# Differential proteomic profiling of primary and recurrent chordomas

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Abstract. Chordomas are locally destructive tumors with high rates of recurrence and a poor prognosis. The mechanisms involved in chordoma recurrence remain largely unknown. In the present study, we examined the proteomic profile of a chordoma primary tumor (CSO) and a recurrent tumor (CSR) through mass spectrum in a chordoma patient who underwent surgery. Bioinformatic analysis of the profile showed that 359 proteins had a significant expression difference and 21 pathways had a striking alteration between the CSO and the CSR. The CSR showed a significant increase in carbohydrate metabolism. Immunohistochemistry (IHC) confirmed that the cancer stem cell marker activated leukocyte cell adhesion molecule (ALCAM or CD166) expression level was higher in the recurrent than that in the primary tumor. The present study analyzed the proteomic profile change between CSO and CSR and identified a new biomarker ALCAM in recurrent chordomas. This finding sheds light on unraveling the pathophysiology of chordoma recurrence and on exploring more effective prognostic biomarkers and targeted therapies against this devastating disease.

### Introduction

Chordoma is a rare slow-growing neoplasm thought to arise from cellular remnants of the notochord. The incidence of chordoma is approximately 1 case in one million individuals and it accounts for  $\sim$ 1-4% of all tumor cases. However, these rare tumors present very significant treatment challenges (1). It

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is notable that 67% of the surgically managed patients suffer local recurrence, and the disease-free survival at 5 years is almost 0% (1,2).

Complete surgical resection followed by radiation therapy offers the best chance of long-term control for chordoma. However, incomplete resection of the primary tumor makes controlling the disease more difficult and increases the odds of recurrence. Tumors at certain sites such as skull base chordoma can hardly be removed completely due to the complicated peritumoral tissue structure which makes the tumor difficult to be exposed. Thus, recurrence of skull base chordoma is as high as 85%. Chordomas are relatively radioresistant as well, requiring high doses of radiation to be controlled. The proximity of chordomas to vital neurological structures such as the brain stem and nerves limits the dose of radiation that can safely be delivered. A strategy to control the recurrence of chordomas is vital for improving the survival rate of patients. It is imperative to explore chordoma recurrence mechanisms at the molecular level and to search for alternative therapeutic methods including chemotherapy to treat it. Recently, Zhou et al assessed the chordoma proteome in chordoma tumor tissues and identified ENO1, PKM2, and gp96 proteins as being upregulated in chordomas. They reported that the expression of these proteins was higher in recurrent than that in the primary chordomas (3). However, the molecular mechanisms involved in chordoma recurrence remain unstudied, and the detailed changes in proteomic profiling in the process of recurrence remain unclear.

We analyzed 3,296 proteins identified by mass spectrum in a chordoma patient original tumor (CSO) and recurrent tumor (CSR) tissue. Bioconductor's Global Anova test was applied to compare the overall proteomic profiling changes between CSO and CSR which indicated there was a significant difference (P≤0.1). Bioconductor's multtest found that 359 proteins exhibited the highest expression difference between CSO and CSR. KEGG database analysis of the 359 proteins revealed that 21 pathways had a significant change between CSO and CSR. Immunohistochemistry (IHC) further verified that cancer stem cell marker activated leukocyte cell adhesion molecule (ALCAM or CD166) was markedly upregulated in the CSR tissue.

#### Materials and methods

Tissue specimen processing. Chordoma specimens were obtained from a resected tumor following an institutional review board approved protocol. The histological composition of the samples was assessed by examining adjacent sections. Tumor samples were dissected and only tissue that was superfluous to that required for pathological evaluation was taken. The samples were immediately snap-frozen in liquid nitrogen and stored at -80°C. Approximately 800  $\mu$ g of tissue samples was cut into small pieces with a scalpel and transferred into a mortar filled with liquid nitrogen. The tissue was ground to a fine powder with a pestle in the continuous presence of liquid nitrogen and transferred into a reaction tube with extraction buffer [2 M thiocarbamide, 7 M urea and 10  $\mu$ M proteinase inhibitor (Roche Diagnostics, Indianapolis, IN, USA)] at 4°C. The solution was centrifuged at 16,000 x g at 4°C for 15 min, and the supernatant ( $\sim$ 300-400  $\mu$ l) was stored frozen at -200°C.

Proteome analysis. The protein concentration was determined by Amersham 2D Quant kit (GE Healthcare Bio-Sciences, Piscataway, NJ, USA). The protein samples were further lysed to peptides and prepared for proteome analysis as described previously (4). Peptides were analyzed using strong cation (SCX)/reversed phase, upgrade performance liquid chromatography (Nano-RPLC)/ESI/MS/MS. Samples were analyzed using a LTQ Orbitrap XL (Thermo Electron Corp., Bremen, Germany) mass spectrometer. MS/MS spectra data were searched against the Swiss-Prot Human (2009.02.10, 20331 sequences) database or IPI database using Bioworks Browser 3.3.1 SP1. The identified proteins were quantified by APEX software (5,6). To control the false-positive rate, finally the quantitative results by false-discovery rate (FDR) 1% or less (false-positive rate of 1% or less) as the standard filter.

Immunohistochemistry. The paraffin sections were dried in an oven at 65°C for 1 h. The paraffin sections were then dewaxed in xylene and rehydrated in a series of ethanol solutions. The endogenous peroxidase activity was blocked by a 10-min pre-incubation with 3% H<sub>2</sub>O<sub>2</sub>. The paraffin sections were preheated at 100°C in antigen retrieval solution containing EDTA (pH 8.0) for 30 min and blocked by non-immune goat serum at room temperature for 15 min to decrease unspecific staining. Incubation with mouse polyclonal anti-ALCAM (1:1,000 dilution) was performed overnight at 4°C. After being washed 3 times with 1X PBS buffer for 3 min, the sections were incubated with the secondary (link) antibody (biotinylated mouse-anti-human IgG) for 30 min at room temperature. After reacting with the streptavidin-biotin-peroxidase complex for 20 min, the immunoreactivity was determined by 3,3'-diamino-benzidine tetrahydrochloride and H<sub>2</sub>O<sub>2</sub> at room temperature according to the manufacturer's instructions. The positive reaction was manifested as brown (DAB) staining. The sections were counterstained in Mayer's hematoxylin. The selected sections were scanned at x400 magnification to visualize the localization and distribution of ALCAM.

Statistical analysis. Statistically significant proteins were identified by first performing a two-tailed Student's t-test with the 'multtest' package in rat the respective time-points

by comparing protein abundance between CSO and CSR. Multiple hypothesis testing was then implemented with the 'rawp2adjp' function in R by correcting the P-values according to Benjamini and Hochberg procedures (7) to control the FDRs to  $\leq 1\%$ . Proteins with a FDR  $\leq 1\%$ , peptide count  $\geq 3$ , and fold-change  $\geq 2$  were identified as statistically significant. Throughout the present study, upregulation is defined as higher protein abundance measured in CSR relative to CSO and downregulation refers to fewer proteins measured with CSR. Moreover, a positive expression ratio represents upregulation and negative represents downregulation. Blast2Go (8) was used as a comprehensive bioinformatics tool for the functional annotation of the protein sequences in the present study such as determining gene ontology terms. The metabolic map at CSR was generated by first using the statistically significant proteins to identify the key pathways. Once the pathways were identified, all of the detected proteins in the same pathway were evaluated to determine whether they were upregulated or downregulated relative to the control. If ≥50% of the proteins in the pathway was regulated similarly in the same direction, then the pathway would be designated as upregulated or downregulated according to the majority. The cellular pathways are displayed using the iPath 2.0 platform (9).

#### **Results and Discussion**

In order to obtain a comprehensive proteomic profile of CSO and CSR and to investigate the mechanism of recurrence, the proteome of patient CSO and CSR tumor tissue samples was analyzed with LC/MS. In total, 3,296 unique protein sequences were identified. Bioconductor's Global Anova package was used to determine the significance of the protein expression change between CSO and CSR. A protein expression difference with P≤0.1±0.002 was defined as statistically significant (Table I). Furthermore, we applied Bioconductor's multtest package to analyze the identified 3,296 proteins and found that a large number of proteins (359) showed significant changes in expression with BH  $\leq$ 0.01 and P $\leq$ 0.01 between CSO and CSR. These proteins were involved in central metabolism, genetic information transcription and other processes essential to cell functions. Of these 359 proteins, there were 244 downregulated proteins (CSR/CSO value ≤0.1) and 115 upregulated proteins (CSR/CSO≥9) (Fig. 1; Table III and IV). This analysis discovered many significant proteins which have never been reported before in recurrent chordomas.

Among the top downregulated proteins, podocan is involved in negative regulation of cell migration and proliferation, concomitant with increased p21 expression which is a tumor-suppressor gene and can mediate cellular senescence (10). ZO-1 has been shown to be downregulated in poorly differentiated, highly invasive breast cancer cell lines (11), and downregulation of complement factor I (CFI) is regarded as a potential suppressive protein for gastric cancer identified by serum proteome analysis (12). Downregulation of osteomodulin (OMD) is referred to in the context of uterine serous papillary carcinoma. It was further disclosed that activation of OMD or/and PRELP gene expression or function can suppress cancer initiation and development (13). FK506-binding protein 4 (FKBP4) was reported to have cancer-specific methylation which usually inactivates this

Table I. Global test for differential gene expression.

ANOVA	SSQ	DF	MS
Effect	0.002073207	3296	6.29007E-07
Error	0.00065985	13184	5.00493E-08
Test Result			
F.value	12.56774798		
p.perm	0.1		
p.approx	0.002139239		

Table II. KEGG database identified 21 pathways with Seqs  $\geq 3$ .

Pathways	#Seqs	#Enzs
Purine metabolism	10	11
Arginine and proline metabolism	5	6
Starch and sucrose metabolism	5	6
Pyruvate metabolism	4	4
Glycolysis/gluconeogenesis	4	4
Oxidative phosphorylation	4	2
Alanine, aspartate and glutamate metabolism	4	5
Cysteine and methionine metabolism	4	5
Aminoacyl-tRNA biosynthesis	3	3
Butanoate metabolism	3	3
Methane metabolism	3	3
Streptomycin biosynthesis	3	3
Amino sugar and nucleotide sugar metabolism	3	5
Propanoate metabolism	3	4
Valine, leucine and isoleucine degradation	3	3
β-alanine metabolism	3	4
Drug metabolism - other enzymes	3	2
Glyoxylate and dicarboxylate metabolism	3	3
Phosphatidylinositol signaling system	3	3
Inositol phosphate metabolism	3	3
Pyrimidine metabolism	3	3

gene in breast cancer tissues (14). We are the first to report downregulation of these tumor-suppressing proteins in recurrent chordomas. The markedly decreased expression of these tumor-suppressing proteins suggests that recurrent chordomas are more aggressive than primary chordomas. While the top upregulated proteins included myosin-7 (MYH7) which is related to eukaryotic cell motility; CD166 (ALCAM) has been regarded as a potential cancer stem cell marker (15). Splicing factor, arginine/serine-rich 2 (SFRS2) demonstrated Wnt signaling-dependent activation which promotes cell migration (16); and Ras-related protein Rab-11A (RAB11A) can differentially modulate epidermal growth factor-induced proliferation and motility in immortal breast cells (17). These newly identified upregulated molecules in recurrent chordomas by our analysis may be possible biomarkers for diagnosis and/ or targets for treating recurrent chordomas. Our study also provides valuable information for future studies on chordoma recurrence mechanisms which remain unelucidated.

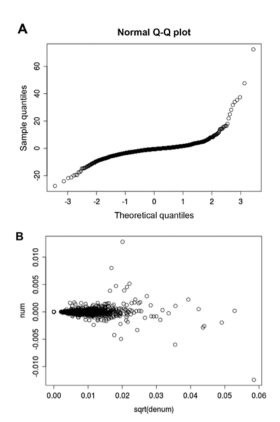


Figure 1. (A) Normal Q-Q plot of t-statistics for the protein data. The points that deviate markedly from an otherwise linear relationship correspond to those genes whose expression levels differ between the CSO and CSR groups. (B) Numerator vs. square root of denominator of the t-statistics for the protein data. The majority of proteins which do not reveal any difference in expression between CSO and CSR groups are represented by points distributed on the y-axis around zero. Some of the points that deviate from this area represent the corresponding proteins whose expression levels are higher or lower in the CSR group than that in the CSO group. CSO, chordoma primary tumor; CSR, recurrent tumor.

To investigate which signaling pathways have alterations in CSR, we searched the 359 identified proteins with significant change in the KEGG database and found that there were over 21 pathways with Seqs ≥3 between CSO and CSR as shown in Table II. Eight pathways were markedly upregulated in CSR compared with CSO (num upregulated protein ≥60%) and nine pathways were apparently downregulated in CSR compared with CSO (num downregulated protein ≥60%) (Fig. 2). Notably, most of the upregulated pathways (6 of 8) and cellular components are involved in carbohydrate metabolism indicating that carbohydrate metabolic activity was higher in recurrent chordomas than in primary chordomas (Fig. 2). Fig. 3 shows that the top 3 upregulated pathways including butanoate, inositol phosphate and glyoxylate and dicarboxylate metabolism are all involved in carbohydrate metabolism. Those 3 pathways were upregulated by 85.7, 75 and 73.3%, respectively, in the CSR compared with the CSO (Fig. 3). The glycolysis/gluconeogenesis pathway was also upregulated (Fig. 2). It has been reported that a high glycolytic rate has advantages for malignant cells (18). High glycolytic activity produces high levels of lactate and H+ ions which are transported outside the cell where they directly promote tumor aggressiveness through invasion and metastasis, two other hallmarks of cancer (19). Additionally, the genes and pathways that upregulate

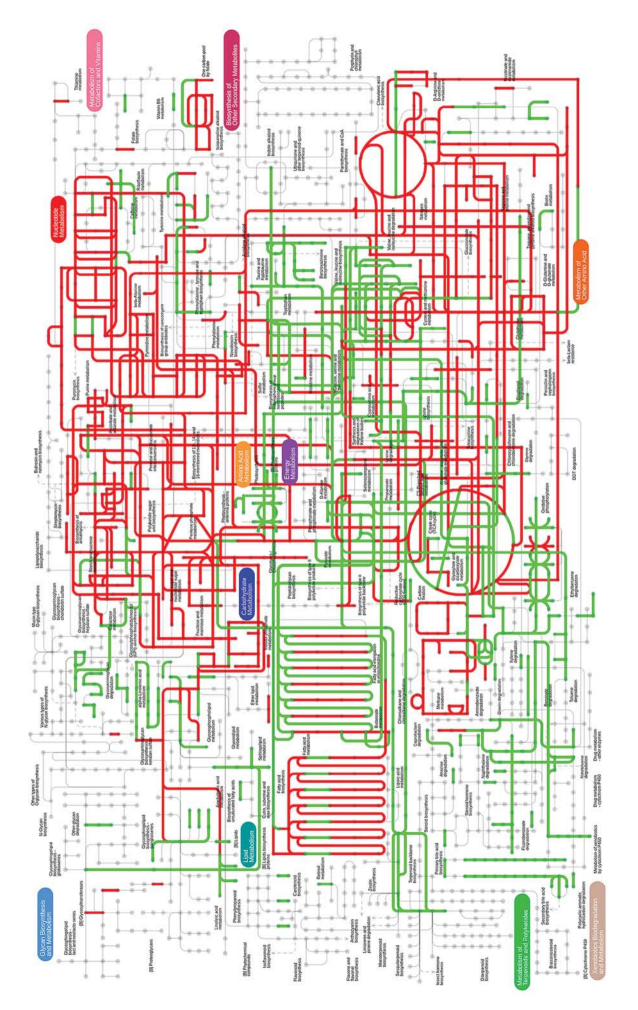
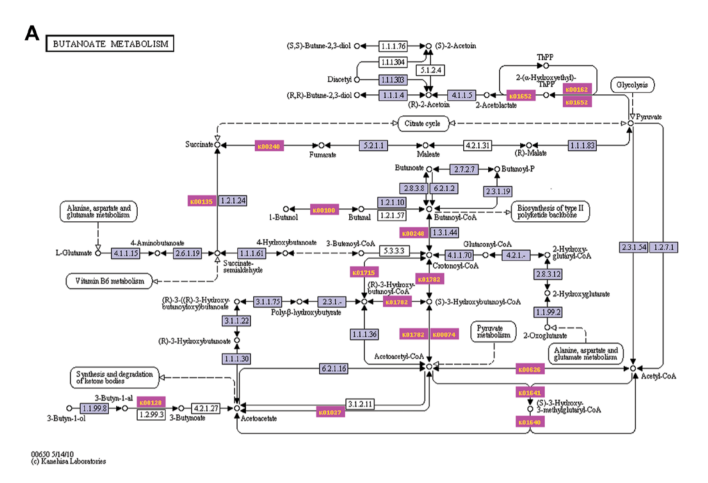


Figure 2. An overview of how the metabolic pathways were changed between CSO and CSR. Red color represents downregulation, green represents upregulation and grey represents no change. Key metabolic pathways are highlighted with bold font. CSO, chordoma primary tumor; CSR, recurrent tumor.



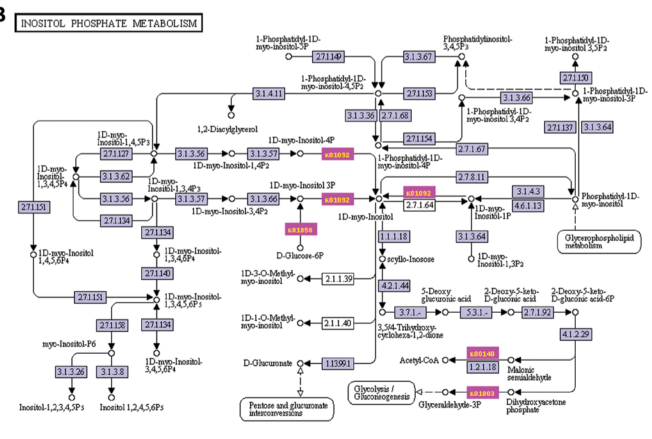


Figure 3. Top upregulated pathways in the KEGG database. (A) Carbohydrate metabolism, butanoatemetabolism pathway. (B) Carbohydrate metabolism, inositol phosphate metabolism.

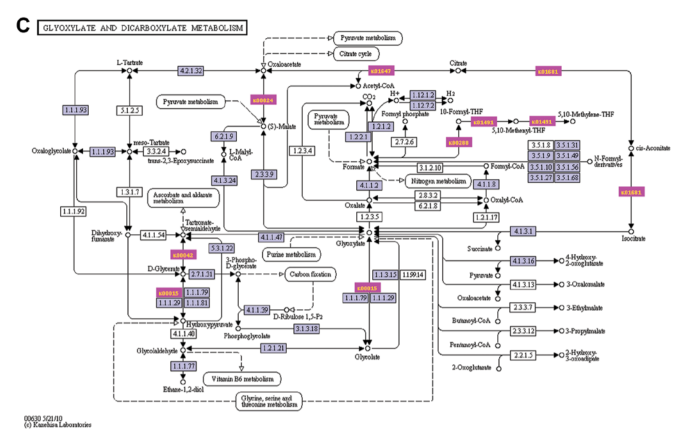
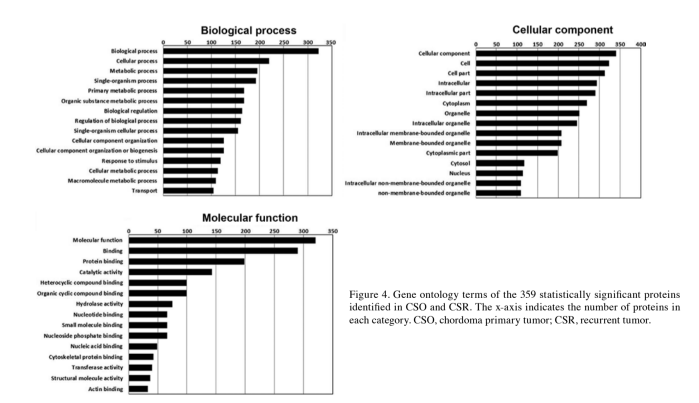


Figure 3. Continued. Top upregulated pathways in the KEGG database. (C) Carbohydrate metabolism, glyoxylate and dicarboxylate metabolism.



glycolysis are themselves anti-apoptotic (20). The increased carbohydrate metabolism in recurrent chordomas suggests that recurrent chordomas have a more aggressive phenotype and are resistant to therapy. The other 2 upregulated pathways in

the recurrent chordomas are involved in energy metabolism and amino acid metabolism. In contrast to the upregulated pathways, the downregulated pathways participate in various biological processes and molecular functions including

Table III. Downregulated proteins (244) (CSR/CSO value ≤0.1). Table III. Continued.

Protein	Protein_description	Protein	Protein_description
Q7Z5L7	Podocan GN=PODN	O94769	Extracellular matrix protein 2 GN=ECM2
Q7L5N1 P31939	COP9 signalosome complex subunit 6 GN=COPS6 Bifunctional purine biosynthesis protein PURH	O94903	Proline synthetase co-transcribed bacterial homolog protein GN=PROSC
	GN=ATIC	O95302	FK506-binding protein 9 GN=FKBP9
O94808	Glucosamine-fructose-6-phosphate aminotransferase	O95373	Importin-7 GN=IPO7
Q8TD55	(isomerizing) 2 GN=GFPT2 Pleckstrin homology domain-containing family O	O95425	Supervillin GN=SVIL
	member 2 GN=PLEKHO2	O95433	Activator of 90 kDa heat shock protein ATPase homolog 1 GN=AHSA1
O75400	Pre-mRNA-processing factor 40 homolog A GN=PRPF40A	O95757 O95810	Heat shock 70 kDa protein 4L GN=HSPA4L Serum deprivation-response protein GN=SDPR
Q6KB66	Keratin, type II cytoskeletal 80 GN=KRT80	O95816	BAG family molecular chaperone regulator 2
Q07157	Tight junction protein ZO-1 GN=TJP1		GN=BAG2
P10155	60 kDa SS-A/Ro ribonucleoprotein GN=TROVE2	O95965	Integrin β-like protein 1 GN=ITGBL1
P05156 Q99983	Complement factor I GN=CFI Osteomodulin GN=OMD	O96005	Cleft lip and palate transmembrane protein 1
Q99983 Q02790	FK506-binding protein 4 GN=FKBP4	D01614	GN=CLPTM1
Q9Y240	C-type lectin domain family 11 member A	P01614 P01781	Ig κ chain V-II region cum Ig heavy chain V-III region GAL
Q 12.0	GN=CLEC11A	P02724	Glycophorin-A GN=GYPA
Q9H8Y8	Golgi reassembly-stacking protein 2	P02750	Leucine-rich α-2-glycoprotein GN=LRG1
	GN=GORASP2	P04433	Ig κ chain V-III region VG (fragment)
P27658	Collagen α-1(VIII) chain GN=COL8A1	P05543	Thyroxine-binding globulin GN=SERPINA7
P27169	Serum paraoxonase/arylesterase 1 GN=PON1	P05546	Heparin cofactor 2 GN=SERPIND1
Q99729	Heterogeneous nuclear ribonucleoprotein A/B GN=HNRNPAB	P07093	Glia-derived nexin GN=SERPINE2
Q9H4A4	Aminopeptidase B GN=RNPEP	P07358	Complement component C8 β chain GN=C8B
P50570	Dynamin-2 GN=DNM2	P08174	Complement decay-accelerating factor GN=CD55
Q14157	Ubiquitin-associated protein 2-like GN=UBAP2L	P08253	72 kDa type IV collagenase GN=MMP2
P02788	Lactotransferrin GN=LTF	P08493	Matrix Gla protein GN=MGP
Q96S97	Myeloid-associated differentiation marker	P08708	40S ribosomal protein S17 GN=RPS17
	GN=MYADM	P10253 P10451	Lysosomal α-glucosidase GN=GAA Osteopontin GN=SPP1
O60841	Eukaryotic translation initiation factor 5B	P10431 P10600	Transforming growth factor β-3 GN=TGFB3
Q96C19	GN=EIF5B EF-hand domain-containing protein D2	P11234	Ras-related protein Ral-B GN=RALB
Q90C19	GN=EFHD2	P12004	Proliferating cell nuclear antigen GN=PCNA
P50225	Sulfotransferase 1A1 GN=SULT1A1	P12107	Collagen α-1(XI) chain GN=COL11A1
A0AVT1	Ubiquitin-like modifier-activating enzyme 6	P15104	Glutamine synthetase GN=GLUL
	GN=UBA6	P19367	Hexokinase-1 GN=HK1
A0MZ66 A1L4H1	Shootin-1 GN=KIAA1598 Scavenger receptor cysteine-rich domain-containing	P20036	HLA class II histocompatibility antigen, DP $\alpha$ chain GN=HLA-DPA1
A8MWD9	protein LOC284297 Small nuclear ribonucleoprotein G-like protein	P20591	Interferon-induced GTP-binding protein Mx1 GN=MX1
O00154	Cytosolic acyl coenzyme A thioester hydrolase	P20851	C4b-binding protein β chain GN=C4BPB
O00461	GN=ACOT7 Golgi integral membrane protein 4 GN=GOLIM4	P22102	Trifunctional purine biosynthetic protein adenosine-3 GN=GART
O14791	Apolipoprotein L1 GN=APOL1	P23193	Transcription elongation factor A protein 1
O43592	Exportin-T GN=XPOT	1 20170	GN=TCEA1
O43670	Zinc finger protein 207 GN=ZNF207	P23497	Nuclear autoantigen Sp-100 GN=SP100
O43847	Nardilysin GN=NRD1	P26373	60S ribosomal protein L13 GN=RPL13
O60240	Perilipin GN=PLIN	P26599	Polypyrimidine tract-binding protein 1 GN=PTBP1
O60684	Importin subunit α-7 GN=KPNA6	P26639	Threonyl-tRNA synthetase, cytoplasmic GN=TARS
O60687	Sushi repeat-containing protein SRPX2 GN=SRPX2	P28300 P31153	Protein-lysine 6-oxidase GN=LOX S-adenosylmethionine synthetase isoform type-2
O60831	PRA1 family protein 2 GN=PRAF2		GN=MAT2A
O75094	Slit homolog 3 protein GN=SLIT3	P32321	Deoxycytidylate deaminase GN=DCTD
O75110	Probable phospholipid-transporting ATPase IIA	P35542	Serum amyloid A-4 protein GN=SAA4
0.55000	GN=ATP9A	P35625	Metalloproteinase inhibitor 3 GN=TIMP3
075339	Cartilage intermediate layer protein 1 GN=CILP	P35858	Insulin-like growth factor-binding protein complex
O75592	Probable E3 ubiquitin-protein ligase MYCBP2 GN=MYCBP2	P36969	acid labile chain GN=IGFALS  Phospholzinid hydroperavide glutathione perovidese
O76021	Ribosomal L1 domain-containing protein 1	1 20404	Phospholzipid hydroperoxide glutathione peroxidase, mitochondrial GN=GPX4

Table III. Continued.

# Table III. Continued.

<ul> <li>P41218 Myeloid cell nuclear differentiation antigen GN=MNDA</li> <li>P41240 Tyrosine-protein kinase CSK GN=CSK</li> <li>P45877 Peptidyl-prolyl cis-trans isomerase C GN=PPIC</li> <li>P46108 Proto-oncogene C-crk GN=CRK</li> <li>P46109 Crk-like protein GN=CRKL</li> <li>P48556 26S proteasome non-ATPase regulatory subunit 8 GN=PSMD8</li> <li>P49321 Nuclear autoantigenic sperm protein GN=NASP</li> <li>P49354 Protein farnesyltransferase/geranylgeranyltransferase type-1 subunit α GN=FNTA</li> <li>P49458 Signal recognition particle 9 kDa protein GN=SRP9</li> <li>P49591 Seryl-tRNA synthetase, cytoplasmic GN=SARS</li> <li>P7050 Raftlin GN=PRPSAP1</li> <li>Q15008 Collectin 1 GN=PRPSAP1</li> <li>Q15008 Collectin 1 GN=RFTN1</li> <li>Q3200 Collectin GN=RC3H1</li> <li>Q406K74 Microtubule-associated protein 1S GN=MAP1S</li> </ul>				
Pal 218   Myeloid cell nuclear differentiation antigen	Protein	Protein_description	Protein	Protein_description
GN=MNDA		<u>*</u>	Q14558	Phosphoribosyl pyrophosphate synthetase-associated
Pal 240   Tymsine-protein kinase CSK GN=CSK   Polico-prospect C-ck GN=CRK   Protein-oncogene C-ck GN=CRK   CRK   Protein-oncogene C-ck GN=CRK   CRK   CRK   Crk GN=CRK   CRK   Protein-oncogene C-ck GN=CRK   CRK   Crk GN=CRK   CRK   Crk GN=CRK   CRK   Crk GN=CRK	P41218		O14699	÷
PASS77 Peptidyl-probly lcis-trans isomerase C GN-PPIC Proto-oncogene C-erk CN-ECRK P46109 Ctk-like protein GN-CRKL Q15181 Longanic pyrophosphatase GN-PPA1 (15405 Sproteasome non ATPase regulatory subunit 8 GN-PSMD8 Nuclear autoantigenic sperm protein GN-ANSP Q15405 Sproteasome non ATPase regulatory subunit 8 GN-PSMD8 Nuclear autoantigenic sperm protein GN-ANSP Q15405 Q15	P41240		•	
P46109 Ch. like protein GN-CRK   Q15121   Astrocytic phosphoprotein PEA 15 GN-PEA15   Q15818   Q15818   Q15818   Q15818   Q15828   Q15907   Q159				
P49556 20S proteasome non ATPase regulatory subunit 8 (Ne-PSMD8 (Ne-PSMD8 (Ne-PSMD8 (Ne-PSMD8 (Ne-PSMD8))) and the part of the		* * * *	-	
GN=PSMD8   Nuclear automitigenic sperm protein GN=NASP     P49321   Potein famesyltransferase/geranyltgeranyltransferase type-1 subunit α GN=FNTA     P49458   Signal recognition particle 9 kDa protein GN=SRP     P49459   Seryl-IRNA synthetase, cytoplasmic GN=SRP     P49591   Seryl-IRNA synthetase, cytoplasmic GN=SRP     P49591   Seryl-IRNA synthetase, cytoplasmic GN=SRP     P59135   Histamine N methyltransferase (M=IRNMT     P59137   P59137   P59137   P59138   Histamine N methyltransferase (N=IRNMT     P59138   P59138   Histamine N methyltransferase (N=IRNMT     P59139   P59138   GN=RDAB     GN=NUD172   P592 and LIM domain protein 4 GN=PDLIM4     ISC -nucleosyl-terraphosphatase (asymmetrical)     GN=NUD172   GN=SRB     GN=NEABC     P51148   Ras-related protein Rab-5C GN=RAB5C     P51812   Ribosomal protein S6 kinase or 3 GN=RPS6KA3     P59539   Septmine synthase GN=SMS     P59539   Septmine synthase GN=GME     GN=DRG     P59530   Septmine synthase GN=GME     GN=DRG     GN=DRG     P59530   Septmine synthase GN=GME     GN=DRG     GN=DRG     P59530   Septmine synthase GN=GME     GN=DRG     GN=DRG     GN=DRG     GN=DRG     GN=DRG     GN=GMF     GN=G	P46109	Crk-like protein GN=CRKL	-	
Pag351   Nuclear autoantigenic sperm protein GN=NASP	P48556		•	
Pay 154 Protein famesyltransferase type-I subunit α GN=PNTA pay 158 type-I subunit α GN=PNTA pay 158 type-I subunit α GN=PNTA pay 159 Seryl-tRNA synthetase, cytoplasmic GN=SARS pay 150 Seryl-tRNA synthetase, c	D40221		•	
type-1 subunif α GN=FNTĀ Signal recognition particle 9 kDa protein GN=SRP9 P49458 Signal recognition particle 9 kDa protein GN=SRP9 P49591 Sey1-lRNA synthetase, cytoplasmic GN=SRP5 P50133 Histamine N methyltransferase GN=HNMT P50833 Bis (5-nucleosyl)-tetraphosphatase (asymmetrical) GN=MVIDT2 P51148 Ras-related protein Rab-5C GN=RAB5C GN=DRG2 SP51812 Ribosomal protein 4 GN=PBLIM4 P55182 Ribosomal protein 56 kinase α-3 GN=RPS6KA3 P55193 Developmentally-regulated GTP-binding protein 2 GN=DRG2 SP5196 Addin GN=MLLT4 SP5197 Addin GN=MLLT4 SP5197 Addin GN=MLLT4 SP5197 Addin GN=MLLT4 SP5197 Addin GN=MLLT4 SP5198 Addin GN=MLLT4 SP5198 Addin GN=MLLT4 SP5199 Addin GN=MLD14 SP5190 Addin GN=MLD14 SP5190 Addin GN=MLD15 AGN=SPB11 SP6122 Ras-related protein Rap-2b GN=RAP2B SP6123 Ras-related protein Rap-2b GN=RAP2B SP6133 GN=SPB07 SP6137 Pefoldin subunit 3 GN=VBP1 SP6137 Pefoldin subunit 3 GN=VBP1 SP6138 Pefoldin subunit 3 GN=VBP1 SP6139 Okribosomal protein L15 GN=RPS11 SP6227 AdS ribosomal protein S13 GN=RPS13 SP6230 40S ribosomal protein S13 GN=RPS13 SP6230 40S ribosomal protein S13 GN=RPS13 SP6230 40S ribosomal protein S14 GN=RPS11 SP62304 Small nuclear ribonucleoprotein GN=SNRPD2 SP62304 Small nuclear ribonucleoprotein GN=SNRPD2 SP62304 Small nuclear ribonucleoprotein GN=SNRP13 SP62304 Small nuclear ribonucleoprotein SP2 GN=SNRP2 SP62305 Small nuclear ribonucleoprotein SP2 GN=SNRP2 SP			-	
P49491   Sejnal recognition particle 9 kDa protein GN=SRP9   P49591   Seryl-tRNA synthetase, cytoplasmic GN=SARS   Q66K74   Microtubule-associated protein 1S GN=MAP1S   Microtubule-associated protein 1S GN=MAP1S   Microtubule-associated protein 1S GN=MAP1S   Microtubule-associated protein 1S GN=MAP1S   Ankyrin repeat and SOCS box-containing protein 1 GN=SRI   Microtubule-associated protein 1S GN=MAP1S   Ankyrin repeat and SOCS box-containing protein 1 GN=SRI   Microtubule-associated protein 1S GN=MAP1S   Ankyrin repeat and SOCS box-containing protein 1 GN=SRI   Microtubule-associated protein 1S GN=MAPIS   Ankyrin repeat and SOCS box-containing protein 2 GN=SRI   Microtubule-associated protein 1S GN=MADC1   Microtubule-associated protein 1S GN=MAPIS   Ankyrin repeat and SOCS box-containing protein 2 GN=ASBI   Microtubule-associated protein 1S GN=AMDC1   Microtubule-associated protein 1S GN=AMDIS   Ankyrin repeat and SOCS box-containing protein 2 GN=ASBI   Microtubule-associated protein 1S GN=AMDIS   Ankyrin repeat and SOCS box-containing protein 2 GN=ASBI   Microtubule-associated protein 1S GN=ASBI   Mi	P49334		-	-
P49591   Seyl-IRNA synthetase, cytoplasmic GN=SARS   Q66K74   Microtubule, associated protein IS GN=MAPIS   GN=MAPIS   GN=VSB   Histamine N-methyltransferase GN=HNMT   Q60Z8   Ankyrin repeat and SOCS box-containing protein I GN=ASBI	P49458	* *	-	
PS0135   Histamine N-methyltransferase GN-HNMT   PD2 and LIM domain protein 4 GN=PDLIM4   GN=NDID12   GN=SETX   GN=CASP6   GN=CASP6   GN=CASP6   GN=CASP6   GN=CASP6   GN=CASP6   GN=GMFB   G			Q66K74	Microtubule-associated protein 1S GN=MAP1S
PSOSS3   Bis (5-nucleosyl)-tetraphosphatase (asymmetrical)   GN=NUDT2	P50135		Q6ZVZ8	Ankyrin repeat and SOCS box-containing protein 18
Probable helicase senataxin GN=SETX   Q86UB   Serine/threonine-protein kinase tousled-like 2   GN=TLK2   GN=DRC2	P50479	PDZ and LIM domain protein 4 GN=PDLIM4	0.5520.4	
P51812   Ribosomal protein S6 kinase α-3 GN=RPS6KA3   Spermine synthase GN=SMS   Q86W92   Liprin-β-1 GN=PPFIBP1   GN=PPFIBP1   GN=PPFIBP1   Histone-arginine methyltransferase CARM1   GN=CARM1   GN=GARM1   GN=CARM1   GN=CARM1   GN=CARM1   GN=CARM1   GN=CARM1   G	P50583		Q7Z333	Probable helicase senataxin GN=SETX
PS1981   Ribosomal protein So kinase α-3 GN=RP50A-3	P51148	Ras-related protein Rab-5C GN=RAB5C	Q86UE8	
P55039   Developmentally-regulated GTP-binding protein 2 GN=DRG2   P55196   Afadin GN=MLLT4   Q8IWE2   Q8IWU6   Caspase-6 GN=CASP6   Q8IWE2   GN=DRG2   Q8IWE2   GN=DRG2   Q8IWE2   GN=DRG2   Q8IWE2   GN=DRG2   Q8IWE2   GN=DRG2   Q8IWE2   GN=DRG2   Q8IWE3   GN=DRG2   Q8IWE2   GN=DRG2   Q8IWE3   GN=DRG2   GN=DRG2   Q8IWE3   GN=DRG2   GN=D		<u>-</u>	086W02	
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P55212 Caspase-6 GN=CASP6 P60983 Glia maturation factor β GN=GMFB P61221 ATP-binding cassette sub-family E member 1 CN=ABCE1 P61225 Ras-related protein Rap-2b GN=RAP2B P61313 60S ribosomal protein L15 GN=RPL15 P6178 Prefoldin subunit 3 GN=VBP1 P6179 Nuclear transport factor 2 GN=NUTF2 P62195 26S protease regulatory subunit 8 GN=PSMC5 P62267 40S ribosomal protein S13 GN=RPS13 P62277 40S ribosomal protein S13 GN=RPS13 P62304 Small nuclear ribonucleoprotein E GN=SNRPE P62316 Small nuclear ribonucleoprotein E GN=SNRPE P6236 Small nuclear ribonucleoprotein E GN=SNRPE P62874 40S ribosomal protein S28 GN=RPS28 P62879 60S ribosomal protein S28 GN=RPS28 P6289 60S ribosomal protein L3 GN=RPL31 P80217 Interferon-induced 35 kDa protein GN=IF135 P80303 Nucleobindin-2 GN=NUCB2 P82987 ADAMTS-like protein 3 GN=ADAMTSL3 P80318 Antigen peptide transporter 1 GN=TAP1 Q00341 Vigilin GN=MLDLBP Q03123 Protein Rap Order SQN=SNRPD Q01246 1,4-α-glucan-branching enzyme GN=GBE1 Q04446 1,4-α-glucan-branching enzyme GN=GBE1		GN=DRG2		GN=CARM1
Posopas of lian maturation factor β GN=GMFB Pologas of lian maturation factor β GN=CMFB Pologas of Sribosomal protein L15 GN=RPL31 Pologas of Sprotease regulatory subunit 8 GN=PSMC5 Pologas of Sribosomal protein S23 GN=RPS23 Pologas of Sribosomal protein S23 GN=RPS13 Pologas of Sribosomal protein S23 GN=RPS13 Pologas of Sribosomal protein S24 GN=RPS24 Pologas of Sribosomal protein S28 GN			-	
Floops   Gilla maturation factor β GN=CMPB		•	-	
GN=ABCEI  P61225 Ras-related protein Rap-2b GN=RAP2B P61313 GOS ribosomal protein L15 GN=RPL15 P61758 Prefoldin subunit 3 GN=VBP1 P61758 Prefoldin subunit 3 GN=VBP1 P61759 Nuclear transport factor 2 GN=RUTF2 P6170 Nuclear transport factor 2 GN=RUTF2 P62195 26S protease regulatory subunit 8 GN=PSMC5 P62260 40S ribosomal protein S23 GN=RPS23 P62277 40S ribosomal protein S13 GN=RPS13 P62280 40S ribosomal protein S11 GN=RPS11 P62304 Small nuclear ribonucleoprotein GN=SNRPE P62304 Small nuclear ribonucleoprotein SmD2GN=SNRPD2 P62370 60S ribosomal protein L23a GN=RPL23A P62287 40S ribosomal protein S24 GN=RPS24 P62874 40S ribosomal protein S24 GN=RPS24 P62875 40S ribosomal protein S24 GN=RPS24 P62876 40S ribosomal protein S24 GN=RPS24 P62877 40S ribosomal protein S24 GN=RPS24 P62879 60S ribosomal protein S24 GN=RPS24 P62879 60S ribosomal protein S24 GN=RPS24 P62899 60S ribosomal protein S24 GN=RPS24 P80217 Interferon-induced 35 kDa protein GN=IF135 P80303 Nucleobindin-2 GN=NUCB2 P80303 Nucleobindin-2 GN=NUCB2 P80304 ADAMTS-like protein 3 GN=ADAMTSL3 P80305 Nucleobindin-2 GN=NUCB2 P82987 ADAMTS-like protein 3 GN=ADAMTSL3 P03518 Antigen peptide transporter 1 GN=TAP1 Q0341 Tyrosine-protein phosphatase non-receptor type 11 GN=PTPN11 Q0361 Tyrosine-protein phosphatase non-receptor type 11 GN=PTPN11 Q03818 Antigen peptide transporter 1 GN=CHBP Q6681 Q966R1 Q79682 Q9684D Q79683 Q9684D Q79684 Q9684D Q79685 Q9684D Q79685 Q96864 Q9687D Q796865 Q96867D Q796867 Q9687D Q79687D Q79		•		
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P80303Nucleobindin-2 GN=NUCB2GN=CTHRC1P82987ADAMTS-like protein 3 GN=ADAMTSL3Q96CV9Optineurin GN=OPTNP83110Probable serine protease HTRA3 GN=HTRA3Q96FW1Ubiquitin thioesterase OTUB1 GN=OTUB1Q00341Vigilin GN=HDLBPQ96GS4Uncharacterized protein C17orf59 GN=C17orf59Q03518Antigen peptide transporter 1 GN=TAP1Q96HF1Secreted frizzled-related protein 2 GN=SFRP2Q044461,4-α-glucan-branching enzyme GN=GBE1Q96HN2Putative adenosylhomocysteinase 3 GN=AHCYL2Q06124Tyrosine-protein phosphatase non-receptor type 11 GN=PTPN11Q96JQ2Calmin GN=CLMNQ08J23tRNA (cytosine-5-)-methyltransferase NSUN2 GN=NSUN2Q96MM6Heat shock 70 kDa protein 12B GN=HSPA12BQ12965Myosin-Ie GN=MYO1EQ96PX9Pleckstrin homology domain-containing protein 7 GN=MBOAT7Q13123Protein Red GN=IKQ96PX9Pleckstrin homology domain-containing family G member 4B GN=PLEKHG4BQ13315Serine-protein kinase ATM GN=ATMQ96RF0Sorting nexin-18 GN=SNX18Q13838Spliceosome RNA helicase BAT1 GN=BAT1Q96RL7Vacuolar protein sorting-associated protein 13A			-	÷
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Q12965 Myosin-Ie GN=MYO1E domain-containing protein 7 GN=MBOAT7 Q13123 Protein Red GN=IK Q13315 Serine-protein kinase ATM GN=ATM Q13838 Spliceosome RNA helicase BAT1 GN=BAT1 Q14011 Cold inducible RNA hinding protein GN=CIPRP Q96RF0 Q96RL7 Vacuolar protein sorting-associated protein 13A	Q08J23	tRNA (cytosine-5-)-methyltransferase NSUN2	-	
Q13123 Protein Red GN=IK Q13315 Serine-protein kinase ATM GN=ATM Q13838 Spliceosome RNA helicase BAT1 GN=BAT1 Q14011 Cold inducible RNA hinding protein GN=CIPRP Q96PX9 Pleckstrin homology domain-containing family G member 4B GN=PLEKHG4B Sorting nexin-18 GN=SNX18 Q96RF0 Q96RL7 Vacuolar protein sorting-associated protein 13A	Q12965			domain-containing protein 7 GN=MBOAT7
Q13315 Serine-protein kinase ATM GN=ATM Q13838 Spliceosome RNA helicase BAT1 GN=BAT1 Q14011 Cold inducible RNA hinding protein GN=CIPRP Q96RF0 Q96RL7 Wacuolar protein sorting-associated protein 13A	-		Q96PX9	
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	Q13838		-	· ·
	Q14011	Cold-inducible RNA-binding protein GN=CIRBP	Q90KL/	

Table III. Continued.

# Table III. Continued.

D4.:			
Protein	Protein_description	Protein	Protein_description
Q99426	Tubulin folding cofactor B GN=TBCB	Q9Y5X3	Sorting nexin-5 GN=SNX5
-	Legumain GN=LGMN	Q9Y6K5	2'-5'-Oligoadenylate synthetase 3 GN=OAS3
-	Protein C10 GN=C12orf57	Q9Y6R7	IgGFc-binding protein GN=FCGBP
•	COP9 signalosome complex subunit 8 GN=COPS8	O43765	Small glutamine-rich tetratricopeptide
-	Vacuolar protein-sorting-associated protein 25	043703	• • • • • • • • • • • • • • • • • • • •
	GN=VPS25	060740	repeat-containing protein α GN=SGTA
		O60749	Sorting nexin-2 GN=SNX2
	3-hydroxybutyrate dehydrogenase type 2 GN=BDH2	Q12765	Secernin-1 GN=SCRN1
	Dual specificity protein phosphatase 23 GN=DUSP23	Q8N0U8	Vitamin K epoxide reductase complex subunit 1-lik protein 1 GN=VKORC1L1
	Complement C1q tumor necrosis factor-related protein 5 GN=C1QTNF5	Q9NYL4	FK506-binding protein 11 GN=FKBP11
	Arsenite-resistance protein 2 GN=ARS2		
-	AP-1 complex subunit mu-1 GN=AP1M1	T.1.1. IV	II
	Inosine triphosphate pyrophosphatase GN=ITPA	Table IV.	Upregulated proteins (115).
~	Ester hydrolase C11orf54 GN=C11orf54		
-	TRIO and F-actin-binding protein GN=TRIOBP	Protein	Protein_description
	GDP-fucose protein O-fucosyltransferase 1 GN=POFUT1	O00148	ATP-dependent RNA helicase DDX39 GN=DDX39
	UPF0554 protein C2orf43 GN=C2orf43	O00330	Pyruvate dehydrogenase protein X component,
-	•	000550	mitochondrial GN=PDHX
-	Phosphopantothenate-cysteine ligase GN=PPCS	O00629	Importin subunit α-4 GN=KPNA4
	Retinoid-inducible serine carboxypeptidase	O00029	Carboxylesterase 2 GN=CES2
	GN=SCPEP1	O14958	Calsequestrin-2 GN=CASQ2
	Progressive ankylosis protein homolog GN=ANKH		•
	Nitrilase homolog 2 GN=NIT2	O43676	NADH dehydrogenase (ubiquinone) 1 β subcomplex subunit 3 GN=NDUFB3
Q9NRN5	Olfactomedin-like protein 3 GN=OLFML3	060794	
Q9NS15	Latent-transforming growth factor β-binding protein 3	O60784	Target of Myb protein 1 GN=TOM1
	GN=LTBP3	O75112	LIM domain-binding protein 3 GN=LDB3
Q9NZL9	Methionine adenosyltransferase 2 subunit β	O75128	Protein cordon-bleu GN=COBL
	GN=MAT2B	O75298	Reticulon-2 GN=RTN2
-	Protein RCC2 GN=RCC2	O75306	NADH dehydrogenase (ubiquinone) iron-sulfur
Q9UBB6	Neurochondrin GN=NCDN		protein 2, mitochondrial GN=NDUFS2
Q9UBR2	Cathepsin Z GN=CTSZ	O94826	Mitochondrial import receptor subunit TOM70
Q9UBW8	COP9 signalosome complex subunit 7α GN=COPS7A	004006	GN=TOMM70A
Q9UDY2	Tight junction protein ZO-2 GN=TJP2	O94906	Pre-mRNA-processing factor 6 GN=PRPF6
	γ-adducin GN=ADD3	O94925	Glutaminase kidney isoform, mitochondrial GN=GL
	Dipeptidyl-peptidase 2 GN=DPP7	O95248	Myotubularin-related protein 5 GN=SBF1
	Enolase-phosphatase E1 GN=ENOPH1	O95299	NADH dehydrogenase (ubiquinone) 1 α
	SH3 domain-binding glutamic acid-rich-like protein 2		subcomplex subunit 10, mitochondrial GN=NDUFA1
-	GN=SH3BGRL2	P02585	Troponin C, skeletal muscle GN=TNNC2
	Angiopoietin-related protein 2 GN=ANGPTL2	P05166	Propionyl-CoA carboxylase β chain,
-	• •		mitochondrial GN=PCCB
	Hippocalcin-like protein 4 GN=HPCAL4	P06732	Creatine kinase M-type GN=CKM
	Neurogenic locus notch homolog protein 3	P07451	Carbonic anhydrase 3 GN=CA3
	GN=NOTCH3	P08590	Myosin light chain 3 GN=MYL3
-	Myosin-VI GN=MYO6	P10916	Myosin regulatory light chain 2, ventricular/cardiac
	NFU1 iron-sulfur cluster scaffold homolog, mitochondrial GN=NFU1		muscle isoform GN=MYL2
Q9UNF0	Protein kinase C and casein kinase substrate in neurons	P11217	Glycogen phosphorylase, muscle form GN=PYGM
	protein 2 GN=PACSIN2	P11233	Ras-related protein Ral-A GN=RALA
	Sorting nexin-7 GN=SNX7	P13805	Troponin T, slow skeletal muscle GN=TNNT1
_	Serine/threonine-protein phosphatase 6 regulatory	P13807	Glycogen (starch) synthase, muscle GN=GYS1
	subunit 1 GN=SAPS1	P14649	Myosin light chain 6B GN=MYL6B
	Nuclear migration protein nudC GN=NUDC	P19237	Troponin I, slow skeletal muscle GN=TNNI1
	Integral membrane protein 2B GN=ITM2B	P23327	Sarcoplasmic reticulum histidine-rich
O0V287	Peptidyl-prolyl <i>cis-trans</i> isomerase-like 1 GN=PPIL1		calcium-binding protein GN=HRC
	repugyi-proivi cis-irans isomerase-like I GN=PPILI	P28289	Tropomodulin-1 GN=TMOD1
Q9Y3C6			
Q9Y3C6 Q9Y4E8	Ubiquitin carboxyl-terminal hydrolase 15 GN=USP15	P29218	
Q9Y3C6 Q9Y4E8 Q9Y5K8	Ubiquitin carboxyl-terminal hydrolase 15 GN=USP15 V-type proton ATPase subunit D GN=ATP6V1D	P29218 P30038	Inositol monophosphatase GN=IMPA1
Q9Y3C6 Q9Y4E8 Q9Y5K8 Q9Y5U9	Ubiquitin carboxyl-terminal hydrolase 15 GN=USP15	P29218 P30038	

Q96EY8

Q9BQS8

Cob(I)yrinic acid a,c-diamide adenosyltransferase,

FYVE and coiled-coil domain-containing protein 1

mitochondrial GN=MMAB

GN=FYCO1

Table IV. Continued.

Table IV. Continued.

Protein	Protein_description	Protein	Protein_description	
P35080 P35609	Profilin-2 GN=PFN2 α-actinin-2 GN=ACTN2	Q9BWD1	Acetyl-CoA acetyltransferase, cytosolic GN=ACAT2	
P42704	Leucine-rich PPR motif-containing protein, mitochondrial GN=LRPPRC	Q9GZV1	Ankyrin repeat domain-containing protein 2 GN=ANKRD2	
P45378	Troponin T, fast skeletal muscle GN=TNNT3	Q9HC07	Transmembrane protein 165 GN=TMEM165	
P48788	Troponin I, fast skeletal muscle GN=TNNI2	Q9NP98	Myozenin-1 GN=MYOZ1	
P50461	Cysteine and glycine-rich protein 3 GN=CSRP3	Q9NPC6	Myozenin-2 GN=MYOZ2	
P51553	Isocitrate dehydrogenase (NAD) subunit γ, mitochondrial GN=IDH3G	Q9NTI5	Sister chromatid cohesion protein PDS5 homolog FGN=PDS5B	
P52179	Myomesin-1 GN=MYOM1	Q9NZQ9	Tropomodulin-4 GN=TMOD4	
P54296	Myomesin-2 GN=MYOM2	Q9UBF9	Myotilin GN=MYOT	
P63316	Troponin C, slow skeletal and	Q9UKS6	Protein kinase C and casein kinase substrate	
000073	cardiac muscles GN=TNNC1	003/225	in neurons protein 3 GN=PACSIN3	
Q00872 Q02045	Myosin-binding protein C, slow-type GN=MYBPC1 Myosin light chain 5 GN=MYL5	Q9Y235	Probable C-)U-editing enzyme APOBEC-2 GN=APOBEC2	
Q09013	Myotonin-protein kinase GN=DMPK	Q9Y2J8	Protein-arginine deiminase type-2 GN=PADI2	
Q10589	Bone marrow stromal antigen 2 GN=BST2	Q9Y639	Neuroplastin GN=NPTN	
Q13061	Triadin GN=TRDN	P12883	Myosin-7 GN=MYH7	
Q14118	Dystroglycan GN=DAG1	P31415	Calsequestrin-1 GN=CASQ1	
Q14324	Myosin-binding protein C, fast-type GN=MYBPC2	P20929	Nebulin GN=NEB	
Q15111	Inactive phospholipase C-like protein 1 GN=PLCL1	Q8WZ42	Titin GN=TTN	
Q16630	Cleavage and polyadenylation specificity factor subunit 6 GN=CPSF6	P05976	Myosin light chain 1, skeletal muscle isoform GN=MYL1	
Q16775	Hydroxyacylglutathione hydrolase GN=HAGH	P02144	Myoglobin GN=MB	
Q5BKX8	PTRF/SDPR family protein	P11532	Dystrophin GN=DMD	
Q5T1J5	Coiled-coil-helix-coiled-coil-helix	Q14315	Filamin-C GN=FLNC	
	domain-containing protein 9,	Q9UHQ9	NADH-cytochrome b5 reductase 1 GN=CYB5R1	
0.51.7777.5	mitochondrial GN=CHCHD9	Q9NZ01	Synaptic glycoprotein SC2 GN=GPSN2	
Q5VTT5	Myomesin-3 GN=MYOM3	Q13740	CD166 antigen GN=ALCAM	
Q5VXT5 Q5W0V3	Synaptophysin-like protein 2 GN=SYPL2 UPF0518 protein FAM160B1 GN=FAM160B1	O95817	BAG family molecular chaperone regulator 3 GN=BAG3	
Q6ZMU5	Tripartite motif-containing protein 72 GN=TRIM72	P25786	Proteasome subunit α type-1 GN=PSMA1	
Q702N8	Xin actin-binding repeat-containing protein 1	Q01130	Splicing factor, arginine/serine-rich 2 GN=SFRS2	
00677704	GN=XIRP1	P12235	ADP/ATP translocase 1 GN=SLC25A4	
Q86TD4	Sarcalumenin GN=SRL	P13929	β-enolase GN=ENO3	
Q86UW8	Hyaluronan and proteoglycan link protein 4 GN=HAPLN4	O75923	Dysferlin GN=DYSF	
Q86VU5	Catechol-O-methyltransferase	P53634	Dipeptidyl-peptidase 1 GN=CTSC	
Q00 V 03	domain-containing protein 1 GN=COMTD1	P23258 O75746	Tubulin γ-1 chain GN=TUBG1	
Q8IWX7	Protein unc-45 homolog B GN=UNC45B	073740	Calcium-binding mitochondrial carrier protein Aralar1 GN=SLC25A12	
Q8IZL8	Proline-, glutamic acid- and leucine-rich protein 1 GN=PELP1	O94919	Endonuclease domain-containing 1 protein GN=ENDOD1	
Q8N1G4	Leucine-rich repeat-containing protein 47	P24043	Laminin subunit α-2 GN=LAMA2	
OONEOC	GN=LRRC47	P11216	Glycogen phosphorylase, brain form GN=PYGB	
Q8NE86	Coiled-coil domain-containing protein 109A GN=CCDC109A	P12829	Myosin light chain 4 GN=MYL4	
Q8NF37	1-acylglycerophosphocholine O-acyltransferase 1 GN=LPCAT1	P55042 P62491	GTP-binding protein RAD GN=RRAD Ras-related protein Rab-11A GN=RAB11A	
Q8NFW1	Collagen α-1(XXII) chain GN=COL22A1	Q14BN4	Sarcolemmal membrane-associated protein	
Q8NI60	Chaperone activity of bc1 complex-like, mitochondrial GN=CABC1	Q96JG9	GN=SLMAP Zinc finger protein 469 GN=ZNF469	
Q8WW22	DnaJ homolog subfamily A member 4 GN=DNAJA4			
Q92629	δ-sarcoglycan GN=SGCD			
Q92029 Q96A32	Myosin regulatory light chain 2, skeletal		metabolism, amino acid metabolism, carbohydrate	
<b>√</b> 70/132	muscle isoform GN=MYLPF		n, genetic information processing and translation	
O06EV8	Coh(I)vrinia acid a a diamida adanaevitranefareca	and biosyn	thesis of other secondary metabolites.	

To further determine the cellular function change between CSO and CRO, the statistically significant 359 proteins identified were classified according to gene ontology

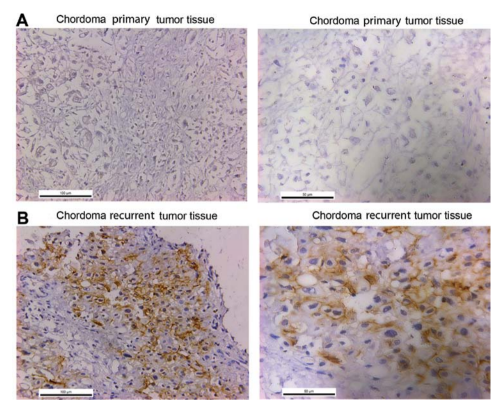


Figure 5. Immunohistochemistry of ALCAM in CSO and CSR. CSO, chordoma primary tumor; CSR, recurrent tumor.

terms. The proteins were found to be involved in either a biological process, a cellular component and/or a molecular function (Fig. 4), indicating that there were diverse cellular components and catalytic activity change in recurrent chordomas compared to the original chordoma. Furthermore, when the identified proteins in the recurrent chordoma were mapped to the corresponding metabolic pathways, many key cellular pathways including amino acid, carbohydrate metabolism, energy, and nucleotide metabolism were found to be downregulated (Fig. 4).

In order to confirm our results from the proteome, we examined tumor-related protein expression from the list of upregulated proteins through IHC in the chordoma patient primary and CSR tissues. ALCAM (or CD166) is a 100-105 kDa type I transmembrane glycoprotein that is a member of the immunoglobulin superfamily of proteins (21). ALCAM has been reported as a cancer stem cell marker for non-small cell lung cancer (NSCLC) (15). Physiologically, it plays a role in the development of different tissues during embryogenesis. It is also expressed in various malignant lesions such as melanoma and esophageal, gynecologic, prostate, and pancreatic cancers, and its expression is associated with diverse outcomes in different tumors (22-32). But the expression of ALCAM in chordomas has never been reported and the association between ALCAM and chordoma prognosis is not fully elucidated. In our study, we firstly detected ALCAM expression by using IHC in chordoma patient primary and CSR tissues. Fig. 5A clearly demonstrates that the primary chordoma tumor was negative for staining; however, the CSR had strong expression of ALCAM which suggests that ALCAM is a positive biomarker for recurrent chordomas and may play important roles for chordoma recurrence (Fig. 5B).

In conclusion, we analyzed the proteomic profile of a chordoma patient CSO and CSR and identified 359 proteins and 21 pathways with significant changes between CSO and CSR. Many of these molecular changes are reported in chordomas for the first time. Further investigation of the potential roles of these proteins in chordoma aggression is of interest. We also firstly found that the recurrent chordoma tumor showed enhanced carbohydrate metabolism, and the cancer stem cell marker ALCAM (CD166) expression level was increased markedly in CSR. The present study can serve as the basis for further research of recurrent chordomas.

## Acknowledgements

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## References

- 1. Stacchiotti S, Casali PG, Lo Vullo S, *et al*: Chordoma of the mobile spine and sacrum: a retrospective analysis of a series of patients surgically treated at two referral centers. Ann Surg Oncol 17: 211-219, 2010.
- DeLaney TF, Liebsch NJ, Pedlow FX, et al: Phase II study of high-dose photon/proton radiotherapy in the management of spine sarcomas. Int J Radiat Oncol Biol Phys 74: 732-739, 2009.
- 3. Zhou H, Chen CB, Lan J, *et al*: Differential proteomic profiling of chordomas and analysis of prognostic factors. J Surg Oncol 102: 720-727, 2010.
- Wisniewski JR, Zougman A, Nagaraj N and Mann M: Universal sample preparation method for proteome analysis. Nat Methods 6: 359-362, 2009.

- Vogel C and Marcotte EM: Calculating absolute and relative protein abundance from mass spectrometry-based protein expression data. Nat Protoc 3: 1444-1451, 2008.
- Lu P, Vogel C, Wang R, Yao X and Marcotte EM: Absolute protein expression profiling estimates the relative contributions of transcriptional and translational regulation. Nat Biotechnol 25: 117-124, 2007.
- Benjamini Y and Hochberg Y: Controlling the false discovery rate - a practical and powerful approach to multiple testing. J R Stat Soc B 57: 289-300, 1995.
- 8. Conesa A, Gotz S, Garcia-Gomez JM, Terol J, Talon M and Robles M: Blast2GO: a universal tool for annotation, visualization and analysis in functional genomics research. Bioinformatics 21: 3674-3676, 2005.
- Yamada T, Letunic I, Okuda S, Kanehisa M and Bork P: IPath2.0: interactive pathway explorer. Nucleic Acids Res 39: W412-W415, 2011.
- Shimizu-Hirota R, Sasamura H, Kuroda M, Kobayashi E and Saruta T: Functional characterization of podocan, a member of a new class in the small leucine-rich repeat protein family. FEBS Lett 563: 69-74, 2004.
- Hoover KB, Liao SY and Bryant PJ: Loss of the tight junction MAGUK ZO-1 in breast cancer: relationship to glandular differentiation and loss of heterozygosity. Am J Pathol 153: 1767-1773, 1998.
- 12. Liu W, Liu B, Xin L, Zhang Y, Chen X, Zhu Z and Lin Y: Down-regulated expression of complement factor I: a potential suppressive protein for gastric cancer identified by serum proteome analysis. Clin Chim Acta 377: 119-126, 2007.
- 13. Ohnuma SI: Cancer diagnosis and treatment. Google Patents, 2011
- 14. Ostrow KL, Park HL, Hoque MO, *et al*: Pharmacologic unmasking of epigenetically silenced genes in breast cancer. Clin Cancer Res 15: 1184-1191, 2009.
- Tachezy M, Zander H and Wolters-Eisfeld G: Activated leukocyte cell adhesion molecule (CD166): an 'inert' cancer stem cell marker for non-small cell lung cancer? Stem Cells 32: 1429-1436, 2014.
- Ibrahim SA, Reed KR, Clarke AR, Pritchard DM and Jenkins JR: Sa1992 SFRS2 and CDC5L interaction: an insight into downstream events following APC deletion during colorectal carcinogenesis. Gastroenterology 144: S353-S354, 2013.
- Palmieri D, Bouadis A, Ronchetti R, Merino MJ and Steeg PS: Rab11a differentially modulates epidermal growth factor-induced proliferation and motility in immortal breast cells. Breast Cancer Res Treat 100: 127-137, 2006.
- 18. Klement RJ and Kammerer U: Is there a role for carbohydrate restriction in the treatment and prevention of cancer? Nutr Metab 8: 75, 2011.

- Walenta S, Wetterling M, Lehrke M, Schwickert G, Sundfør K, Rofstad EK and Mueller-Klieser W: High lactate levels predict likelihood of metastases, tumor recurrence, and restricted patient survival in human cervical cancers. Cancer Res 60: 916-921, 2000.
- 20. Seyfried TN and Shelton LN: Cancer as a metabolic disease. Nutr Metab 7: 7, 2010.
- Skonier JE, Bowen MA, Emswiler J, Aruffo A and Bajorath J: Recognition of diverse proteins by members of the immunoglobulin superfamily: delineation of the receptor binding site in the human CD6 ligand ALCAM. Biochemistry 35: 12287-12291, 1996
- Weidle HU, Eggle D, Klostermann S and Swart GW: ALCAM/ CD166: cancer-related issues. Cancer Genom Proteom 7: 231-243, 2010
- 23. Van Kilsdonk JW, Wilting RH, Bergers M, van Muijen GN, Schalkwijk J, van Kempen LL, and Swart GW: Attenuation of melanoma invasion by a secreted variant of activated leukocyte cell adhesion molecule. Cancer Res 68: 3671-3679, 2008.
- Ofori-Acquah SF and King JA: Activated leukocyte cell adhesion molecule: a new paradox in cancer. Transl Res 151: 122-128, 2008.
- Ihnen M, Müller V, Wirtz RM, et al: Predictive impact of activated leukocyte cell adhesion molecule (ALCAM/CD166) in breast cancer. Breast Cancer Res Treat 112: 419-427, 2008.
- 26. Mezzanzanica D, Fabbi M, Bagnoli M, et al: Subcellular localization of activated leukocyte cell adhesion molecule is a molecular predictor of survival in ovarian carcinoma patients. Clin Cancer Res 14: 1726-1733, 2008.
- 27. Tachezy M, Zander H, Marx AH, Stahl PR, Gebauer F, Izbicki JR and Bockhornet M: ALCAM (CD166) expression and serum levels in pancreatic cancer. PLoS One 7: e39018, 2012.
- 28. Tachezy M, Effenberger K, Zander H, et al: ALCAM (CD166) expression and serum levels are markers for poor survival of esophageal cancer patients. Int J Cancer 131: 396-405, 2012.
- 29. Tachezy M, Zander H, Gebauer F, Marx A, Kaifi JT, Izbicki JR and Bockhorn M: Activated leukocyte cell adhesion molecule (CD166) its prognostic power for colorectal cancer patients. J Surg Res 177: e15-e20, 2012.
- Tachezy M, Zander H, Marx AH, et al: ALCAM (CD166) expression as novel prognostic biomarker for pancreatic neuroendocrine tumor patients. J Surg Res 170: 226-232, 2011.
- 31. Ishiguro F, Murakami H, Mizuno T, *et al*: Membranous expression of activated leukocyte cell adhesion molecule contributes to poor prognosis and malignant phenotypes of non-small-cell lung cancer. J Surg Res 179: 24-32, 2013.
- 32. Simon R, Mirlacher M and Sauter G: Immunohistochemical analysis of tissue microarrays. Methods Mol Biol 664: 113-126, 2010.