100 CS chains. The C-terminal G3 domain has EGF-like repeats, a complement regulatory module, and a C-type lectin module (66) and interacts with tenascins. The globular domains are *N*-glycosylated, and the extended domains also carry keratan sulfate, mucin-type *O*-glycans, and *O*-mannose glycans (67). While this provides a parts list for aggrecan, we have little information on how the variation of glycosylation modulates the functions of aggrecan in a weight-bearing tissue such as cartilage versus brain. Further, while it is clear that CS structure varies among brain regions, there is little information on how aggrecan structure varies and impacts on the fine structures of the resulting matrisome molecular networks. The same reasoning applies to the other brain matrisome network glycoproteins, proteoglycans, and collagens.

It is clear, however, that matrisome glycosylation changes during development and with disease states. This can be seen from the alteration of CS sulfation during development in many tissues, including cartilage (68) and brain (69, 70). Further, staining of brain tissue with *Wisteria floribunda* agglutinin (WFA), a lectin that binds GalNAc residues, has been used to identify brain region

pathologies associated with schizophrenia (71-74). The extreme complexities of matrisome proteins concerning glycosylation, cross-linking and other PTMs, drives the need for many more validated antibody reagents than are available to date (64). Such lectin and/or antibody staining studies do not define the glycosite changes that underlie dysregulated interactions with lectin-containing binding partners that give rise to pathologies.

Matrisome proteomics

Matrisome sample preparation methods. As summarized in Table 1 and reviewed in detail (75), methods for enrichment of matrisome proteins for proteomics studies include tissue decellularization and extraction of matrisome components from tissue homogenates. Proteomics researchers often use decellularization to remove cellular components prior to solubilization of matrisome molecules (76-84). The use of a chaotrope solubilizes some matrisome components, leaving an insoluble pellet rich in fibrillar collagens (85). The yield of such matrisome components improves with chemical digestion with cyanogen bromide (86). Alternatively, hydroxylamine cleavage at Asn-Gly sites has also been used to solubilize matrisome from insoluble pellets prior to tryptic digestion and proteomics (82, 86). For matrisome protein quantification among tissue sample cohorts, Hansen *et al.* have developed targeted proteomics (87-89).

Proteomics data acquisition methods for the analysis of matrisome proteins. Present discovery proteomics methods suffice to identify matrisome proteins based on the presence of minimally modified peptides using database searching. Such peptides have been used in targeted proteomics assays for quantification of matrisome molecules based on inferred core protein abundances (87-89). Data-independent analysis (DIA) has the advantage that all precursor ions are subjected to collisional dissociation. Using the sequential window acquisition of all theoretical fragment ion spectra (SWATH)-MS DIA method (90), fragment ion spectra for all precursors are

acquired within the specified m/z range and retention time window. Interpretation of such datasets in which tandem mass spectra show product ions from co-eluting peptides requires the use of spectral libraries (91). Huang *et al.* developed a spectral library of 201 matrisome proteins and compared the performance of SWATH versus data-dependent acquisition (DDA) for analysis of unfractionated tissue extracts (92). They reported a 15-20% improvement in peptide reproducibility and a 54% increase in a number of matrisome proteins identified relative to DDA. Önnarfjord *et al.* used high pH reversed-phase fractionation of tryptic digests as a workflow for cartilage proteomics (93). Due to the additional fractionation step, they reported 653 proteins identified. They used DDA data to build spectral libraries for interpretation of a subset of identified proteins using DIA. They showed that DIA produced a more precise measurement of peptide abundances than DDA.

Naba *et al.* used a commercial Cytosol/Nucleus/Membrane/Cytoskeleton compartmental protein extraction kit to enrich intracellular and matrisome proteins in separate fractions from tissue (18, 94-96). The tryptic digests of the matrisome -enriched pellets were solubilized using urea prior to LC-tandem MS (97). This approach resulted in the identification of approximately 250 matrisome proteins from tissue (94, 95, 98, 99).(100).

Mayr *et al.* extracted matrisome proteins from vascular tissue using decellularization, solubilization using a three-stage extraction (salt, detergent, guanidine HCl) and MS-based quantification (101-105). In studies of human veinous tissue, they reported the identification of ~150 matrisome proteins. They identified a proteomics 4-biomarker signature for atherosclerotic plaques from a comparison of vascular matrisome in human carotid artery specimens (106). In this work, they report 110 matrisome -associated proteins from guanidine HCl extraction and 87 from the salt fractions with an overlap of 51. They also performed matrisome proteomics studies of restenosis and thrombosis following coronary stent implantation in pigs, for which they report the identification of 151 matrisome proteins (107, 108).

Berretta *et al.* demonstrated using a combination of immunohistochemistry and proteomics that matrisome molecule expression is brain region-dependent (34). For this work, fresh rat brains were dissected, and regions were snap frozen. Tryptic peptides were fractionated using ERLIC, and the resulting fractions analyzed using LC-MS. The fold change abundances of a set of 17 matrisome molecules, including tenascins, hyalactans, link proteins, and others were reported for a set of five rat brain regions.

MALDI imaging of matrisome proteins. As described in detail in recent reviews (109-111), MALDI imaging mass spectrometry (IMS) produces 2-dimensional maps of the distributions of ions desorbed from the surface of tissue slides. The advantage is that the maps can be produced at approximately 25 μm or better resolution and demonstrate impressive ion-specific spatial resolution patterns. The disadvantages are that in the absence of a separation step, the dynamic range of protein/peptide detection is limited, and identification of observed proteins or peptides can be cumbersome. Drake *et al.* have demonstrated the use of MALDI-imaging to visualize proteins and peptides from matrisome-rich tissues, including cardiac (112). They demonstrated the use of matrix metalloproteinase enzymes to localize collagen and elastin peptides on the surfaces of the tumor and cardiac tissue slides (113). This group also pioneered MALDI-based imaging of glycans at approximately 25 μm spatial resolution on tissue slides (114, 115) for which they derivatized sialic acid residues to prevent dissociation resulting from the MALDI process.

Matrisome glycomics and proteomics from histological slides. We developed a workflow for profiling GAGs, *N*-glycans, and proteins from tissue slides (116, 117). We applied the method to comparative glycomics profiling from invertebrates (118-120), mammalian organ tissues (121-123), skeletal muscle (124), kidney tissues (125, 126) leukocytes (127), stem cell niche (128), and tumor tissues (117, 129). We have investigated brain aging (130) and neuropathological diseases including glioma (117). Our method provides a readout of GAG quantities, domain structures, and non-reducing end structures using simple enzyme digestions with minimal need

for workup. The final proteomics of tryptic peptides identifies ~1200 proteins from the 10 nL tissue volume, providing deeper coverage than can be obtained from an MS imaging approach.

This approach requires small tissue volumes, minimal sample workup, and reduces the effort required per biospecimen for glycomics and proteomics studies. Fresh frozen slides are washed with a series of solvents, thereby denaturing tissue proteins. Formalin-fixed, paraffin embedded tissue slides require dewaxing, re-hydration, and high pH antigen retrieval prior to enzymatic digestion. While proteins are denatured in both cases, the observed glycomics and proteomics profiles reflect tissue processing biases that remain to be studied in detail. Nonetheless, the analysis of matrisome molecules from tissue slides offers an attractive option to extraction from wet tissue in terms of lower sample quantities and effort required. For example, in a study of aging rat brain from tissue slides with no enrichment, we observed 9-11% of total proteins of extracellular origin, corresponding to 15 matrisome molecules (130).

Matrisome glycoproteomics

Application of a conventional discovery proteomics workflow with database searching identifies Matrisome molecules based on the presence of unmodified peptides. While homogeneous PTMs including phosphorylation, acetylation, methylation, and ubiquitination are amenable to proteomics database searching (131, 132), glycosylation is heterogeneous as a rule. This multiplies the number of PTM forms of a given matrisome molecule glycopeptide, thereby dividing the precursor ion signals, and multiplying the size of the proteomics search space and the difficulty of assigning the glycopeptide with confidence (133-139). As shown in Figure 3, the presence of complex glycosylation alters the collisional dissociation pattern of peptides significantly. Glycopeptide collisional dissociation tandem mass spectra show low m/z oxonium signature ions that indicate the presence of glycosylation. The spectra also show peptide+saccharide ions, the abundances of which depend on the extent of vibrational excitation of the precursor ions. If relatively low collision energies are used, then product ions resulting from losses of saccharide

units are abundant. At higher collision energies, peptide plus from one to a few monosaccharide units are observed. Under such conditions, dissociation of the peptide backbone is often observed albeit at relatively low abundances. Thus, the most confident collisional tandem mass spectra for glycopeptide precursor ions contain all three ion types as shown for example for an aggrecan glycopeptide in Figure 3 (140).

Lectin enrichment of glycopeptides. Investigators have used WFA and concanavalin A (ConA) lectin enrichment of guanidine HCl extracts to enrich glycoproteins from human cardiac tissue from which they reported identification of 65 glycosylation sites from 35 extracellular proteins (141, 142). Wheat germ agglutinin (WGA) was used to enrich glycoproteins from mouse brain to identify O-mannosylated peptides from neurofascin 186 (143) and PNN associated hyaluronan proteoglycans (67). O-Mannosylated peptides have been enriched from tissue extracts digested using trypsin and peptide-N-glycosidase F using ConA lectin chromatography (144, 145). A set of 16 O-mannosylated glycoproteins were identified, several belonging to the cadherin superfamily, using this approach.

Proteoglycan glycoproteomics. Enzymatic digestion of GAG chains leaves a glycopeptide with linker saccharide attached to the core protein. Such linker glycopeptides can be identified by the presence of a diagnostic oxonium ion for CS and HS proteoglycans (146). The linker saccharide glycopeptides detected for CSPGs were modified with sulfate, phosphate, fucose and/or sialic acid (147-149). This approach has been used to analyze PGs from biological fluids including urine and cerebrospinal fluid (149). We used a similar approach for analysis of purified proteoglycans including aggrecan, decorin, brevican and neurocan (140). In order to interpret the glycopeptide tandem mass spectra automatically from the LC-tandem MS datasets, we optimized our GlycReSoft software (150, 151) for interpretation of linker-glycopeptides.

We have observed glycopeptides abundances too low for confident identification when analyzing proteolytic digests from tissue slides. It, therefore, appears that enrichment steps will be

necessary to allow glycoproteomics from the tissue. While such enrichment remains a challenge from small tissue volumes such as obtained from tissue slides, it seems feasible from wet tissue extracts.

Conclusions

For researchers interested in profiling abundances of core matrisome proteins, the use of decellularization or enrichment methods combined with targeted MS or DIA MS seems appropriate. As in other areas of proteomics, the use of multidimensional separations increases the number of proteins identified at the expense of analysis time and cost. One of these separation dimensions can be designed to enrich glycopeptides, thus increasing the ability to detect matrisome determinants of molecular networks. Such enrichment steps are most readily applied to tissue extracts. The analysis of tissue slides has potential benefits in terms of throughput, cost, and applicability to pathological workflows. The tissue volume, however, is rather low, making use of enrichment steps challenging. On the other hand, tissue slides can be microdissected (152), increasing the ability to select cell populations of interest for subsequent proteomics. Robotic approaches for manipulation of microdissected tissue have been described (153, 154). It may, therefore, be feasible to use glycopeptide enrichment in such robotic workflows to enable the application of glycoproteomics LC-MS methods to microdissected tissue. This will enable profiling of designated matrisome glycosites as a means for assessing changes to extracellular networks during disease mechanisms.

Acknowledgements

The authors acknowledge support from NIH grants P41GM104603 and U01CA221234.

Figures

Figure 1. Brain matrisome types include blood-brain barrier, interstitial matrix, and perineuronal nets. These structures are composed of matrisome molecules including hyaluronan, collagens, glycoproteins, and proteoglycans. Matrisome structure is spatially and temporally regulated, dynamic, and becomes altered during the pathogenesis of neuropsychiatric and neurodegenerative diseases.

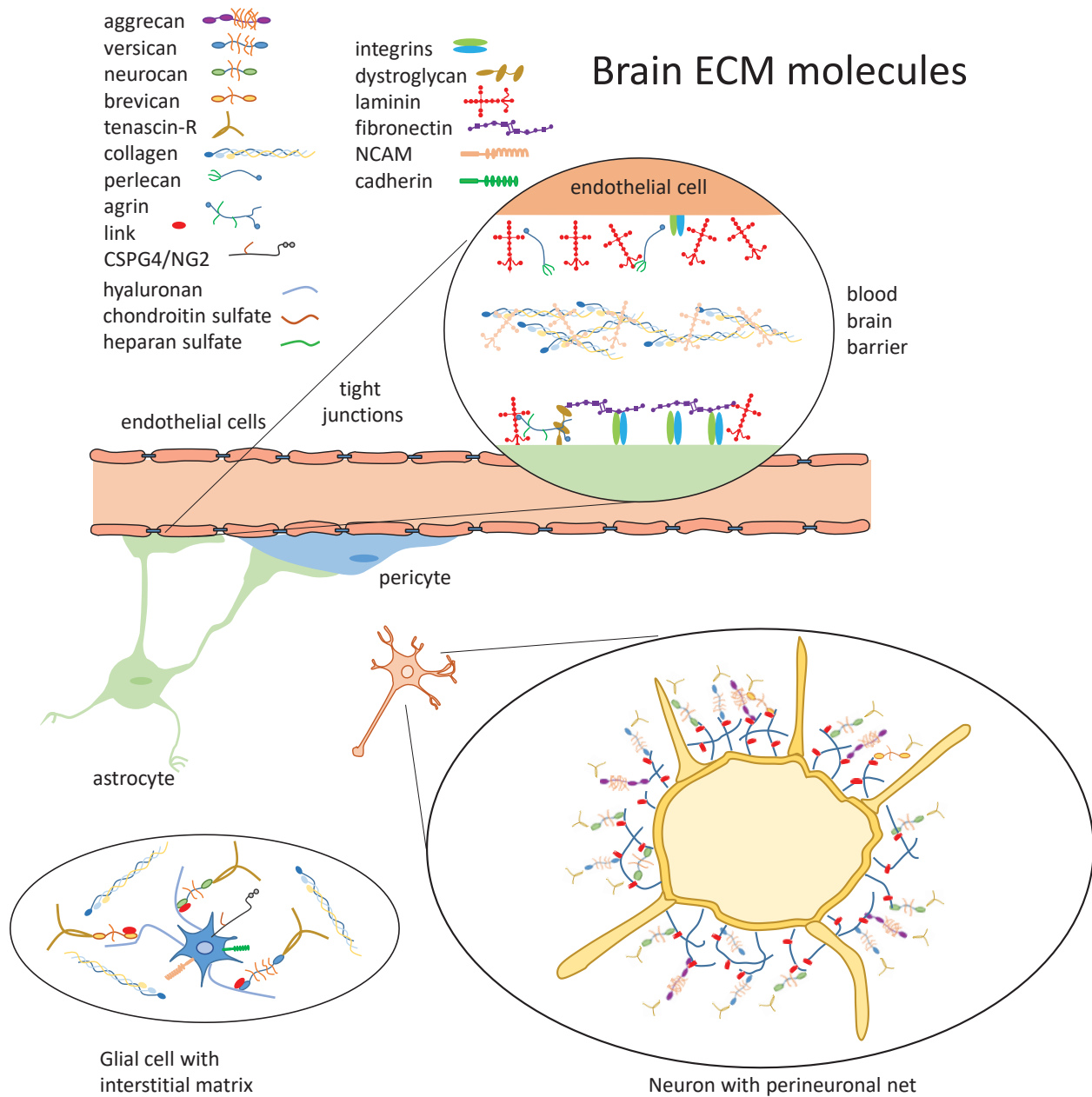


Figure 2. Model for the structure of aggrecan, showing glycosylation with CS, HS, *N*-glycans, mucin-type *O*-glycans, and *O*-mannose glycans.

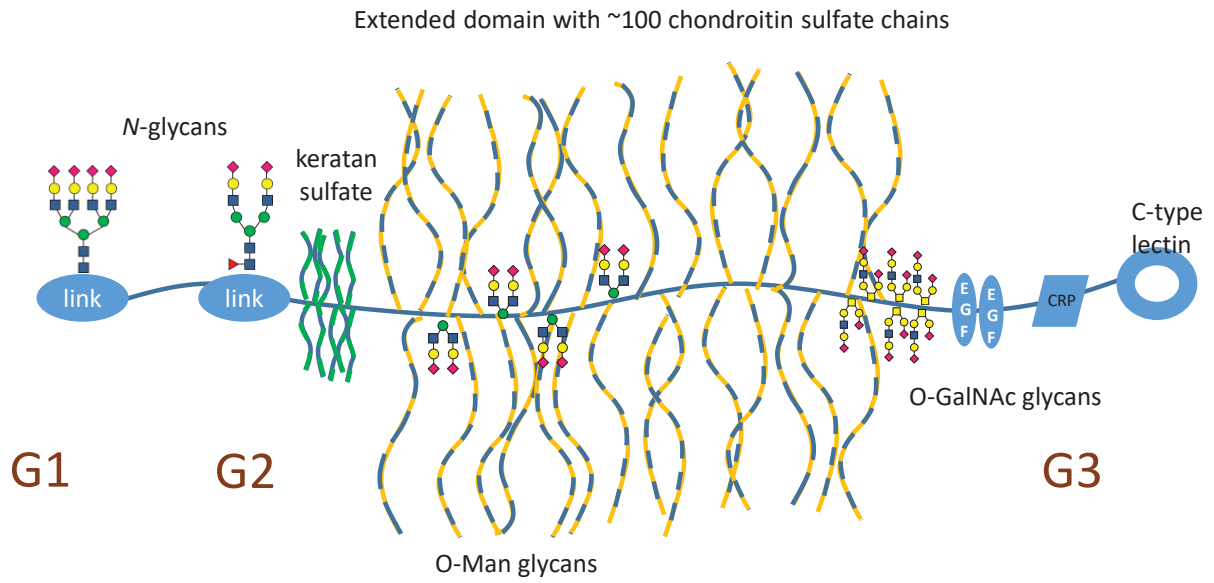


Figure 3. Higher energy collisional dissociation tandem mass spectrum of an aggrecan glycopeptide showing the presence of oxonium ions (green), peptide+saccharide ions (golden), and peptide backbone dissociation ions (red or blue) (140).

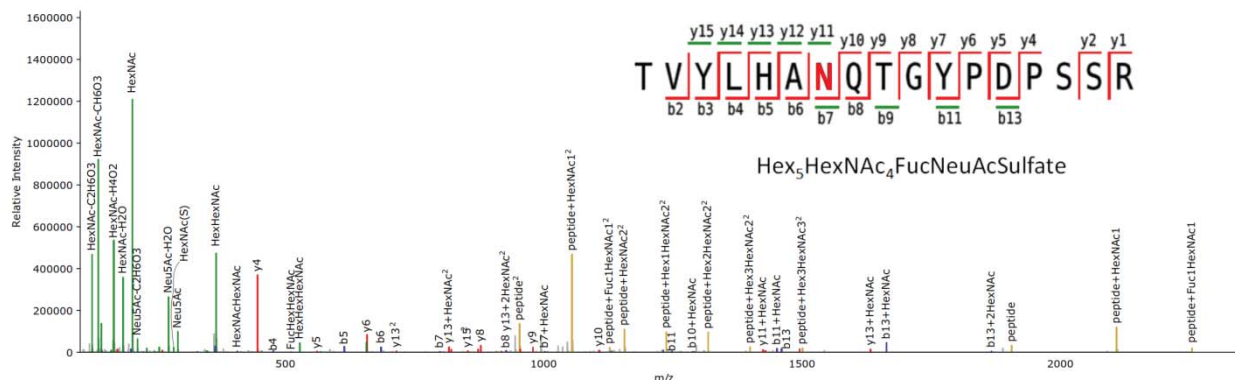


Table 1. An overview of proteomics approaches for matrisome Proteins.

Enrichment method prior to LC-MS/MS analysis	Tissue type	Number of matrisome proteins identified	Reference	Limitations
High pH reversed phase fractionation	Human knee articular cartilage	196 (out of total 653 protein IDs)	Önnerfjord <i>et al.</i> (93)	Additional fractionation step
Commercial compartmental protein extraction kit and urea solubilization	Murine normal and melanoma tumor tissues	200	Naba <i>et al.</i> (18)	Semi-quantitative estimates
Decellularization, solubilization using a three-stage extraction (salt, detergent, guanidine HCl)	Atherosclerotic plaque tissue	~200	Mayr <i>et al.</i> (106)	False positives and limited list of proteins
Decellularization, solubilization using a three-stage extraction (salt, detergent, guanidine HCl)	Pig coronary arteries	151	Mayr <i>et al.</i> (155)	limited availability of porcine antibodies
ERLIC off-line fractionation	Rat brain	17	Berretta S <i>et al.</i> (34)	Additional fractionation step
On-slide tissue digestion (no enrichment)	Rat brain (Striatum and Substantia-nigra)	15	Zaia J <i>et al.</i> (130)	Low protein coverage

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