Abstract

Compositions and methods for detection of anti-citrullinated protein antibodies (ACPAs) in rheumatoid arthritis (RA) patients. Patient samples known or suspected of containing ACPAs were probed against citrullinated proteins, and antibody responses to 150 citrullinated proteins in 20 RA patients were investigated. Unique antibody reactivity patterns in both clinical anti-cyclic citrullinated peptide assay positive (CCP+) and negative (CCP-) RA patients were observed. At individual antigen levels, six novel antibody/antigen complexes were discovered and validated against specific citrullinated antigens (Myelin Basic Protein (MBP), osteopontin (SPP1), flap endonuclease (FEN1), insulin like growth factor binding protein 6 (IGFBP6), insulin like growth factor 1 (IGF1) and stanniocalcin-2 (STC2)) in RA patients. Identification of immune-dominant epitope(s) for citrullinated MBP was also performed. The identified biomarkers have high specificity, especially MBP.

Specification includes a Sequence Listing.
Figure 1:

1. cDNA printed in PDMS microwells using piezo printer
2. Add in vitro protein expression lysates and cover wells with tag ligand coated slide
3. Expressed tagged proteins will be captured by ligands on the cover slide
4. Discard slide with PDMS microwells
5. Contra-captured protein microarray
6. On-array citrullination post-translationally
7. Binding of ACPA in serum to one citrullinated protein specifically
8. Detection of ACPA by dye-labelled secondary antibody

Legend: plasmid = tag ligand = tag = expressed proteins = arginine = citrulline
ACPA = dye-labelled secondary antibody = PDMS microwells = capture slide
Figure 2:
Figure 3:

Figure 4:
Figure 7:

MBP Isoform 1

MBP Isoform 2

Serum reactivity to deletion mutants

MBP (1.1) N 1-C 22
MBP (1.2) N 1-C 53
MBP (1.3) N 1-C 78
MBP (1.4) N 1-C 109
MBP (1.5) N 1-C 124

MBP (1.6) N 1-C 158
MBP (1.7) N 1-C 176
MBP (1.8) N 1-C 197

8 MBP Isoform 1
C-Terminal deletions

MBP (2.1) N 134-C 158
MBP (2.2) N 134-C 178
MBP (2.3) N 134-C 193

MBP (2.4) N 134-C 212
MBP (2.5) N 134-C 222
MBP (2.6) N 134-C 244
MBP (2.7) N 134-C 267
MBP (2.8) N 134-C 283

8 MBP Isoform 2
C-Terminal deletions
FIG. 8
IDENTIFICATION AND MEDICAL APPLICATIONS OF
ANTI-CITRULLINATED-PROTEIN
ANTIBODIES IN RHEUMATOID ARTHRITIS

CROSS-REFERENCE TO RELATED
APPLICATION

[0001] This application represents the national stage entry of
PCT International Application No. PCT/US2017/057492,
filed on Jun. 14, 2017, and claims priority to U.S. Provisi-
onal Application No. 62/350,064, filed on Jun. 14, 2016,
the disclosure of which is incorporated herein by reference.

STATEMENT REGARDING FEDERALLY
SPONSORED RESEARCH

[0002] This invention was made with government support
under R21 AR062220 and R42 GM106704 awarded by the
National Institutes of Health. The government has certain
rights in the invention.

REFERENCE TO A SEQUENCE LISTING
SUBMITTED VIA EFS-WEB

[0003] The content of the ASCII text file of the sequence
listing named “112624_00872_Sequence_Listing_ST25.txt”
which is 1.21 kb in size was created on Jun. 14, 2017
and electronically submitted via EFS-Web herewith the
application is incorporated herein by reference in its entirety.

FIELD OF THE INVENTION

[0004] This disclosure relates to biomarker complexes and
detection in the field of rheumatoid arthritis.

BACKGROUND OF THE INVENTION

[0005] Citrullination is a post translational modification
(PTM) that converts arginine to citrulline catalyzed by
peptidyl arginine deiminase (PAD). Anti-citrullinated pro-
tein antibodies (ACPA) have been specifically detected
in rheumatoid arthritis (RA) patients and show utility in RA
risk assessment and diagnosis. ACPA levels parallel with RA
disease activity, prognosticate erosive diseases, and serve as
a surrogate marker for treatment efficacy. Clinically, ACPA
are assayed using cyclic citrullinated peptide (CCP), which
measures a generalized reactivity with citrulline containing
peptides but does not provide information about reactivity
to disease-specific RA antigens.

[0006] Rheumatoid arthritis (RA) is a common autoim-
nune disease affecting 1% of the world population and is
characterized by the synovial inflammation and joint
destruction. Despite decades of research, the cause for
rheumatoid arthritis remains unknown. RA has been
observed to be heterogeneous through differences in clinical
presentation, and the production of different autoantibodies,
such as rheumatoid factor (RF) and ACPA. ACPA have been
specifically detected in RA patients. ACPA levels correlate
with RA disease activity, prognosticate erosive diseases and
serve as a surrogate marker for treatment efficacy. The exact
mechanism for ACPA production in RA is not clearly
understood because citrullination itself is not RA specific.
It has been hypothesized that genetic factors, such as specific
risk alleles for HLA class II genes and genes involved in
immune regulation, together with external factors, such as

smoking and viral infections contribute to the break of
the immune tolerance leading to the generation of ACPA.

SUMMARY OF THE INVENTION

[0007] Embodiments herein relate to a protein array plat-
form that enables the modification of many proteins in
parallel and assesses their immunogenicity without the need
to express, purify, and modify proteins individually. Anti-
citrullinated protein antibodies (ACPAs) in rheumatoid
arthritis (RA) were used as a model modification. We first
profiled antibody responses to 190 citrullinated proteins in
20 RA patients using a newly developed modification to
nuclease acid programmable protein arrays (NAPPA) to iden-
tify candidate RA specific antigens. ELISA assays of prom-
ising candidates were performed on 100 RA patients and 50
controls. From protein microarray screening experiments,
using a discovery set of 20 patients and 10 controls, 6
antigens showing higher reactivity in RA cases relative to
controls were subsequently selected for evaluation in a large
sample set (n=150) using enzyme-linked immunosorbance
assay (ELISA).

[0008] A previously unknown 6-antigen set—Myelin
Basic Protein (MBP), osteopontin (SFP1), lIap endonuclease
(FEN1), insulin like growth factor binding protein 6 (IG-
FBP6), insulin like growth factor I (IGF1), and stannio-
celin-2 (STC-2)—was identified in cyclic citrullinated peptide
positive (CCP+) RA samples with a sensitivity of 82%, 59%,
42%, 56%, 52% and 66% respectively at 95% specificity.
Additionally the 6-AAB antigens showed 22%, 12%, 10%,
18%, 12% and 12% sensitivity at 95% specificity for cyclic
citrullinated peptide negative (CCP-) samples. Thus, we
discovered previously unknown antigen complexes, i.e.,
AABs, associated with rheumatoid arthritis, with related
embodiments directed to diagnostics and rheumatoid arthri-
tis pathogenesis.

[0009] Embodiments herein include identification of
immune-dominant epitope(s) for citrullinated antigen MBP
using a protein array. Epitope mapping of MBP was per-
formed as the prevalence to anti-citrullinated (anti-cit)-MBP
(82%) was much higher than the other known or novel
citrullinated antigens (40-60%) in our sample set. We have
shown the two reactive epitopes on MBP: pattern 1, MBP
N144-C176 with the corresponding sequence HOSKY-
LATASTMDHARJIGILPRUHDTGILDSCG (SEQ ID NO. 1)
and in pattern 2, response was observed for polypeptides
MBP N168-C198, which includes the sequence D1GILD-
SIGRFFGGRAGPKRSGKVSSEE (SEQ ID NO. 2).

BRIEF DESCRIPTION OF THE DRAWINGS

[0010] FIG. 1: Principle of the contra capture protein array
platform.

[0011] FIG. 2: A. Heat map depicting overall reactivity of
20 CCP+ and CCP– serum samples to 190 genes printed on
array. CCP– RA samples were assayed as pools of two
samples as annotated on the top of the figure. Blue, low
reactivity; red, high reactivity. B. Example array images
probed with CCP+RA patient serum samples.

[0012] FIG. 3: Assaying 6 novel antigens against valida-
tion serums. Heat map depicting auto-reactivity against
citrullinated antigens assayed by ELISA in different groups
of the independent sample set.
FIG. 4: Sero-profiling of ACPs in RA patients on contra capture protein arrays against 190 antigens. Example array images probed with CCP+RA patient serum samples (S366 and S451).

FIG. 5: Analysis and verification of new targets identified on contra capture arrays by ELISA. 11 antigens identified on protein array assayed, by ELISA, against 30 serum samples from discovery set.

FIG. 6: Validation, by ELISA, of novel ACPs identified on contra capture arrays. Blinded validation of selected novel ACPs, by ELISA, using an independent set of 150 sera comprising 50 CCP+, 50 CCP− and 50 control samples. HC, healthy control. * indicates t-test p<0.05, ** p<0.01 and ***p<0.0001.

FIG. 7: Epitope mapping for MBP. Eight C-terminal deletion fragments were constructed for each isoform of MBP. Two response patterns were observed. In pattern 1, reactivity to deletion mutants MBP 1.7, MBP 1.8, MBP 2.2 and MBP 2.3 were observed. In pattern 2, reactivity to deletion mutants MBP 1.8 and MBP 2.3 were observed, but not to MBP 1.7 and MBP 2.2. MBP 1.7 and MBP 2.2 shared the same sequence and MBP 1.8 and MBP 2.3 shared same sequence. Red vertical lines indicate the immunodominant epitopes for the pattern 1 (sample S010) and for the pattern 2 (sample S922).

FIG. 8: Epitope mapping, by ELISA, for MBP using serial deletion fragments and tiling fragments. Top, diagram for the two MBP isoforms and serial deletion mutants used in this study. Middle, quantitative analysis of sero-reactivity to the deletion mutants for S922 and S100 serum samples on arrays. Bottom, sero-reactivity to the tiling fragments of the overlapping region of MBP isoform 1 and 2. S922 reacted with T-2.3 and S100 reacted with T2.2. This matched the results on arrays with deletion mutants.

DETAILED DESCRIPTION OF THE INVENTION

Only a few citrullinated antigens have been reported and validated in RA, such as fibrinogen (Fib), enolase-1 (ENO1) and vimentin (VIM). The identification of additional citrullinated antigens and elucidation of their immunodominant epitopes will help develop more sensitive diagnostic assays, and by comparing citrullinated antigens, the assays will achieve higher disease specificity. Therefore, there is a need for identifying more citrullinated antigens to improve performance of current diagnostic tests and also identification of more antigens should help to elucidate the cause of RA.

Citrullinated proteins can trigger host immune responses and elicit antibodies against them in rheumatoid arthritis. The identification of the disclosed autoantibodies (AAbs) and their corresponding antigens impact our knowledge of immunity, leading to early diagnostics and benefitting immunotherapy. Thus, the embodiments herein relate to ACPA’s and novel complexes of ACPA’s with antigen, as well as their application in diagnostics, research, and treatment.

For example, embodiments herein relate to a composition that includes an in vitro antigen/auto antibody complex associated with rheumatoid arthritis, wherein the complex includes an antigen selected from the group consisting of one or more of Myelin Basic Protein (MBP), osteopontin (SPP1), flap endonuclease (FEN1), insulin like growth factor binding protein 6 (IGFBP6), insulin like growth factor 1 (IGF1), and stanniocalcin-2 (STC2).

Moreover, embodiments herein relate to a composition that includes an in vitro antigen/auto antibody complex associated with rheumatoid arthritis, wherein the complex includes reactive epitopes of citrullinated MBP selected from the group consisting of MBP N134-C158 (HGSKYLATASTM-DHARHGFIPHPRTDGLDSDG) (SEQ ID NO. 1) and MBP N144-C176 (DTGILDSGRFFGGIDRGAPKRGSGKVSSEE) (SEQ ID NO. 2).

Further embodiments relate to a method of detecting rheumatoid arthritis in a patient, including the steps of contacting a patient sample known or suspected to contain antibodies with a substrate including one or more of an antigen selected from the group consisting of Myelin Basic Protein (MBP), osteopontin (SPP1), flap endonuclease (FEN1), insulin like growth factor binding protein 6 (IGFBP6), insulin like growth factor 1 (IGF1), and stanniocalcin-2 (STC2); and detecting an antibody/antigen complex with one or more of the aforesaid antigens.

Additional embodiments are directed to a method of differentiating rheumatoid arthritis in a patient, including the steps of contacting a patient sample known or suspected to contain antibodies with a substrate including one or more of an antigen selected from the group consisting of Myelin Basic Protein (MBP), osteopontin (SPP1), flap endonuclease (FEN1), insulin like growth factor binding protein 6 (IGFBP6), insulin like growth factor 1 (IGF1), and stanniocalcin-2 (STC2); and detecting an antibody/antigen complex with one or more of the aforesaid antigens.

Previous studies were mostly performed on western blots using either serum samples or cell lysates followed by separation by either one dimensional (1D) or 2D gel electrophoresis followed by the identification/characterization of reactive protein bands/spots using mass spectrometry. To further confirm the presence of antibodies against a citrullinated protein or a citrullinated epitope on a protein, immunoassays were performed against in vitro citrullinated recombinant proteins using PAB or synthetic peptides with arginine substituted by citrulline.

This approach has proven to be effective, and has the advantage that it examines proteins from natural cell lysates. However, highly abundant proteins in such lysates can obscure the detection of low abundance proteins, identifying the relevant protein in a reactive spot is cumbersome because multiple proteins are often present in the same location, and interpretation of gels can be complicated because multiple isoforms of the same protein may migrate differently. While all these technical challenges can be overcome, the level of effort needed to thoroughly analyze even a single sample prevents this approach from use in large clinical sample sets.

An alternative strategy is the development of high-throughput, unbiased methods to test and identify different candidate antigens in rheumatoid arthritis. Protein microarrays were developed as a method to suit the need of studying antigens in an unbiased high-throughput way.

Embodiments herein are described that relate to high-throughput protein arrays where all proteins are modified simultaneously, probed with samples from rheumatoid arthritis (CCP+, CCP−) and control to identify citrullinated proteins against which there are antibodies present in samples (FIG. 1). We started with a screen for citrullinated proteins specific antibodies in patients with rheumatoid
arthritis using a contra capture nucleic acid programmable protein arrays (NAPPA) displaying 190 full length proteins. Candidate citrullinated proteins were further assessed and validated using a blinded ELISA in an independent set of cases and controls which were age, gender and race matched.

[0028] The 6 antigens that we have identified as ones which get citrullinated and have antibodies against them in rheumatoid arthritis with potential of improved sensitivity of current diagnostics assays are (Table 1): MBP, SPP1, FEN1, IGFBP6, IGF1 AND STC2. The identified candidates achieved 86% sensitivity in CCP+ and 22% sensitivity in CCP− at a 95% specificity.

[0029] We also used our platform to identify immuno dominant epitope(s) for citrullinated antigen MBP. C-terminal deletion mutants for both myelin basic protein (MBP) isoforms were designed based on the following rule: if two arginines were separated by fewer than 9 amino acids, they were treated as a unit; if two arginines were separated by more than 9 amino acids, the amino acids in the middle was used to divide the two arginines into two mutants.

**TABLE 1**

<table>
<thead>
<tr>
<th>Antigen</th>
<th>CCP+ (n = 50)</th>
<th>CCP− (n = 50)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MBP</td>
<td>82%</td>
<td>22%</td>
</tr>
<tr>
<td>SPP1</td>
<td>50%</td>
<td>12%</td>
</tr>
<tr>
<td>FEN1</td>
<td>42%</td>
<td>10%</td>
</tr>
<tr>
<td>IGFBP6</td>
<td>56%</td>
<td>18%</td>
</tr>
<tr>
<td>IGF1</td>
<td>52%</td>
<td>12%</td>
</tr>
<tr>
<td>STC2</td>
<td>66%</td>
<td>12%</td>
</tr>
</tbody>
</table>

[0030] Using an immuno-proteomics approach, we profiled antibody responses against specific citrullinated proteins in CCP+, CCP− rheumatoid arthritis patients. The performance of the candidates identified using protein arrays were confirmed by ELISA using an expanded sample set including subjects from CCP+(n=50), CCP−(n=50), control (n=50) and RA related diseases such as Relapsing Polychon-

tritis (RP) (n=50) and Inflammatory Bowel Disease (IBD) (n=25). Citrullinated MBP was assayed against all 5 serum groups with sensitivities varying from 4% to 82% at 95% specificity. We did not observe significant anti-cit-MBP antibody reactivity in RP and IBD.

[0031] The identified candidates highlight the need for identifying more individual antigens in rheumatoid arthritis as it is yet to be understood what causes rheumatoid arthritis. More importantly we believe that the global profiling of ACPAs against antigens in the human proteome and the mapping of their immunodominant epitopes will provide a more comprehensive picture of ACPA responses in RA. The clinical characteristics of RA are heterogeneous, which may be in part due to the specificity of the immune response directed against post translationally modified antigens. To our knowledge, this is one of the first studies that citrullinated all proteins simultaneously to identify specific citrullinated antigens in a high throughput unbiased way. To ensure accurate estimations of responses when analyzing ELISA results, we also estimated the background associated with the supporting reagent for each plasma sample, which provides the most rigorous assay in similar studies.

[0032] Two isoforms of MBP were used for epitope mapping. Isoform 1 (UniProt: P02686-2) represents the N-terminal 197 AAs (N1-C197) and isoform 2 (UniProt: P02686-6) the C-terminal 160 AAs (N134-C293) of the full length MBP, with a partial overlap between them (MBP 134-197). We observed two representative immune response patterns to these MBP deletion mutants. In pattern 1, a response was always detected for polypeptides including MBP N144-C176 and in pattern 2, response was observed for polypeptide MBP N168-C198.

[0033] In view of the above, one point of novelty is the identification of the 6 autoantigens against which antibodies were detected as candidates in rheumatoid arthritis (Table 1).

**TABLE 2**

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Discovery</th>
<th>Validation</th>
<th>IC</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Subjects</td>
<td>12</td>
<td>50</td>
<td>50</td>
<td>0.87 (one-way ANOVA)</td>
</tr>
<tr>
<td>Mean age Sex</td>
<td>60.08 (31-83)</td>
<td>62.5 (34-89)</td>
<td>34.1 (22-57)</td>
<td>52.22 (18-82)</td>
</tr>
<tr>
<td>Male</td>
<td>3</td>
<td>0</td>
<td>7</td>
<td>7</td>
</tr>
<tr>
<td>Female</td>
<td>9</td>
<td>8</td>
<td>43</td>
<td>43</td>
</tr>
<tr>
<td>No data</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Race</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>White</td>
<td>10</td>
<td>8</td>
<td>49</td>
<td>48</td>
</tr>
<tr>
<td>Asian</td>
<td>2</td>
<td>0</td>
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<td>0</td>
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<tr>
<td>Indian</td>
<td>0</td>
<td>0</td>
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<td>0</td>
</tr>
<tr>
<td>Other</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>No data</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
</tr>
</tbody>
</table>

Many of them have not been previously associated with rheumatoid arthritis. In addition, we also were able to identify certain samples as CCP+ which were initially classified as CCP−. Upon rechecking after our analysis the samples initially classified as CCP− using the commercial tests were categorized again as CCP+. Another point of novelty is the method by which the C-terminal mutants for
both myelin basic protein were produced and mapping of reactive epitopes in rheumatoid arthritis which has not been done before.

[0034] Currently, the diagnostic sensitivities for anti-CCP assay in RA patients range from 66-79% at 86-95% specificity and there are no high-throughput methods to modify hundreds of protein simultaneously and analyze them. Moreover, the antigens and complexes of antigen with antibody described herein are believed to be novel in the context of rheumatoid arthritis detections and patient differentiation, for example, differentiating RA from osteoarthritis and other conditions.

NON-LIMITING EXAMPLES

Characteristics of Plasma Samples

[0035] A total of 30 serum samples were initially obtained from the Benaroya Research Institute, for discovery study. This set consisted of three different groups (CCP+RA (n=10), CCP− RA (n=10) and Healthy controls (n=10)) used for the study on protein array and initial testing on ELISA (Table 2). For the validation study a total of 150 samples (CCP+(n=50), CCP− (n=50) and Healthy controls (n=50)) were obtained and these samples were age, sex and race matched (Table 2).

[0036] The patients with rheumatoid arthritis serum samples were obtained from the clinics at the Benaroya Research Institute after receipt of informed patient consent at Benaroya Research Institute under protocols approved by the Institutional Review Board at Arizona State University and Benaroya Research Institute. Regular controls were used in the study and all samples used in the validation study were matched.

[0037] In the discovery sample set for protein array experiment, 10 samples per group (CCP+, CCP− and control) were selected. However after experiments two CCP− samples were reclassified as CCP+ and changed the original intended sample numbers per group. For validation purpose, additional 150 patient serum was used with 100 from rheumatoid arthritis patients and 50 controls which were matched.

Protein Array Experiments

[0038] Open reading frames were obtained from DNASTAR (dnastr.org). All genes of interest were cloned in the nucleic acid-programmable protein array (NAPPA) compatible expression vectors, pJFT7_nHALO or pJFT7_eHALO. Plasmid DNA was prepared and mixed with NAPPA printing buffer prior to printing as previously described. Protein arrays were produced using the following method: plasmid DNA encoding proteins with a HaloTag are printed in polydimethylsiloxane (PDMS) microwells using a piezo-electric printer. PDMS microwell substrates were blocked with superblock (Thermo Scientific) for 15 minutes at room temperature with gentle rocking and rinsed thoroughly with deionized water. After the application of in vitro protein expression lysates (Human in vitro transcription and translation kit—Thermo Scientific) into the PDMS wells, the wells were covered with a HaloTag ligand (Promega) coated hydrogel slides (SCIOITT). The hydrogel slide and the PDMS microwell substrate were held together by placing them between the plates of the hybridization chamber DT-1001 (Die-Tech) clamped together using screws from DT-2002 (Die-Tech). The substitution of screws helped us to accommodate the PDMS microwell substrate and capture slide setup (~3.5 mm) better between the hybridization plates. This setup was then placed at 30 °C for 3 hrs and proteins expressed in the wells were covalently immobilized on the HaloTag ligand coated hydrogel slides through the HaloTag on each protein. After the capture of expressed proteins, the cover slide are used for subsequent PTM. PTM was followed by blocking with 5% milk in phosphate buffered saline with Tween 20 (milk-PBST), slides were incubated with serum samples (1:500) for 16 h at 4 °C, followed by 3 times wash with 5% milk-PBST. Then slides were incubated with Alexa Fluor 647 labeled Goat α Human IgG (Invitrogen) at 23 °C for 1 h. Slides were then washed, dried and scanned by Tecan scanner under consistent settings.

Protein Array Image Analysis and Quantification

[0039] Arrays were scanned using Tecan PowerScanner and intensity data were quantified using the ArrayPro image analysis software (MediaCybernetics). Local background subtracted median intensity for replicate spots were used for the analysis. Spots with obvious defects were excluded and corresponding spots from replicate arrays were used for analysis instead. GraphPad Prism software was used for generating jitter plots and un-paired t-test analysis. Heat maps of sample reactivity to proteins were generated using MeV software and a hierarchical cluster analysis using Pearson correlation was performed on the log transformed data.

Candidate Selection

[0040] Protein antigens were selected for subsequent ELISA confirmation based on the reactivity that was observed on arrays. The criteria’s for selection are: 1) They showed reactivity to serum samples in their citrullinated forms while there was absence of reactivity to their native forms and the same gene was observed for more than 1 sample; 2) If the reactivity was stronger to citrullinated forms of protein compared to the reactivity to native proteins and if the same pattern was observed in more than 1 sample. Totally, 6 protein antigens were selected.

Epitope Mapping

[0041] Deletion mutants for myelin basic protein (MBP) isoforms was designed based on the following rule: if two arginines were separated by fewer than 9 amino acids. These rules were designed so that enough numbers of native amino acids surrounding arginines was maintained in our deletion mutants to facilitate their recognition after citrullination by serum antibodies. This also limited the total number of deletion mutants to be assayed for this preliminary study. Based on these rules, primers for each mutant were designed (Integrated DNA Technologies) and mutant genes were PCR amplified and cloned into pJFT7_nHALO expression vector using Gateway recombinational cloning. In total, eight deletion mutants were constructed for each MBP isoforms.

ELISA Assays

[0042] ELISA assays were performed to verify selected Ab responses towards citrullinated protein antigens using freshly produced human proteins as previously described. We adapted RAPID ELISA to allow citrullination of target
antigen before assessing its sero-reactivity. HaloTag ligand coated 96 well plates (Promega) were pre-blocked overnight with superblock (Thermo Scientific). Proteins were expressed using hela lysate in vitro transcription-translation system (Thermo Scientific). Protein expression mixture was added into the plates and incubated for 1 hr at room temperature with shaking at 500 RPM to allow the antigen to bind to the HaloTag ligand in each well. Plates were washed and covalently bound antigens were citrullinated by rmPAD2 at 55°C for 3 hrs. Citrullinated antigens were incubated with diluted serum samples (1:1000) overnight at 4°C followed by the addition of Horseradish peroxidase conjugated goat anti human IgG (Jackson ImmunoResearch) with shaking at 500 RPM. The plates were developed with the TMB substrate (Thermo Scientific) for 20 minutes and stopped by addition of 2M sulfuric acid. Absorbance at 450 nm was read on Perkin Elmer EnVision Multilabel Reader.

Statistics and Data Analysis

[0043] To identify candidates we looked for either: 1) citrullinated protein reactivity to antibodies in the serum compared to no reactivity to their native forms or 2) if reactivity to citrullinated proteins was stronger than reactivity to native proteins in more than one sample.

[0044] For all spots identified, local background subtracted median intensity for replicate spots were used for the analysis. Spots with obvious defects were excluded and corresponding spots from replicate arrays were used for analysis instead.

[0045] A heatmap was developed to display differential Ab responses of the 30 sera to all 190 citrullinated proteins (FIG. 2). A heatmap was also generated to show the 6 selected targets in 150 rheumatoid arthritis patients and controls using the confirmation ELISA results (FIG. 3). The heatmap color was scaled according to strength of the reactivity.

[0046] We categorized subjects as Ab responders from ELISA analysis of each antigen if there was a significant difference between CCP+ vs control and CCP+ vs CCP-. We performed t-test analysis for the groups to verify differences and validate.

Identification of Candidate Antigens Associated with Rheumatoid Arthritis

[0047] To identify rheumatoid arthritis associated candidate Abs, we first performed comprehensive profiling of antibodies against 190 full-length human proteins in serum samples from 20 rheumatoid arthritis and 10 control patients on contra capture NAPA. Based on the array data, we selected 11 antigens (SP1, FEN1, IGFBP6, IGF1, STC2, cyclin-A1 (CCNA1), calumenin (CALU), t-actin-capping protein subunit alpha-1 (CAPZA1), protein S100-A11 (S100A11), peroxiredoxin-2 (PRDX2) and glutamate decarboxylase 2 (GAD65)) that showed high antibody reactivity against their citrullinated form in several RA patients and assessed the antibody reactivity of their citrullinated form in all thirty clinical samples by ELISA (FIG. 4 and FIG. 5). Of these, SP1 is a protein selected based on its high expression in synovial fluid in patients with erosive RA. To our knowledge it is not known whether SP1 is citrullinated in RA. Antibodies against citrullinated SP1 showed the best differential reactivity between RA patients and healthy controls.

Validation of Candidate Antigens Associated with Rheumatoid Arthritis

[0048] To validate these novel ACPA, we performed a blinded study of responses in ELISA to SPPI, FEN1, IGFBP6, IGF1 and STC2 in an independent set of 150 serum samples from groups of subjects with three different clinical characteristics: CCP+RA patients (50); CCP−RA patients (50); and healthy controls (50) (Table 2). We observed significant differences for the antigens’ reactivity between the CCP+RA and healthy control groups (FIG. 6). At 95% specificity, the sensitivity of predicting RA with the new ACPAs was 50%, 42%, 56%, 52% and 66% in CCP+RA patients for SPPI, FEN1, IGFBP6, IGF1 and STC2, respectively (Table 1). Surprisingly, response levels to these five ACPAs were also higher in a number of CCP−RA patients, who were clinically categorized as not responsive to standard citrullinated antigens. The sensitivities in the CCP−RA patients were 12%, 10%, 18%, 12%, and 12%, respectively (Table 1). We also assayed anti-cit-MBP antibodies in these samples by ELISA. At 95% specificity, the sensitivity for cit-MBP was 82% for CCP+ serum and 22% for CCP− serum samples.

Epitope Mapping

[0049] It is often useful to determine which epitopes are recognized by antibodies in an immune response. For epitope mapping, ELISA is usually performed on overlapping peptides. This requires the synthesis of multiple native and modified peptides and is a costly process. Recognizing that our platform can manipulate epitopes at the DNA level, followed by display and citrullination of the peptides expressed in vitro, we performed epitope mapping to identify immune-dominant citrullinated epitopes for MBP in RA.

[0050] Epitope mapping of MBP was performed as it is a poorly studied citrullinated antigen for RA. Two isoforms of MBP were used for epitope mapping. Isoform 1 (UniProt: P02686-2) represents the N-terminal 197 AAs (N1-C197) and isoform 2 (UniProt: P02686-6) the C-terminal 160 AAs (N134-C293) of the full length MBP, with a partial overlap between them (MBP 134-197; FIG. 7). C-terminal serial deletion mutants of these two MBP isoforms were constructed, expressed, citrullinated, and probed with MBP reactive serum samples on arrays.

[0051] We observed two representative immune response patterns to these MBP deletion mutants. Within each pattern, the reactivity to isoform 1 and isoform 2 deletion mutants was consistent in the overlapping regions (FIG. 7). In pattern 1, a response was always detected for polypeptides including MBP N144-C176 with the corresponding sequence HGSKYLAATASTMTHIRGLFPRHRDTGLDSIG (SEQ ID NO. 1) and in pattern 2, reactivity was observed for polypeptides MBP N168-C198, which includes the sequence DTGLDSIGRFDDGDRGAPKRDSGKVSEEE (SEQ ID NO. 2). To pinpoint the reactive epitope in the region N131-C205, we constructed tiling fragments of MBP T (2.1) N134-C158, T (2.2) N144-C176, T (2.3) N168-C198, and assayed antibody reactivity using all serum samples. We found several serum samples reacting to region MBP N168-C198 and the rest to region MBP N131-C205 (Suppl. FIG. 5 FIG. 8). No reactivity was seen on other deletion mutants of either isoform lacking this sequence. Thus we were able to use this platform to rapidly identify two distinct B cell epitopes in citrullinated MBP.
Using an immuno-proteomics approach, we applied serum samples to arrays displaying hundreds of citrullinated proteins in parallel. We were able to detect specific antibody reactivity to known citrullinated proteins in RA patients. We provided the first glimpse of reactivity patterns of 20 RA patients against 190 individual citrullinated antigens. We observed two clusters of samples clinically classified as CCP+ and CCP−; additionally, each sample seemed to have its own unique reactivity pattern (Fig. 2).

We discovered and validated six novel ACPAs in RA patients against antigens MBP, SPP1, FEN1, IGFBP6, IGF1 and STC2. Antibodies against citrullinated SPP1 are particularly interesting because SPP1, also known as osteopontin, is a candidate marker elevated in synovial fluid of patients with erosive RA, though it has never been shown to be a citrullinated antigen. We observed immune reactivity specific to citrullinated SPP1 in 50% of CCP+ patients and 12% of CCP− patients at a specificity of 95% in our blinded validation study. SPP1 is recognized as a potential proinflammatory cytokine associated with inflammatory processes. Its discovery as a specific antibody target in RA, from among almost 200 candidates, suggests the possibility of a role that it or the immune response to it might play in the disease.

Anti-cit-MBP has the highest prevalence among the ACPAs that we assayed in this study. Citrullinated MBP has been reported in other diseases such as multiple sclerosis (MS) and other demyelinating diseases and is believed to be involved in the pathogenesis of CNS autoimmune diseases. Here, we detected anti-cit-MBP antibodies in a large fraction of RA patients (~80% of CCP+ samples), which agrees with a previous report on the expression of MBP in the synovial lining and the presence of anti-cit-MBP in RA. In conjunction with the fact that the prevalence to anti-cit-MBP (82%) was much higher than the other known or novel citrullinated antigens (40-60%) in our sample set, it seems likely that these two citrullinated epitopes on MBP, MBP T (2.1) N134-C158 (HGSKYLA6TMDHARHGFLPRHRTGDLSGIS) (SEQ ID NO. 1), T (2.2) N144-C176 (DTGILDSSGRFFGDRGPKRGRSGKVSSEE) (SEQ ID NO. 2) may represent novel immunodominant epitopes in RA, whose biological implications warrant further investigations.

We also calculated the sensitivities and specificities of all 6 candidate antigens that went into the validation study. The physiological relevance of these novel ACPAs warrants future investigation. More importantly, we believe that the global profiling of ACPAs against antigens in the human protome and the mapping of their immunodominant epitopes will provide a more comprehensive picture of ACPA responses in RA. The clinical characteristics of RA are heterogeneous, which may be in part due to the specificity of the immune response directed against post translationally modified antigens.

Assessing reactivity to immune-dominant antigens/epitopes in longitudinal samples collected from high-risk subjects or in patients with different clinical parameters (such as environmental exposures) will promote a better understanding of RA pathogenesis and stratification of RA patients into subtypes based on their response signatures, which based on this initial study may impact the CCP− RA population as it is now defined.

One of the strengths of this study is that protein arrays display many proteins, which make them suitable to screen thousands of proteins against multiple samples for discovery studies. A key expectation is that they will discover new candidate targets which will need to be validated in many more samples. To address the need to test a small number of candidate antigens against many samples (hundreds to thousands), we adapted this method to produce an ELISA assay that supports protein modification and is compatible with biomarker validation/verification studies. This method dovetails with arrays by exploiting the same plasmid clones, without the need to transfer the gene, and it avoids the need to optimize and execute the expression and purification of recombinant proteins as needed by conventional ELISA. We created a system to citrullinate multiple proteins simultaneously and screen them without bias to identify novel antigens in RA. By increasing the number of proteins used for screening, our chances of identifying novel candidates increase accordingly. By performing an unbiased scan we identified MBP where prevalence to anti-cit-MBP (82%) was much higher than the other known and established citrullinated antigens.

To evaluate these Abs against citrullinated antigens performance, we used more clinically relevant ELISA assays in large sample sets, and performed independent blind validation. Our results were also consistent with results obtained on protein arrays.

In summary, we have developed a platform that can assay reactivity to many post-translationally modified antigens in parallel. We screened 190 proteins, and utilized miPAD2 to determine reactivity in RA sera to known and novel citrullinated protein epitopes. We observed unique antibody reactivity patterns in both clinical anti-cyclic citrullinated peptide assay positive (CCP+) and negative (CCP−) RA patients.

At individual antigen levels, we detected antibodies against known citrullinated autoantigens and discovered and validated six novel antibodies against specific citrullinated antigens: Myelin Basic Proteins (MBP), osteopontin (SPP1), flap endonuclease (FEN1), insulin like growth factor binding protein 6 (IGFBP6), insulin like growth factor 1 (IGF1) and stanniocalcin-2 (STC2) in RA patients. We also identified immune-dominant epitope(s) for citrullinated antigen MBP in RA samples—MBP, MBP T (2.1) N134-C158 (HGSKYLA6TMDHARHGFLPRHRTGDLSGIS) (SEQ ID NO. 1), T (2.2) N144-C176 (DTGILDSSGRFFGDRGPKRGRSGKVSSEE) (SEQ ID NO. 2).

The following claims are not meant to be limited to the particular embodiments and examples herein.
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What is claimed is:
1. A composition, comprising an in vitro antigen/autoantibody complex associated with rheumatoid arthritis, wherein said complex includes an antigen selected from the group consisting of one or more of Myelin Basic Protein (MBP), osteopontin (SPP1), flap endonuclease (FEN1), insulin like growth factor binding protein 6 (IGFBP6), insulin like growth factor 1 (IGF1), and stanniocalcin-2 (STC2).

2. The composition of claim 1, wherein said antigen is Myelin Basic Protein.

3. A composition, comprising an in vitro antigen/auto antibody complex associated with rheumatoid arthritis, wherein said complex includes reactive epitopes of citrullinated MBP selected from one or more of the sequences MBP N134-C158 (HGSKYLA STMDIARHGeF LPRHRDTGILDSIG) (SEQ ID NO. 1) and MBP N144-C176 (DTGILDSIGRFFG GDRGAPKR GS GKVSSSE) (SEQ ID NO. 2).

4. A method of detecting rheumatoid arthritis in a patient, comprising the steps of contacting a patient sample known or suspected to contain antibodies with a substrate comprising one or more of an antigen selected from the group consisting of Myelin Basic Protein (MBP), osteopontin (SPP1), flap endonuclease (FEN1), insulin like growth factor binding protein 6 (IGFBP6), insulin like growth factor 1 (IGF1), and stanniocalcin-2 (STC2); and detecting an antibody/antigen complex with one or more of said antigens.

5. The method of claim 4, further including contacting a patient sample known or suspected to contain antibodies with a substrate comprising one or more of citrullinated MBP selected from one or more of the sequences MBP N134-C158 (HGSKYLA STMDIARHGeF LPRHRDTGILDSIG) (SEQ ID NO. 1) and MBP N144-C176 (DTGILDSIGRFFG GDRGAPKR GS GKVSSSE) (SEQ ID NO. 2).

6. The method of claim 4, further comprising treating a patient positive for RA with a therapeutic agent and detecting a quantity of antibody/antigen complex to determine if said quantity of complex increases, decreases, or remains the same.

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