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# Cardiac extracellular proteome profiling and membrane topology analysis using glycoproteomics

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#### Abstract

**Purpose**—Extracellular proteins are easily accessible, which presents a sub-proteome of molecular targets that have high diagnostic and therapeutic potential. Efforts have been made to catalogue the cardiac extracellular matridome and analyze the topology of identified proteins for the design of therapeutic targets. Although many bioinformatics tools have been developed to predict protein topology, topology has been experimentally validated for only a very small portion of membrane proteins. The aim of this study was to use a glycoproteomics and mass spectrometry approach to identify glycoproteins in the extracellular matridome of the infarcted LV and provide experimental evidence for topological determination.

**Experimental design**—Glycoproteomics analysis was performed on eight biological replicates of day 7 post-MI samples from wild type mice using solid-phase extraction of glycopeptides, followed by mass spectrometric identification of *N*-linked glycosylation sites for topology assessment.

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**Results**—We identified hundreds of glycoproteins and the identified *N*-glycosylation sites provide novel information on the correct topology for membrane proteins present in the infarct setting.

**Conclusions and clinical relevance**—Our data provides the foundation for future studies of the LV infarct extracellular matridome, which may facilitate the discovery of drug targets and biomarkers.

#### Keywords

Extracellular matridome; glycoprotein; membrane orientation; matrix metalloproteinase; proteomics; myocardial infarction; left ventricle

#### 1 Introduction

The extracellular matridome consists of all proteins expressed outside of cells, including transmembrane proteins, cell surface proteins, extracellular matrix (ECM) proteins, and secreted proteins [1]. The composition includes receptors, growth factors, cytokines, chemokines, hormones, enzymes, and fibrillar components [2]. The cardiac extracellular proteome provides the left ventricle (LV) with mechanical support, coordinates the signal transduction capabilities, and regulates cell functions by modifying biological processes [3]. Myocardial infarction (MI) is associated with extensive extracellular protein turnover as old ECM is replaced by an infarct scar that is primarily composed of ECM. MI is a highly prevalent cardiovascular disease, with over 1.5 million new patients diagnosed each year in the U.S. [4]. LV remodeling following MI depends on the balance between ECM degradation and deposition, as too much degradation can lead to LV aneurysms or rupture and too much deposition can lead to a stiff LV that provides a substrate for the development of heart failure [5]. Matrix metalloproteinase-9 (MMP-9) is a member of the family of enzymes that break down ECM and has been shown to play a critical role in LV remodeling post-MI [6, 7].

Knowledge of protein structure provides crucial information for understanding protein function, including information on location and availability for post-translational modification that is necessary for selecting optimal antigen sites and identifying drug targets. A common feature for optimal antigens and drug targets is easy accessibility, making extracellular protein analysis highly relevant to drug development for LV remodeling post-MI [8]. Therefore, efforts have been made to determine the spatial orientation of target proteins. In addition to the traditional methods for topology evaluation, membrane protein side accessibility is becoming a tool to evaluate protein orientation. Examples of this concept include *N*-glycosylation sites, antibody epitopes, iodinatable sites, and proteolytic sites [8]. The reason *N*-glycosylation sites can be used for topology evaluation is based on the fact *N*-linked glycosylation occurs only in extracellular domains of membrane proteins [9].

The goal of this study was to examine the geographic location of proteins in the LV infarct region, focusing on the extracellular matridome. Glycoproteomic analysis was performed on eight biological replicates of day 7 post-MI samples from wild type mice using solid-phase

extraction of glycopeptides, followed by mass spectrometric identification of *N*-linked glycosylation sites for topology assessment [10, 11]. The logic of this examination was based on the concept that extracellular proteins are frequently *N*-glycosylated [12–14]; therefore, preferential isolation of N-linked glycoproteins would greatly enrich for the extracellular matridome [1, 13, 15]. In the present study, we identified 1352 *N*-linked glycosylation sites, and 56% of them were from membrane proteins proteins. Since membrane *N*-glycans always face the extracellular space, the identification of these glycosites provides topological information and helps to determine the protein orientation.

#### 2 Materials and methods

#### 2.1 Mice

C57BL/6J male and female mice, 3–6 months of age, were used in this study (n=8, 4 male and 4 female). Mice were kept in a light-controlled environment with a 12:12 hour light-dark cycle and given free access to standard mice chow and water. All animal procedures were approved by the Institutional Animal Care and Use Committee at the University of Texas Health Science Center at San Antonio and the University of Mississippi Medical Center in accordance with the "Guide for the Care and Use of Laboratory Animals". The mice underwent permanent coronary artery ligation surgery, to produce myocardial infarction, as described previously [16, 17]. At day 7 post-MI, the mice were sacrificed.

#### 2.2 Tissue samples and protein extraction

The infarct region of the LV was collected at 7 days post-MI as described previously [17], and the LV tissue was homogenized first in phosphate buffered saline (PBS; 16  $\mu$ L per mg LV wet weight) with 1x protease inhibitors (Roche, Basel, Switzerland) and centrifuged to remove the soluble fraction. The insoluble pellet was homogenized in Reagent 4 (16  $\mu$ L per mg LV wet weight; Sigma, St. Louis, MO) with 1x protease inhibitors [18]. Because the insoluble fraction is enriched for ECM, we used that fraction for the glycoproteomic analysis.

#### 2.3 Trypsin digestion

Total protein from the insoluble fraction (1 mg) was denatured and reduced by adding 8 M urea and 12 mM Tris (2-carboxyethyl) phosphine hydrochloride (TCEP) in 1 M ammonium bicarbonate and incubating at 37°C for 1 h. The samples were alkylated with 16 mM iodoacetamide at room temperature in the dark for 30 min. The samples were diluted 5-fold with water prior to the addition of trypsin (Promega, Madison, WI), which was added at a ratio of 1:50 (w/w, enzyme: protein). The samples were digested at 37°C overnight with gentle shaking. The samples were centrifuged to remove precipitate. Peptides were cleaned by Sep-Pak Vac C18 cartridge (Waters, Milford, MA). Peptide concentration was determined by BCA assay (Thermo Fisher Scientific, Rochford, IL).

#### 2.4 N-Linked Glycopeptide Capture

*N*-linked glycopeptides were isolated from the tryptic peptides using the solid phase extraction of glycopeptides (SPEG) method previously reported [10, 11]. Briefly, 0.8 mg of peptides was oxidized by 10 mM sodium periodate (Bio-Rad, Hercules, CA) at room

temperature for 1 h. Glycopeptides were covalently conjugated to a solid support via hydrazide chemistry (Bio-Rad, Hercules, CA), with incubation at room temperature overnight. The hydrazide beads were washed with 1.5 M NaCl, and water prior to release of formerly *N*-linked glycopeptides from solid support by PNGase F (New England Biolabs, Ipswich, MA) at 37°C overnight. The eluent was purified by Sep-Pak Vac C18 cartridge (Waters, Milford, MA) and re-suspended in 20 μL of 0.4% acetic acid.

#### 2.5 Mass spectrometry analysis

The peptides were analyzed by LC-MS/MS using a Q Exactive (ThermoFisher, Waltham, MA) coupled with a 15 cm  $\times$  75 µm C18 column (5 µm particles with 100 angstrom pore size). A nano UPLC at 300 nL/min with a 90 min linear acetonitrile gradient (from 5–35% B over 90 min; A = 0.2% formic acid in water, B = 0.2% formic acid in 90% acetonitrile) was used. A top 20 data dependent MS/MS with exclusion for 25 s was set. The samples were run with HCD fragmentation at normalized collision energy of 30 and an isolation width of 2 m/z. A lock mass of the polysiloxane peak at 371.10123 was used to correct the mass in MS and MS/MS. Target values in MS were 1e6 ions at a resolution setting of 70,000 and in MS2 1e5 ions at a resolution setting of 17,500.

MS/MS spectra were searched with SEQUEST using Proteome Discoverer (version 1.3; Thermo Fisher) against the mouse IPI database (version 3.87) containing 59,534 sequences. For this database search, the precursor mass tolerance and fragment mass tolerance were set at 15 ppm and 0.05 Da, respectively. Trypsin was specified as the protease. The fixed modification was set as carbamidomethylation (C), and other modifications were set as flexible modification as follows: deamidation (N) and oxidation (M). Full-tryptic end and two missed cleavage sites were allowed. A decoy version of the IPI mouse database was used to estimate peptide and protein false discovery rate. The False Discovery Rate was set at 0.01 to eliminate low-probability protein identifications.

#### 2.6 Protein classification

Signal peptides were predicted using SignalP 4.1 [19]. Transmembrane (TM) helices were predicted using the TMHMM program (version 2.0; CBS prediction servers), which predicts protein topology and the number of TM helices [20]. Information from SignalP and TMHMM were combined to classify proteins into one of the following 6 categories: (i) secreted proteins that contain predicted cleavable signal peptides and no predicted TM segments; (ii) single TM type I proteins that contain predicted single TM helix with an extracellular (or luminal) N-terminus; (iii) single TM type II proteins that contain predicted single TM helix with an extracellular (or luminal) C-terminus; (iv) multiple TM proteins that contain multiple TM helices; or (v) ambiguous proteins [21].

#### 3 Results

#### 3.1 Extracellular matridome of cardiac tissue

To profile the extracellular matridome of post-MI LV tissue, the infarct region was analyzed. Since extracellular proteins are more likely to be glycosylated compared to intracellular proteins [8], the glycopeptide capture method was used to enrich for

extracellular proteins [10, 11]. Of the identified peptides, 95.6% were de-glycosylated and had *N*-linked glycosylation motif (NXS/T, where X is any amino acid except proline). We also determined the extent of spontaneous deamidation on Asn residues in the *N*-glycosylation motif by profiling the un-conjugated fraction of hydrazide beads with or without PNGase F treatment [22]. The rate of spontaneous deamidation was 1.5% in our sample sets. The deamidated peptides identified in this negative control were not included in the identification list (Supplementary Table S1).

A total of 1352 unique *N*-linked glycosylation sites were identified, with a 1% false discovery rate (FDR), and these sites represented 694 unique glycoproteins (Supplemental Table S1). A unique glycosylation site was defined as a peptide containing a unique glycosylation motif, regardless of the length of the peptide and other modification of amino acid other than on asparagine in the motif. As summarized in Figure 1A, 406 proteins (59%) were identified as having one unique glycosite, whereas 288 (41%) were identified as having two or more unique glycosites. The majority of single-glycosites (80%) were identified more than two times (either in multiple identifications of the same peptide sequence with varying charge states or in different peptides). For the single-glycosite peptides only identified once, the annotated MS/MS spectra are provided in Supplemental Table S2. Many ECM proteins were identified with multiple peptides, such as collagens, fibronectin, and laminins (Supplemental Table S2).

The prevalence of identified glycopeptides in eight biological replicates is illustrated in Figure 1B. A total of 907 glycosites (67%) were identified in at least 4 biological replicates, whereas 200 glycosites (15%) were only identified in one of the biological replicates. Notably, 486 glycosites (36%) were identified from all eight samples. This large overlap indicates that our data were highly consistent in biological replicates, implicating good coverage of the *N*-glycoproteome in the mouse LV infarct region.

#### 3.2 Functional analysis and KEGG pathway analysis of identified glycoproteins

To obtain an overview of the molecular functions and associated pathways, the identified glycoproteins were analyzed using David bioinformatics tools (6.7) [23, 24]. With 1% FDR, many functions related to extracellular activities were significantly enriched (p < 0.01), including calcium ion binding, exopeptidase activity, endopeptidase inhibitor activity, peptidase activity, carboxypeptidase activity, ECM structural constituent, metalloexopeptidase activity, and metallocarboxypeptidase activity (Table 1 and Supplemental Table S3). KEGG pathway analysis using DAVID revealed similar results [25]. The *N*-glycoproteome enriched the pathways of ECM-receptor interaction, cell adhesion molecules, arrhythmogenic right ventricular cardiomyopathy, hypertrophic cardiomyopathy, and dilated cardiomyopathy (Table 1 and Supplemental Table S3). These results indicate that the *N*-glycoproteome of the LV infarct region has important myocardial functions, and the identified extracellular proteome contains a large number of cardiac specific glycoproteins.

#### 3.3 New information for protein topology provided by glycosite identification

Based on the information from SignalP and TMHMM, the proteins were classified into one of five categories: secreted, single transmembrane protein (TM) type I, single TM type II, multiple TM, or ambiguous [21]. Of 694 identified proteins, 388 (56%) were TM proteins including single TM type I, type II, and multiple TM (Figure 1C). The identification of Nlinked glycosylation site provided topological information for the transmembrane proteins, since N-linked glycosylation only occurs in extracellular domains of plasma membrane proteins. Because the orientation of single TM protein determines the protein type, this information is important to corroborate the orientation prediction of single TM proteins. Our study identified 153 single TM type I proteins and 107 single TM type II proteins based on the prediction using TMHMM. The single TM type I proteins have extracellular (or luminal) N-terminus, while the single TM type II proteins have extracellular (or luminal) C-terminus. The identified N-glycosites matched with the TMHMM predicted single TM protein type for 252 of the 260 proteins (Supplementary Table S4). The remaining 8 single TM proteins had identified N-glycosites that TMHMM predicted to be cytoplasmic domains (Supplementary Table S4). For these 8 proteins, we checked the topology prediction in Swiss-Prot, and found that our results matched Swiss-Prot prediction (which was of TMHMM prediction) for 7 of the 8 proteins. The final protein was not recorded in Swiss-Prot.

Many integrins were identified in the most enriched KEGG pathway, ECM-receptor interaction. They are the major adhesion receptor for many ECM proteins, and their primary role is to link ECM to the intracellular signaling network [26]. Therefore, it is very important to determine their topology in order to fully understand the signaling interactions. Fourteen integrins are predicted as single transmembrane proteins. Table 2 shows the predicted topology of these integrins, how many N-glycosites are predicted in the extracellular domain, and how many of the predicted N-glycosites identified in this study are in the extracellular domain. The identified glycosylation sites were all in the predicted extracellular domains, supporting the topology prediction. The  $\beta_1$  Integrin is further illustrated in Figure 2. The  $\beta_1$  Integrin has fourteen predicted N-linked glycosites, which are located in the potential extracellular domain (Figure 2). However, out of fourteen predicted N linked glycosites in  $\beta_1$  integrin, only four have supporting experimental evidence according to Swiss-prot (highlighted in green). Our study identified four novel N-glycosites for  $\beta_1$  Integrin (namely, N212, N406, N481, N520, highlighted in yellow) in addition to one of the four N-glycosites recorded in Swiss-prot (Figure 2). These glycosites confirm the extracellular domain prediction of  $\beta_1$  integrin.

#### 4 Discussion

This study employed a glycoproteomic analysis followed by mass spectrometry and a topology bioinformatic assessment to profile the extracellular matridome of the LV infarct region. The most significant findings were: 1) 1352 unique *N*-linked glycosylation sites (representing 694 unique glycoprotein groups) were identified, with the majority (77%) being membrane proteins and secreted proteins; 2) the extracellular proteins identified included collagens, fibronectin, and laminins. These proteins were highly prevalent in the LV infarct; and 3) analysis of *N*-linked glycosites provides experimental evidence of

membrane protein orientation. Out of 260 single TM proteins, the *N*-glycopeptides identified in the present study confirmed 259 proteins for the topology prediction using TMHMM or Swiss-Prot. TMHMM and Swiss-Prot had opposite topology predictions for seven proteins. Combined, these results provide the first glycoproteomic analysis of the extracellular matridom of the LV infarct region and validate the assumption that extracellular proteins are greatly enriched by the glycoprotein isolation approach used. These results also provide useful geographic information on membrane proteins, which will help in mechanistic and drug discovery studies, particularly for the proteins having controversial topology prediction.

Using a glycoproteomic approach to profile the extracellular matridome allows enrichment for extracellular proteins, targeting proteins with important signaling roles. In the functional analysis and pathway annotation of the identified proteins, functions related to extracellular proteins and cardiac signaling pathways were significantly enriched. For example, MMP-9 was identified as a central player, and roles for this metalloproteinase in the post-MI setting have been well documented [7, 27–29].

Using the N-linked glycosylation site analysis provided in vivo topologic evidence that has not been previously available. Zielinska et al demonstrated that glycosylation sites of membrane proteins always orient toward the extracellular space [30]. Gundry and coworkers found one glycoprotein transmembrane orientation was inconsistent with Swiss-Prot annotation using glycoproteomic analysis, providing complementary information to correct the protein orientation prediction [9]. Rossi et al studied the membrane-bound form of complement protein C9 using glycosylation mapping, anti-peptide antibody binding, and disulfide modification analyses [31]. By deleting two N-glycosites and introducing new Nglycosites, the authors determined the glycosylation required for human C9 activity and its membrane anchoring, which shows that the glycosite identification is a useful tool for protein orientation assignment. The current study identified 1352 unique N-linked glycosylation sites, providing new topologic information (Table 2 and Supplemental table S1). For  $\beta_1$  integrin, there were fourteen predicted *N*-linked glycosites, but only four of them (namely, N363, N366, N376, and N669) were identified previously according to Swiss-prot. In addition, these glycosites (N363, N366, and N376) are very close to each other, which is not good for supporting the large region (728 aa) of the extracelluar domain prediction. Our study identified four novel N-glycosites for  $\beta_1$  Integrin (namely N212, N406, N481, and N520) in addition to one of four N-glycosites (N669) recorded in Swiss-prot. These four novel glycosites were also recently identified by other groups [32, 33]. These five sites were distributed throughout the predicted extracellular domain, which provides supporting evidence for this section being the extracellular domain region of  $\beta_1$  integrin.

In the present data, 406 (59%) glycoproteins were identified by one unique glycosite, whereas 288 (41%) were identified by two or more unique glycosites. There are several possible reasons for a single identification. The first reason is that there really is only one potential *N*-glycosite in the protein. Out of the 406 proteins identified with one unique glycosite, 52 (12.8%) had only one single predicted glycosite, indicating that this reason accounted for a minority of the proteins. A second reason is that the peptide length of some glycopeptides may not be suitable for mass spectrometry analysis. Usually, a peptide length

of 7–35 residues is most suitable for mass spectrometry sequencing technology [34]. A third reason is that some peptides may not be suitable for identification due to poor ionization and fragmentation. A fourth reason is the random sampling issue of mass spectrometry, which basically means that false negatives can occur. Our results indicate that all of the above reasons likely contribute to the results. To assess the quality of identification, the single identified glycosites were counted for the number of peptide-spectra matching. We found that the majority of single glycosites (80%) were identified more than two times (either in multiple identification of the same peptide sequence or in different glycopeptides), indicating the majority of the glycopeptide identification was reliable.

In conclusion, this study identified a large number of extracellular proteins in the LV infarct. The information of experimentally identified *N*-glycosites provides *in vivo* experimental evidence for topology prediction. Our data provides the foundation for future studies of the LV infarct extracellular matridome, which may facilitate the discovery of drug targets and biomarkers.

#### Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

#### Acknowledgments

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#### Abbreviations

ECM	extracellular matrix
LV	left ventricle
ГМ	transmembrane
PNGase F	Peptide-N-Glycosidase F

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#### **Clinical relevance**

The extracellular matridome plays a critical role in remodeling of the left ventricle (LV) following myocardial infarction (MI). The aim of this study was to use a glycoproteomics and mass spectrometry approach to identify glycoproteins in the extracellular matridome of the infarcted LV, to provide experimental evidence for topological determination. The information of protein topology is critical for selecting antigen sites and designing drug targets. The biological reproducibility was also investigated to assess the prevalence of the identified *N*-glycosylation sites. The identification of *N*-glycosylation sites supports the prediction of the membrane protein topology. Combined, this information may facilitate the discovery of drug targets and biomarkers for the post-MI patient.



#### Figure 1.

(A) Pie chart showing the distribution of glycosites identified per protein. Of the proteins identified, 59% had 1 glycosite and 41% proteins had 2 glycosites. (B) Stacked column chart showing the prevalence of identified glycopeptides in each of the eight biological replicates. A total of 36% of the glycosites were identified in all eight samples. (C) Classification of identified glycopeptides according to SignalP and TNHMM. A total of 77% of the glycopeptides were classified as transmembrane (TM) or secreted proteins.

	1 <b>0</b>	2 <b>0</b>	30	4 <b>0</b>	5 <b>0</b>	6 <b>0</b>
1	MNLQLVSWIG	LISLICSVFG	QTDKNRCLKA	NAKSCGECIQ	AGPNCGWCTN	TTFLQEGMPT
61	SARCDDLEAL	KKKGCQPSDI	ENPRGSQTIK	KNK <mark>NVTN</mark> RSK	GMAEKLRPED	ITQIQPQQLL
121	LKLRSGEPQK	FTLKFKRAED	YPIDLYYLMD	LSYSMKDDLE	NVKSLGTDLM	NEMRRITSDF
181	RIGFGSFVEK	TVMPYISTTP	AKLRNPCTSE	Q <mark>N</mark> CTSPFSYK	NVLSLTDRGE	FFNELVGQQR
241	ISGNLDSPEG	GFDAIMQVAV	CGSLIGWR <mark>N</mark> V	TRLLVFSTDA	GFHFAGDGKL	GGIVLPNDGQ
301	CHLENNVYTM	SHYYDYPSIA	HLVQKLSENN	IQTIFAVTEE	FQPVYKELKN	LIPKSAVGTL
360	SG <mark>N</mark> SS <mark>N</mark> VIQL	IIDAY <mark>N</mark> SLSS	EVILENSKLP	DGVTINYKSY	CKNGV <mark>N</mark> GTGE	NGRKCSNISI
421	GDEVQFEISI	TANKCPNKES	ETIKIKPLGF	TEEVEVVLQF	ICKCNCQSHG	IPASPKCHEG
481	NGTFECGACR	CNEGRVGRHC	ECSTDEVNSE	DMDAYCRKE <mark>N</mark>	SSEICSNNGE	CVCGQCVCRK
541	RDNTNEIYSG	KFCECDNFNC	DRSNGLICGG	NGVCRCRVCE	CYPNYTGSAC	DCSLDTGPCL
601	ASNGQICNGR	GICECGACKC	TDPKFQGPTC	ETCQTCLGVC	AEHKECVQCR	AFNKGEKKDT
661	CAQECSHF <mark>N</mark> L	TKVESREKLP	QPVQVDPVTH	CKEKDIDDCW	FYFTYSVNGN	NEAIVHVVET
721	PDCPTGPD <mark>II</mark>	PIVAGVVAGI	VLIGLALLLI	WKLLMIIHDR	REFAKFEKEK	MNAKWDTGEN
781	PIYKSAVTTV	VNPKYEGK				
	_					
21-7	28: Extracel	lular				
729-	751: Helical					

752-798: Cytoplasmic

N : Putative *N*-linked glycosites but not validated

N: Previously validated N-linked glycosites (N363, N366, N376, N669)

Novel N-linked glycosites validated in this study (N212, N406, N481, N520)

#### Figure 2.

The glycopeptide identification supports the topology prediction for the  $\beta_1$  integrin, which is predicted to be a TM protein (with residues 21–728 as the extracellular domain, 729–751 as the TM helices, and 752–798 as the intracellular domain). There were four *N*-glycosites previously identified in other studies (highlighted in green). Five *N*-glycosites were identified in this study, and four of the five *N*-glycosites were not previously reported and highlighted in yellow.

#### Table 1

Molecular functions and KEGG pathways enriched for the identified glycoproteins, according to Gene Ontology (GO) analysis. The molecular functions of the identified glycoproteins are ordered by p value, from lowest to highest value.

Molecular functions enriched for the identified glycop	roteins			_
Term	P Value	FDR	Count	%
Calcium ion binding	1.8E-35	2.6E-32	111	16.6
Carbohydrate binding	1.4E-28	2.0E-25	62	9.3
Polysaccharide binding	3.3E-24	4.9E-21	38	5.7
Pattern binding	3.3E-24	4.9E-21	38	5.7
Glycosaminoglycan binding	8.0E-23	1.2E-19	35	5.2
Heparin binding	1.8E-19	2.6E-16	28	4.2
Exopeptidase activity	1.6E-13	2.4E-10	21	3.1
Endopeptidase inhibitor activity	1.1E-11	1.5E-08	28	4.2
Metallopeptidase activity	1.3E-11	1.9E-08	30	4.5
Peptidase inhibitor activity	8.8E-11	1.3E-07	28	4.2
Peptidase activity	1.7E-10	2.5E-07	55	8.2
Integrin binding	2.5E-10	3.6E-07	12	1.8
Peptidase activity	2.8E-10	4.1E-07	56	8.4
Carboxypeptidase activity	1.2E-09	1.7E-06	13	1.9
Extracellular matrix structural constituent	2.6E-09	3.7E-06	12	1.8
Serine-type endopeptidase inhibitor activity	2.7E-09	3.9E-06	21	3.1
Transmembrane receptor protein tyrosine kinase activity	5.1E-09	7.4E-06	15	2.2
Enzyme inhibitor activity	7.4E-09	1.1E-05	30	4.5
Growth factor binding	2.1E-08	3.1E-05	16	2.4
Extracellular matrix binding	9.0E-08	1.3E-04	10	1.5
ion binding	9.2E-08	1.3E-04	191	28.6
Cation binding	9.9E-08	1.4E-04	189	28.3
Metalloexopeptidase activity	1.1E-07	1.6E-04	11	1.6
Metallocarboxypeptidase activity	4.4E-07	6.4E-04	9	1.3
Scavenger receptor activity	6.4E-07	9.2E-04	11	1.6
Metal ion binding	1.1E-06	1.5E-03	183	27.4
Sugar binding	1.4E-06	2.1E-03	22	3.3
Protein complex binding	2.8E-06	4.0E-03	14	2.1
KEGG Pathway enriched for ident	ified glycop	roteins		
Term	P Value	FDR	Count	%
ECM-receptor interaction	1.5E-35	1.8E-32	45	6.7
Lysosome	5.7E-20	6.6E-17	38	5.7
Focal adhesion	2.4E-17	2.8E-14	45	6.7
Cell adhesion molecules	1.1E-16	1.3E-13	39	5.8

Molecular functions enriched for the identified glycop	roteins			
Term	P Value	FDR	Count	%
Complement and coagulation cascades	1.2E-16	1.3E-13	28	4.2
Arrhythmogenic right ventricular cardiomyopathy	1.4E-14	1.7E-11	26	3.9
Hypertrophic cardiomyopathy	3.1E-14	3.6E-11	27	4.0
Hematopoietic cell lineage	2.3E-12	2.6E-09	25	3.7
Dilated cardiomyopathy	2.7E-12	3.1E-09	26	3.9

# Table 2

The topology of the integrins identified in the ECM-receptor interaction pathway. The numbers represent the amino acid position, with the bracketed numbers indicating the number of identified/predicted glycosites.

Tian et al.

			Confirmed to	oology (identified/	predicted glycosit
Accession number	Protein description	cellular location	Inside	Tmhelix	Outside
IPI00126077	Integrin alpha-2	Single TM type I	1152-1178	1129–1151	1-1128 (1/9)
IPI00126090	Integrin alpha-3	Single TM type I	606–642	583-605	1–582 (2/13)
IPI00121334	Integrin alpha-4	Single TM type I	1008-1039	985-1007	1-984 (1/12)
IPI00115976	Integrin alpha-5	Single TM type II	1026-1053	1003-1025	1-1002 (5/15)
IPI00227969	Integrin alpha 6	Single TM type I	1038-1073	1015-1037	1-1014 (3/8)
IPI00345112	Integrin alpha-8	Single TM type I	1034-1062	1011-1033	1-1010 (1/16)
IPI00417168	Integrin alpha-11	Single TM type I	1165-1188	1142–1164	1–1141 (4/16)
IPI00315155	Integrin alpha-IIb	Single TM type I	1013-1033	990-1012	1–989 ( <b>1/5</b> )
IPI00132286	Integrin alpha-L	Single TM type I	1109-1163	1086-1108	1-1085 (2/16)
IPI00120245	Integrin alpha-V	Single TM type I	1013-1044	990-1012	1–989 (5/13)
IPI00132474	Integrin beta-1	Single TM type I	752–798	729–751	1–728 <b>(5/14)</b>
IP100320605	Integrin beta-2	Single TM type I	726-771	703-725	1-702 (2/6)
IPI00266264	Integrin beta-3	Single TM type I	741–787	718–740	1-717 (1/6)
IPI00229516	Integrin beta-5	Single TM type I	743-816	720–742	1–719 (1/7)