ORIGINAL ARTICLE



Identification of citrullinated peptides in the synovial fluid of patients with rheumatoid arthritis using LC-MALDI-TOF/TOF

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Abstract The objective of the study is to investigate potential citrullinated autoantigens as targets of anti-citrullinated protein antibodies (ACPAs) response in synovial fluids (SFs) of patients with rheumatoid arthritis (RA). SFs from six RA patients and six osteoarthritis (OA) patients as controls were collected. The citrullinated proteins in SFs were extracted by immunoprecipitation with rabbit anti-citrulline antibodies. Matrix-assisted laser desorption/ionization time of flight mass spectrometry/time of flight mass spectrometry (MALDI-TOF/TOF) mass spectrometry was subsequently performed to discover a characteristic neutral loss to finally determine citrullinated autoantigens. A total of 182 citrullinated peptides and 200 citrullinated sites were identified in RA SFs, while 3 citrullinated peptides and 4 citrullinated sites were identified in OA SFs. The 182 citrullinated peptides from RA SFs and

Fei Wang and Fang-Fang Chen contributed equally to this work.

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the 3 citrullinated peptides from OA SFs were derived from 83 and 3 autoantigens, respectively. Eighty-three autoantigens except protein-arginine deiminase type-2 (PADI2) and protein-arginine deiminase type-2 (PADI4) were overcitrullinated compared with controls, and the citrullinated sites of PADI2 and PADI4 were different in two groups. Interestingly, citrullinated histone H3.3 (H3F3A) was found in OA controls, but not in RA groups. The differential citrullinated proteins identified in RA SFs suggested potential autoantigens were targeted for ACPAs response and might contribute to the induction and perpetuation of complement activation and joint inflammation in RA.

Keywords Citrullinated protein · LC-MALDI-TOF/TOF · Rheumatoid arthritis · Synovial fluid

Introduction

Rheumatoid arthritis (RA) is an autoimmune disease characterized by the formation of inflammatory, invasive tissue, and rheumatoid pannus in synovial membranes, subsequently resulting in joint destruction and systemic complications. The related autoimmunity is often associated with certain major histocompatibility complex (MHC) types and the presence of anti-citrullinated protein antibodies (ACPAs) [1]. ACPAs are important biomarkers of RA and can be detected even before the clinical onset of the disease; consequently, they are recognized as a predictive and diagnostic marker. Furthermore, ACPAs in the inflammatory synovium bind to citrullinated autoantigens to form immune complexes (ICs), which lead to the development of inflammation [2–7]. Thus, a simple and effective method is needed to detect citrullinated proteins in the joint fluid from RA patients.

Citrullination is a post-translational modification (PTM) involving the conversion of an arginine residue to a noncoded citrulline residue, catalyzed by peptidylarginine deiminases (PADIs). This PTM leads to the loss of a positive charge and a reduction in hydrogen-bonding ability [8]. The traditional method to detect citrullinated proteins in biological fluids is two-dimensional polyacrylamide gel electrophoresis (2D-PAGE) followed by immunoblotting and Fourier transform ion cyclotron resonance (ICR) mass spectrometry (MS) analysis, which is labor-intensive and time-consuming [9–12]. Moreover, the mass shift of citrullination is very small (+1 Da), which can result in false positives [4,13]. Hao et al. [14] found that one specific signature of citrullination is that the neutral loss of 43 Da from the peptidyl-citrulline can be observed after collision-induced dissociation (CID) during triple quadruple/linear ion trap (Q-Trap) mass spectrometry, which indicates the elimination of isocyanic acid from the citrulline ureido group as shown in (Fig. 1). However, this technology has not been applied to human fluid because the complexity of the protein mixture made detection of lowabundance proteins very difficult. In addition, the low mass resolution of Q-trap MS prevented the reliable peptide and PTM characterization as in the high mass resolution of timeof-flight (TOF) MS. Thus, we first applied liquid chromatography-matrix-assisted laser desorption/ionization (LC-MALDI)-TOF/TOF to detect citrullinated proteins in human RA synovial fluid (SF).

In this study, citrullinated proteins were extracted by immunoprecipitation using agarose-conjugated rabbit anticitrulline antibodies, followed by LC-MALDI-TOF/TOF MS analysis. The peptide sequences and citrullinated sites in RA SFs or osteoarthritis (OA) SFs were identified. The highenergy CID mode of MALDI-TOF/TOF (20 keV collision energy) was adopted to improve fragmentation efficiency for reliable peptide and PTM characterization, thereby enabling the identification of potential autoantigens for ACPAs.

Methods

Human sample collection

Samples of SFs were consecutively obtained from knee joints of patients with RA or patients with OA, as a control, during knee therapeutic arthrocentesis at the Department Osteology, Jinling Hospital, School of Medicine, Nanjing University from January 2011 to January 2012. In order to improve the possibility of the existence of citrullinated antigens, patients of RA fulfilling the criteria (serum ACPA >200 U/ml and synovial fluids ACPA >200 U/ml) were selected and SFs of patients who were diagnosed as OA were collected. Patients who had incomplete data were excluded. The diagnosis of RA was made according to the American College of Rheumatology/

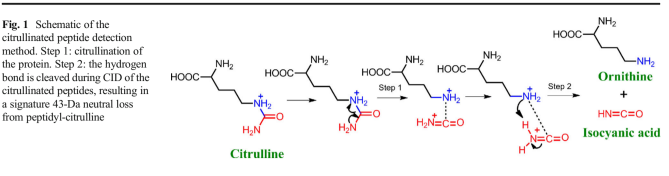
European League Against Rheumatism Collaborative Initiative 2010 criteria [15] and the diagnosis of OA was based on the 1986 clinical and radiological criteria for the diagnosis of knee OA developed by the American College of Rheumatology [16]. Finally, six patients with RA were obtained and six patients with OA were selected by random. The patients' basic information and serologic profile are shown in (Table 1). Informed consent was obtained from all subjects and the study was approved by the local ethics committee (Nanjing, China).

Immunoprecipitation

All SF samples were centrifuged at $25,000 \times g$ for 10 min at 4 °C and the supernatants were stored separately in sterile conditions at -80 °C. EDTA was added at a final concentration of 50 mM, followed by centrifugation at 14,000×g for 10 min at 4 °C, and the supernatant or sample was transferred to a new vial. Protein concentrations were determined by BioSpecnano (Shimadzu Biotech, Kyoto, Japan), and 1 mg was then subjected to immunoprecipitation [17,18]. Anti-citrulline polyclonal antibody (pAb; Abcam, Cambridge, USA) was cross-linked to protein G-Agarose (Sigma Aldrich, St Louis, MO, USA) with stable amide linkages according to the manufacturer's instructions. Citrullinated proteins were immunoprecipitated by incubating the supernatant/sample with agarose-conjugated rabbit anti-citrulline antibodies overnight. The samples were washed three times with 50 mM ammonium acetate pH 7.4 and once with H₂O, and the bound peptides were then eluted with acetonitrile/H₂O (7:3 v/v) containing 5 mM HCl. The eluted peptides were subjected to desalting by C18 tip (SciLifeLab, Shanghai, China). The efficiency of immunoprecipitation was determined on equal amounts of protein/sample (OA, RA, washes of RA, and eluents of RA) by immunoblotting using anti-citrulline pAb.

Protein reduction, alkylation, and enzyme digestion

An additional 937 μ L of 50 mM NH₄HCO₃ was added to the eluted proteins. The proteins were reduced by adding 10 μ L of 0.5 M dithiothreitol (DTT) in 50 mM NH₄HCO₃ to a final concentration of 4 mM and incubated for 20 min at 56 °C. For alkylation, 27 μ L of 0.55 M iodoacetamide was added, and the samples were incubated for 15 min at room temperature in the dark. The final concentration of the extracted proteins was approximately 2.4 mg/mL (total volume approximately 1250 μ L) according to the optical density at 280 nm. ProteaseMax solution (10 μ L of 1 %) was added together with 50 μ L of 1 μ g/ μ L LysC before incubation in a hydrated chamber at 37 °C overnight. The reaction was quenched by adding formic acid (FA; Fluka, Sigma) to a final concentration of 0.5 %. The digested peptides were then subjected to desalting by C18 tip (Shimadzu Biotech, Kyoto, Japan). Finally, the



desalted peptides were immediately applied to a prominence nano2D-HPLC and AccuspotTM system (Shimadzu Biotech, Kyoto, Japan).

2D-NanoLC fractionation

 Table 1
 Clinical and laboratory

 characteristics of the RA and OA

patients

A 5- μ L aliquot of the desalted peptide/sample was loaded directly onto a PolySulfoethyl A Column (1.0 mm × 50 mm, 5 μ m) for the first dimensional strong cation exchange separation (A=10 mM FA, B=600 mM FA). The samples were then subjected to a second dimensional RP separation using a C a pillary EX-Nano MonoCap C-18 column (0.16 mm × 150 mm, 5 μ m): A=5% acetonitrile+0.1% FA, B=95% acetonitrile+0.1% FA. The flow rate of the system was set at 4 μ L/min post-split. The eluent passed through a UV/Vis detector (220 nm) and was mixed with α -cyano 4hydroxycinnamic acid matrix (CHCA; 5 mg/ml in 50/50 acetonitrile/0.1% trifluoroacetic acid) and deposited onto a stainless steel MALDI target using the AccuspotTM LC-MALDI deposition robot [19]. LC-MALDI-TOF/TOF MS analysis

Samples were deposited at a volume of ~1 μ L/spot. The signal-to-noise ratio (S/N) was determined using Launchpad version 2.9.1 software (Shimadzu Biotech, Kyoto, Japan). The limit of detection (LOD) was established with a S/N of 3:1. The m/z value was calibrated with 50 fmol each of human angiotensin II and human adrenocorticotropin fragment 18–39 and 250 fmol each of bovine insulin oxidized beta chain and bovine insulin as external standards. The m/z reported in MALDI-TOF/TOF (MALDI-7090, Shimadzu Kratos, Manchester, UK) was set in positive ion mode and a mass range of 1000–4000 Da. The peptide ions with high S/N (S/N>10:1) were subjected to CID for subsequent MS/MS analysis.

Bioinformatics analysis

The raw MS/MS data were searched using the Mascot engine and then processed with PTM Finder[™] Software (Shimadzu Kratos) using the following criteria: database, Swiss-Prot,

Diagnosis	Sex	Age (years)	ACPA, (U/ml)	RF (IU/ ml)	ESR (mm/h)	CRP (mg/dl)	Disease duration (years)	DAS28
RA	Female	61	>200	86.3	64	25.1	3	4.28
RA	Female	79	>200	397	109	42.7	4	4.93
RA	Female	73	>200	40.6	44	37.5	2	3.84
RA	Male	69	>200	24.5	29	27.3	3	3.43
RA	Male	63	>200	142	44	77.8	5	3.72
RA	Male	70	>200	102	82	31.4	2	4.16
OA	Female	68	< 0.5	<20	8	8.9	2	_
OA	Female	53	< 0.5	<20	7	7.8	3	_
OA	Female	63	< 0.5	<20	5	2.5	2	_
OA	Male	69	< 0.5	<20	3	0.6	5	_
OA	Male	54	< 0.5	<20	7	4.3	4	_
OA	Male	58	< 0.5	<20	14	9.9	3	_

ACPA anticitrullinated protein antibody, RF rheumatoid factor, ESR erythrocyte sedimentation rate, CRP C-reactive protein, DAS28 disease activity score at 28 joints

human; enzyme, LysC; miscleavages, 2; static modifications, carbamidomethylation of cysteine (+57.02 Da); variable modifications, oxidation of methionine (+16.00 Da); neutral loss of isocyanic acid from peptidyl-citrulline (-43.02 Da); precursor ion tolerance, 0.3 Da; fragment ion tolerance, 0.8 Da. At this point, the citrullinated sites were identified by MS/MS analysis of the AA(8)AA ion because an ornithine residue (Orn) was expected to be the product after loss of a carbamyl group. Thus, the neutral loss of isocyanic acid from peptidyl-citrulline could be differentiated from the deamidation of peptidylasparagine or peptidyl-glutamine. All entries were filtered using a false positive rate of 1 % at the peptide levels, and false positives were removed. The citrullinated proteins from the RA SFs were further analyzed with DAVID Bioinformatics Resources (David 6.7 software, Bethesda, Maryland, USA) to understand their biological functions.

Results

A total of 182 citrullinated peptides and 200 citrullinated sites were identified in the RA SFs, while only three citrullinated peptides and four citrullinated sites were identified in the OA SFs (Tables 2 and 3). The 182 citrullinated peptides from the RA SFs were derived from 83 autoantigens, and the three citrullinated peptides from the OA SFs were derived from three autoantigens. The autoantigens in the RA SFs were over-citrullinated compared with the controls. Among these, 26 citrullinated proteins identified here have also been validated in previous studies (Table 4), which suggests that this strategy for identifying citrullinated peptides is highly effective.

Functional analysis of the identified citrullinated proteins in the RA group was performed with David 6.7 software. The categories of "disease" and "gene ontology" reported a significant enrichment of RA and acute inflammatory responseassociated genes, which corresponded to the physiological status of the patients in the present study (Supplement Fig. 1). Furthermore, genes involved in the enriched pathways in the list were associated with complement and coagulation cascades (Supplement Fig. 2). Additionally, proteins that were involved in cell differentiation, metabolism, redox state and apoptosis, regulation and transport, immune response and acute phase, structural and cell adhesion, and other groups based on the NCBI and UniProt database information are shown in (Fig. 2a). Moreover, the proteins were also classified by their subcellular location, as described in (Fig. 2b). The results of our analysis demonstrated that the citrullinated proteins obtained with our protocol provide reliable data on the state of citrullination in RA SF.

Discussion

The major methods used currently to identify citrullinated proteins employ 2D-PAGE followed by immunoblotting and Fourier transform ion cyclotron resonance mass spectrometry analysis. For example, J.B.C. van Beers et al. found 192 proteins including 53 citrullinated proteins with their citrullinated residues in RA SFs [12]. One problem with this method is the small mass shift (+1 Da) from the conversion of peptidyl arginine to Cit, which is challenging for mass detection to distinguish. In the present study, citrullinated proteins were effectively enriched following immunoprecipitation (Supplement Fig. 1). NanoLC was then used to fractionate the tryptic digests of citrullinated proteins to improve the sensitivity and dynamic range of protein identification. With this method, not only are peptides of the same nominal mass isolated by temporal separation, but signal suppression is also reduced because of the separation of low- and highabundance peptides. Importantly, the unique LC-MALDI peak picking algorithm promotes the MS/MS of selected ions at the apex of the eluting chromatographic peak to allow the most efficient data acquisition. This is not often the case with ES LC-MS/MS, where MS/MS acquisition is often taken on the rising edge of the eluting chromatographic peak. In addition, the high-energy CID mode of MALDI-TOF/TOF (20 keV collision energy, MALDI-7090) allowed us to determine the citrullinated sites more easily, according to the characteristic neutral loss of an isocyanic acid group from peptidyl-citrulline.

A number of chaperone molecules were identified within the SFs, particularly heat shock 70 kDa protein 1A/1B (HSPA1A), glucose-regulated protein 78 kDa (GRP78 or HSPA5), and HSP90AA1, members of the stress-inducible heat-shock protein 70 family. Also, we previously found GRP75 (HSP70) and binding immunoglobulin protein (BiP or GRP78) in RA synovial fibroblast-like synoviocytes (FLSs) [20]. Citrullinated BiP induces anti-CCP and anticitrullinated fibrinogen antibodies and exacerbates collageninduced arthritis in mice, and deaminated HSP90 was identified as a diagnostic autoantigen for a potentially serious manifestation of RA [10,21]. Recently, HSPs have been reported that not only act as chaperones during protein folding but also play a role between ubiquitin E3 ligase and the proteasome to inhibit proinflammatory NF-KB signaling [22]. In addition, both canonical and non-canonical NF-KBs are overexpressed in RA and are associated with the persistence of inflammation in RA [23]. Thus, citrullination of HSP may contribute to the chronic inflammation in the synovium or dysregulation of RA synovial fibroblasts, suggesting that citrullination may correlate with complement activation and the perpetuation of RA.

In a previous study from our group, we also reported that the elevated Annexin A11 in FLSs may be associated with the

Table 2 Citrullinated peptides and their deaminized sites identified by MALDI-TOF-MS in the RA SFs

Protein ID	Peptide sequence	Citrullinated sites
A2MG_HUMAN	DNGCF R SSGSLLNNAIK	R1081
	GN R IAQWQSFQLEGGLK	R174
	EQAPHCICANG R QTVSWAVTPK	R853
	FQVDNNN R LLLQQVSLPELPGEYSMK	R1297
ACTG_HUMAN	DLYANTVLSGGTTMYPGIAD R MQK	R312
	AGFAGDDAP R AVFPSIVGRPRHQGVMVGMGQK	R28
ALBU_HUMAN	AWAVARLSQ R FPK	R246
	LCTVATL R ETYGEMADCCAK	R105
	VHTECCHGDLLECADD R ADLAK	R281
	RMPCAEDYLSVVLNQLCVLHEK	R469
	YLYEIAR R HPYFYAPELLFFAK	R169
	CCTESLVNR R PCFSALEVDETYVPK	R509
ANXA1 HUMAN	DITSDTSGDF R NALLSLAK	R177
—	GTDVNVFNTILTT R SYPQLRRVFQK	R228
APOA1 HUMAN	ENGGARLAEYHAK	R212
—	VEPLRAELQEGA R QK	R155
		R51
		R239
APOB HUMAN		R3386
—	LTTNG R FREHNAK	R1689
	AEFTG R HDAHLNGK	R3020
		R207
		R2449
		R4519
		R75
F5GZW3 HUMAN		R486
		R73, R77
		R889
		R198
ARC1B HUMAN		R294, R299
		R100
C1R HUMAN		R149
		R420
		R388
C1S HUMAN		R586
_		R77
—		R880
		R315
		R945
		R1060
		R1310
		R1134
COAR HIMAN		R1134 R218
		R218 R80
		R1675
		R1349
C4BPA_HUMAN		R493
004 183 444		R353
CO6_HUMAN	FRCDSG R CIARK	R150
	Protein ID A2MG_HUMAN ACTG_HUMAN ALBU_HUMAN ALBU_HUMAN ANXA1_HUMAN APOA1_HUMAN APOB_HUMAN APOB_HUMAN C1R_HUMAN C1R_HUMAN C1S_HUMAN C03_HUMAN C03_HUMAN	A2MG_HUMAN DNGCFRSSGSLLNNAIK GNRIAQWQSFQLEGGLK EQAPHCICANGRQTVSWAVTPK FQVDNNNRLLQQVSLPELPGEYSMK ACTG_HUMAN AUXARLSQRFPK LCTVATLRETYQGEMADCCAK VHTECCHGDLLECADDRADLAK RMPCAEDYLSVUNQLCVLHEK YLVELARRHPYTAPELLFAK CCTESLVNRRPCFSALEVDETYVPK ANXA1_HUMAN AWAVARLSQRFPK LCTVATLRETYQGEMADCCAK VHTECCHGDLLECADDRADLAK RMPCAEDYLSVVLNQLCVLHEK YLVELARRHPYTAPELLFAK CCTESLVNRRPCFSALEVDETYVPK ANXA1_HUMAN DITSDTSGDFRNALLSLAK GTDVNVFNTILLTRSYPQLRRVFQK APOA1_HUMAN ENGGARLAEVHAK VEPLRAELQEGARQK DSGRDYVSQFEGSALGK PALEDLRQGLLPVLESFK APOB_HUMAN LEGTTRLITK LITINGRFREHNAK AEFTGRHDAHLNGK GNVATEISTERDLGQCDFFK IREVTQRLNGEIQALELPQK RLIDSIGNYHTFLIYTELLK YTYNYEAESSSGVFGTADSRSATRINCK FSGZW3_HUMAN EEQEVSWTQYTQRK AEFFSSRGJCSSREHQSFRK ELLGKTSVRQGLGPASTTSPSPGPRSK LREAEFQEEKAAGRSVPTTTAGATEAGPLRK QSSQRGLTAREFQNLDK MQTRAGSRESEQGVYTCTAQGIVK CCQ_PINLPNGDFRXTTIMGNTYK CIS_HUMAN SSQWQTPGATRSLSK C03_HUMAN RRHQQTVTIPFK VLLDFQURGEDLEVGK RTAGSRESEQGVYTCTAQGIVK CCQ_HUMAN RRHQQTVTIPFK VLLDGVQNRAEDLVGK TVAVRTLDPERLGREGVQK GYTQLAFRQSDFRX ELLGKTSVRQGLGRASTSSTFFNENK CIS_HUMAN SSQWQTPGATRSLSK C03_HUMAN RRHQQTVTIPFK VLLDGVQNRAEDLVGK TVAVRTLDPERLGREGVQK GYTQLAFRQSSRTQFLQGELCPEFK HUMAN SSGPQUTPGATRSLSK C04B_HUMAN PSSVGLENNEK CO4B_HUMAN PLVNGRLSEVFX VDFTLSSERDFALLSLQVPLK AANQMRNFIVRASCRLEPGNLSK C04B_HUMAN PLVNGRLSEVFX VDFTLSSERDFALLSLQVPLK AANQMRNFIVRASCRLEPFSK ELLGKTSVRQGLFREFQNLSK C04B_HUMAN PLVNGRLSEVFX VDFTLSSERDFALLSLQVPLK AANQMRNFIVRASCRLEPFSK C4BPA_HUMAN PLVNGRLSEVFX VDFTLSSERDFALLSLQVPLK AANQMRNFIVRASCRLEPFSK C4BPA_HUMAN PLVNGRLSEVFX VDFTLSSERDFALLSLQVPLK AANQMRNFIVRASCRLEPFSK C4BPA_HUMAN PLVNGRLSEVFX VDFTLSSERDFALLSLQVPLK AANQMRNFIVRASCRLEPFSK SCWB_HUMAN PLVNGRLSVDK NLWTYYQGCEALCCPFFK

Table 2 (continued)

Gene name	Protein ID	Peptide sequence	Citrullinated sites
		RSENINHNSAFK	R289
		SSRTSNPYRVPANLENVGFEVQTAEDDLK	R225
С9	CO9 HUMAN	NFRTEHYEEQIEAFK	R213
CAT	CATA HUMAN	NAIHTFVQSGSHLAAREK	R522
CD44	CD44 HUMAN	NGRYSISRTEAADLCK	R41
	—	EQWFGN R WHEGY R QTPK	R407, R413
CFH	CFAH HUMAN	RITCRNGQWSEPPK	R1149, R1153
	—	HGGLYHENMRRPYFPVAVGK	R340
		RGYRLSS R SHTL R TTCWDGK	R1210, R1215
		AQTTVTCMENGWSPTP R CIRVK	R441
		IPCSQPPQIEHGTINSSRSSQESYAHGTK	R885
		CNMGYEYSERGDAVCTESGWRPLPSCEEK	R246
CFHR2	FHR2 HUMAN	SHSFRAMCQNGK	R254
CFI	CFAI HUMAN	DNERVFSLQWGEVK	R480
		THRYQIWTTVVDWIHPDLK	R389
CHMP2A	CHM2A HUMAN	MDLLFGR R K	R8
		DLVRTRRYVRK	R71
CLC	LEG10 HUMAN	YQVMVNGQSSYTFDH R IK	R/1 R115
CP	CERU HUMAN	ALYLOYTDETFRTTIEK	R81
CI	CERO_HOWAN	NLASRPYTFHSHGITYYK	R115
		ENLTAPGSDSAVFFEQGTTRIGGSYK	R415
		DNEDFQESNRMYSVNGYTFGSLPGLSMCAEDRVK	R415 R258
		NMATRPYSIHAHGVQTESSTVTPTLPGETLTYVWK	R830
CPB2	CDDD2 HUMAN	DHEELSLVASEAVRAIEK	R342
CTLA4	CBPB2_HUMAN		R542 R51
CILA4	CTLA4_HUMAN	AMHVAQPAVVLASSRGIASFVCEYASPGK	
ENO1		AQLNLATRTWPCTLLFFLLFIPVFCK	R18
ENO1	ENOA_HUMAN	TGAPCRSERLAK	R403
F2		LAQANGWGVMVSHRSGETEDTFIADLVVGLCTGQIK	R372
F2	THRB_HUMAN	YTACETARTPRDK DSTDIDITDNIMECA CVK	R94, R97
DE3		DSTRIRITDNMFCAGYK	R541
FF3	AFF3_HUMAN	DFLTDRSNQSHLVGVPK	R111
		EAAANGGSGPRAPVGSINARTTSDIAK	R745
		YTSEDLTSSSRPNGNSLFTSASSSK	R926
EG.		SPPAAVAVAVSAAAPPPAVPCAPAENAPAPAR R SAGK	R606
FGA	FIBA_HUMAN	NVRAQLVDMK	R160
FOD		GLIDEVNQDFTNRINK	R84
FGB	FIBB_HUMAN	REEAPSLRPAPPPISGGGYRARPAK	R60, R72
FGG		EDGGGWWYNRCHAANPNG R YYWGGQYTWDMAK	R445
FGG	FIBG_HUMAN	YEASILTHDSSIRYLQEIYNSNNQK	R134
		YTGNTY R VGDTYERPK	R106
		WLPSSSPVTGY R VTTTPK	R1573
~~	· · · · · · · · · · · · · · · · · · ·	DNRGNLLQCICTGNG R GEWK	R265
GC	VTDB_HUMAN	HLSLLTTLSNRVCSQYAAYGEK	R218
		HQPQEFPTYVEPTNDEICEAF R K	R149
		RSDFASNCCSINSPPLYCDSEIDAELK	R445
H1FX	H1X_HUMAN	VPWFDQQNG R TYLK	R86
		YSQLVVETI R RLGE R NGSSLAK	R57, R62
H2AFY	H2AY_HUMAN	SIAFPSIGSG R NGFPK	R318
HABP2	HABP2_HUMAN	EEFHEQSF R VEK	R391

Table 2 (continued)

2191

Gene name	Protein ID	Peptide sequence	Citrullinated sites
		FCEIGSDDCYVGDGYSY R GK	R203
		LIANTLCNSRQLYDHMIDDSMICAGNLQK	R480
HIST2H2AC	H2A2C_HUMAN	TRIIPRHLQLAIRNDEELNK	R89
		GNYAE R VGAGAPVYMAAVLEYLTAEILELAGNAA R DNK	R43, R72
HMGB2	HMGB2_HUMAN	MSSYAFFVQTC R EEHK	R24
HNRNPA1L2	RA1L2_HUMAN	GGNFGG R SSGPYGGGGQYFAK	R284
HP	HPT_HUMAN	VSVNE R VMPICLPSK	R261
		YVMLPVADQDQCI R HYEGSTVPEK	R311
HPR	HPTR_HUMAN	VLVNE R VMPICLPSK	R203
HPX	HEMO_HUMAN	NFPSPVDAAF R QGHNSVFLIK	R102
HSP90AA1	Q8TBA7_HUMAN	AQALRDNSTMGYMAAK	R620
	` _	HNDDEQYAWESSAGGSFTV R TDTGEPMGRGTK	R173
HSPA1A	HSP71 HUMAN	LLQDFFNGRDLNK	R357
	—	EIAEAYLGYPVTNAVITVPAYFNDSQ R QATK	R155
HSPA5	GRP78 HUMAN	SDIDEIVLVGGSTRIPK	R368
		RLIGRTWNDPSVQQDIK	R98
Ig kappa chain V-II region RPMI 6410	KV206_HUMAN	VSNRDSGVPDRFSGSGSGTDFTLK	R79
Ig lambda chain V-II region NEI	LV202 HUMAN	RPSGVSNRFSGSK	R56, R63
Ig lambda chain V-II region NIG-84	LV211_HUMAN	LLIYDVNSRPSGISNRFSGSK	R56, R63
IGHA1	IGHA1_HUMAN	YLTWASRQEPSQGTTTFAVTSILRVAAEDWK	R282
IGHG3	IGHG3_HUMAN	SCDTPPPCPRCPEPK	R128
		TPLGDTTHTCPRCPEPK	R113
ING4	ING4_HUMAN	WFCPRCSQERK	R241
ITIH2	ITIH2 HUMAN	RLSNENHGIAQRIYGNQDTSSQLK	R475
111112		TILDDLRAEDHFSVIDFNQNIRTWRNDLISATK	R475 R356, R359
KNG1	VNC1 HUMAN	ICVGCPRDIPTNSPELEETLTHTITK	R350, R359 R268
	KNG1_HUMAN		
KRT33B	KT33B_HUMAN	ETMQFLNDRLASYLEK	R66
LBR	LBR_HUMAN	EARREVEVK	R111
		PLTSFRQRK	R61
		SAR R SASASHQADIK	R96
		ELAVRTFEVTPIRAK	R195
I CD1		APRNDLSPASSGNAVYDFFIGRELNPRIGTFDLK	R353
LCP1	PLSL_HUMAN	GDEEGVPAVVIDMSGL R EK	R316
		ALENDPDCRHVIPMNPNTNDLFNAVGDGIVLCK	R141
LGALS3BP	LG3BP_HUMAN	SGGSDRTIAYENK	R514
LDC1		SQLVYQSR R GPLVK	R436
LRG1	A2GL_HUMAN	ALGHLDLSGNRLRK	R175
		LQVLGKDLLLPQPDLRYLFLNGNK	R239
MAPRE1	MARE1_HUMAN	PLTSSSAAPQRPISTQ R TAAAPK	R168
MMP8	MMP8_HUMAN	FYQLPSNQYQST R K	R52
MNDA	MNDA_HUMAN	INQEEVGLAAPAPTARNK	R119
NCF1	NCF1_HUMAN	SGQDVSQAQ R QIK	R292
		STATDITGPIILQTY R AIANYEK	R162
ORM1	A1AG1_HUMAN	EQLGEFYEALDCLRIPK	R167
PABPC1	PABP1_HUMAN	AVTEMNGRIVATK	R356
		PVRIMWSQ R DPSL R K	R89, R94
		ITGMLLEIDNSELLHMLESPESL R SK	R604
PADI2	PADI2_HUMAN	VGVFYVENPFFGQ R YIHILGR R K	R225, R233
PADI4	PADI4 HUMAN	GFRLLLASP R SCYK	R495

Table 2 (continued)

Gene name	Protein ID	Peptide sequence	Citrullinated sites	
		TLREHNSFVERCIDWNRELLK	R536	
		PFGPVING R CCLEEK	R609	
POFUT2	OFUT2_HUMAN	VFVATDAV R K	R337	
		DFIWGHRQDVPSLEGAV R K	R315	
PPIA	PPIA_HUMAN	EGMNIVEAMERFGSRNGK	R148	
PRG4	PRG4_HUMAN	DQYYNIDVPSRTA R AITT R SGQTLSK	R1386, R1391	
PRKCD	KPCD_HUMAN	QSM R SEDEAK	R132	
		SP R DYSNFDQEFLNEK	R628	
		IIGRCTGTAANS R DTIFQK	R216	
PTPN22	PTN22_HUMAN	GP R NPPPTWNI	R499	
		PAESVQSNNSSSFLNFGFAN R FSK	R491	
SAA2	SAA2_HUMAN	RGPGGAWAAEVISNA R ENIQRLTG R GAEDSLADQAANK	R80, R89	
SERPINA3	AACT_HUMAN	ADLSGITGA R NLAVSQVVHK	R350	
SERPINC1	ANT3_HUMAN	LVSAN R LFGDK	R177	
		IPEATN R RVWELSK	R78	
		ANSRFATTFYQHLADSK	R89	
SLC22A4	S22A4_HUMAN	VPLTTSLFFVGVLLGSFVSGQLSD R FGRK	R166	
STAT4	STAT4_HUMAN	NVSTLSN R RFVLCGTNVK	R173	
		SLQSSSVSE R QRNVEHKVAAIK	R139	
		FHGNPMHVAVVISNCL R EERRILAAANMPVQGPLEK	R110	
TAGLN2	TAGL2_HUMAN	MAN R GPAYGLSREVQQK	R4	
TF	TRFE_HUMAN	CSTSSLLEACTF R RP	R696	
		EGYYGYTGAF R CLVEK	R541	
		ADRDQYELLCLDNTRK	R239, R251	
TNC	TENA_HUMAN	NG R ENFYQNWK	R2033	
		RVTTT R LDAPSQIEVK	R802	
		VEAARNLTVPGSL R AVDIPGLK	R1127	
		PDTEYEVSLISR R GDMSSNPAK	R878	
		ETFTTGLDAP R NLRRVSQTDNSITLEWRNGK	R897	
TNC	TENA_HUMAN	VPEITRTVSGNTVEYALTDLEPATEYTL R IFAEK	R1866	
TNFAIP6	TSG6_HUMAN	NFLAG R FSHL	R273	
VIM	VIME_HUMAN	MALDIEIATY R K	R401	
	_	PDLTAAL R DV R QQYESVAAK	R270, R273	
VPRPB	VPRBP_HUMAN	FISGTPRRK	R707	
	—	SPFGSSF R TFNATDYK	R1334	

Citrullinated residues are indicated with a bold ${\bf R}$

extensive synovial fibroblast-like synoviocytes hyperplasia. Additionally, in the extracellular environment defined as synovial fluid, we found citrullinated AnnexinA1, another member of the annexin superfamily of structurally related Ca^{2+} -dependent phospholipid-binding proteins. Several other studies have demonstrated that AnnexinAl is a

Table 3Citrullinated peptidesand their deaminized sitesidentified by MALDI-TOF-MS inthe OA SFs

Gene name	Protein ID	Peptide sequence	Citrullinated sites
H3F3A	H33_HUMAN	DIQLARRIRGERA	R130
PADI2	PADI2_HUMAN	GFPVVLDSP R DGNLK	R373
PADI4	PADI4_HUMAN	TLPVVFDSP R N R GLK	R372, R374

Citrullinated residues are indicated with a bold R

 Table 4
 26 citrullinated proteins in our study were validated in previous studies

Protein ID	References
Arginine deiminase type-4	[19]
Alpha-1-acid glycoprotein 1	[12]
Alpha-2-macroglobulin	[12]
Annexin A1	[12]
Apolipoprotein A-I	[12]
Apolipoprotein B-100	[12]
Ceruloplasmin	[12]
C4b-binding protein alpha chain	[12]
Complement C2	[12]
Complement C4-B	[12]
Complement factor H	[12]
Enolase	[12,20]
Fibrinogen	[20]
Fibronectin	[12]
Hemopexin	[12]
HSP90	[12]
Histone	[8,21]
Inter-alpha-trypsin inhibitor heavy chain H2	[12]
Myeloid cell nuclear differentiation antigen	[12]
Plastin-2	[12,22]
Proteoglycan 4	[12]
Serotransferrin	[12]
Serum albumin	[12]
Tenascin	[12]
Vitamin D-binding protein	[12]
Vimentin	[11]

glucocorticoid-induced molecule that can be transferred into cartilage and can modulate T cell function and the adaptive immune responses relevant to RA [24,25]. Consistent with this, treatment of mice with dexamethasone promotes potent antiarthritic effects that are dynamically attenuated in AnxA1^{-/-} mice [26]. Our observations on citrullinated Annexin A1 reflect the possibility that citrullinated or non-citrullinated Annexin may be a target to minimize glucocorticoid use in RA.

The different citrullinated sites of PADI2 and PADI4 in the two groups suggest new potential biomarkers for RA. PADI2 and PADI4 are the only PAD isotypes expressed in the synovial tissue of patients with RA, and they were reported to induce differentiation and apoptosis [27]. PADI4, found in the cell nucleus, mediated gene transcription by regulating arginine citrullination and methylation in histones H1, H3, and H4 and was autocitrullinated during cell activation [8,28,29]. Interestingly, citrullinated H3F3A was found in the OA controls, but not in the RA group. In addition, histones H1x and H2A were only citrullinated in the RA group. These

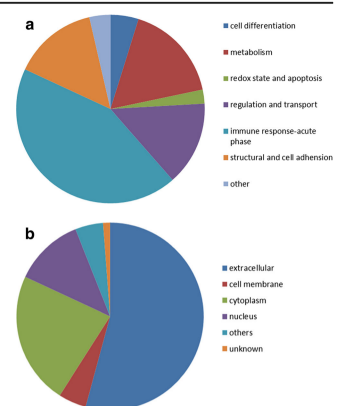


Fig. 2 Classification by a function and b subcellular location of the proteins identified in the RA SFs

results suggest that PADI2 and PADI4 represent a heterogeneous subtype with different citrullinated sites targeting multiple structural domains, where the specific citrullinated site may predict a specific disease. The exact mechanism underlying this phenomenon remains to be elucidated.

Although we identified the potential antigens for ACPA, some limitations remain, including the amount of patients was small, thus we pooled all samples per group to gain more sensitivity and to find more citrullinated antigens; results merely compared with previous studies; the validated process used only one method of mass spectrometry and was only on the basis of mass-spectrometry-based proteomics, so we performed DAVID Bioinformatics Resources to classify these genes corresponding to citrullinated proteins, at the same time, estimate and verify the reliability. Further studies will employ western blot to identify some selected potential autoantigens. At the same time, we will collect samples of synovial fluid or serum of RA patients as more as possible and then test antibodies corresponding to autoantigens in synovial fluid or serum of RA patients in order to obtain reliable results from clinical data. These limitations indicate the need for larger validation studies and prospective SFs studies in groups where larger samples are available.

Overall, we demonstrated a simple and efficient strategy for detecting citrullinated proteins and citrullinated sites in human RA SFs. In addition to the previously detected citrullinated proteins in RA SF, the novel citrullinated proteins identified by the data here may represent new antigens for ACPAs, as well as new markers for diagnosis. More importantly, this data will contribute to the search for the etiopathogenesis of, and new therapeutic targets for RA.

Compliance with ethical standards Informed consent was obtained from all subjects and the study was approved by the local ethics committee (Nanjing, China).

Disclosures None.

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