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Chapter 2

Circulating plasmablasts / plasmacells as a source of anti-citrullinated protein antibodies in patients with rheumatoid arthritis

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Abstract

Objectives: To study the characteristics and phenotype of anti-citrullinated protein antibody (ACPA) specific B cells in peripheral blood of patients with rheumatoid arthritis (RA).

Methods: Peripheral blood B cells from ACPA-positive RA patients were cultured with or without stimulating factors. Following culture, supernatants were assessed for the presence of ACPA-IgG and non-specific total IgG by ELISA.

Results: Following stimulation, ACPA were detectable in up to 100% of culture wells. Of interest, ACPA were also produced spontaneously by unstimulated PBMC. In both cases, the average ACPA titre per culture well correlated with ACPA serum titres. No ACPA production was detectable in B cell cultures from ACPA-negative RA patients or healthy controls. Importantly, FACS-sorting experiments located spontaneous ACPA production to the CD20 negative B cell population corresponding to circulating plasmablasts/cells.

Conclusions: ACPA specific peripheral blood B cells are not confined to the CD20 positive memory pool, as circulating plasmablasts/cells spontaneously producing ACPA are also readily detectable. The latter points to an ongoing B cell immune response against citrullinated proteins and contrasts conventional immune responses against, for example, vaccines, where antigen-specific plasmablasts appear in peripheral blood only shortly after vaccination. These circulating, ACPA specific plasmablasts/cells might represent targets for novel therapeutic interventions.
Introduction
In many autoimmune diseases, including rheumatoid arthritis (RA), autoantibodies are believed to be the causative agent that drives disease pathogenesis. Therefore, novel therapeutic approaches that target autoantibody producing B cells require a thorough understanding of their nature, phenotype and disease specific localization.

Several autoantibodies have been described in RA. Among those, anti-citrullinated protein antibodies (ACPA) exhibit the highest specificity for the disease, predict disease onset and severity, and identify a subgroup of patients eligible for more aggressive treatment.[1-3] Citrullinated antigens are present in the inflamed joints of RA patients, supporting the concept of ACPA specific effects at the site of inflammation.[4] Indeed, recent evidence suggests a role for ACPA in disease pathogenesis.[5-8]

So far, little is known on the characteristics of ACPA producing B cells. A single study detected ACPA production in cultures of synovial fluid mononuclear cells.[9] Another study identified ACPA production by peripheral blood B cells upon stimulation.[10] However, it remained unclear whether ACPA detected in this latter study originated primarily from stimulated, otherwise resting memory B cells, or from circulating, differentiated plasmablasts/cells (PB/PC).

In anti-vaccine immune responses, antigen-specific PB transiently appear in peripheral blood in the week after vaccination. These cells rapidly disappear from the circulation and are thought to home to the bone marrow to become long-lived PC that provide high antibody serum titres for years.[11, 12] Like antibody titres against recall antigens, ACPA serum titres in RA patients are relatively stable and can persist at high titre. Nevertheless, it has been postulated that these high ACPA serum titres in RA patients are due to a persistently active immune response against citrullinated proteins, as suggested by the continuous presence of IgM ACPA in some ACPA-positive patients.[13] If this would be correct, one could predict the presence of ACPA-secreting B cells in peripheral blood. Here, we set out to define ACPA-producing B cells in more detail to increase our understanding of anti-citrullinated protein immunity.

Patients and Methods

Patients and healthy individuals
Peripheral blood and serum samples were obtained from patients with ACPA-positive and ACPA-negative RA visiting the outpatient clinic of the Department of Rheumatology at Leiden University Medical Centre, Leiden, The Netherlands. Patients fulfilled the 1987 criteria for RA at the time of diagnosis and gave written informed consent for sample acquisition. Treatment included disease-modifying anti-rheumatic drugs, biological agents, and glucocorticoids. Permission for conduct of the study was obtained from the ethical review board of Leiden University Medical Centre.
Isolation of PBMC, B cells and B cell subsets

Mononuclear cells were isolated from peripheral blood using Ficoll-Paque gradient centrifugation (LUMC pharmacy). CD19-positive B cells were isolated by magnetic bead-based positive selection using Dynabeads and DETACHaBEAD CD19 (Invitrogen) according to the manufacturer’s instructions.

For selected experiments, B cells were sorted into subsets on a BD FACS Aria flow cytometer. PBMC were surface stained with CD3 Alexa Fluor 700 (clone UCHT1), CD14 Pacific Blue (clone M5E2), CD19 APC-Cy7 (clone SJ25C1), CD20 PerCP (clone L27), and CD27 PE-Cy7 (clone M-T271, all BD Biosciences). Subpopulations were defined as CD3\(^{-}\)CD14\(^{-}\)CD19\(^{+}\)CD20\(^{-}\)CD27\(^{++}\) PB/PC, CD3\(^{-}\)CD14\(^{-}\)CD19\(^{+}\)CD20\(^{+}\)CD27\(^{-}\) naïve and CD3\(^{-}\)CD14\(^{-}\)CD19\(^{+}\)CD20\(^{+}\)CD27\(^{-}\) memory B cells (figure 2.3A).\(^{[14, 15]}\)

Culture conditions

B cells were cultured in Iscove’s Modified Dulbecco’s Medium (IMDM) supplemented with 10% heat-inactivated fetal calf serum (FCS), penicillin/streptomycin (100U/ml), and 2mM Glutamax in 96-wells flat bottom plates at a density of 2\(^{\times}\)10\(^{4}\) cells/well on a layer of irradiated (7000 rad) mouse fibroblast cells stably transfected with human CD40 ligand (CD40L, 5\(^{\times}\)10\(^{3}\) cells/well) in the presence of B cell activating factor (BAFF) (Miltenyi,100ng/ml), IL21 (Invitrogen, 50ng/ml) and anti-IgM F(ab\(^{\prime}\))\(_{2}\)-fragments (JacksonImmunoResearch Laboratories, 5µg/ml). Supernatants were harvested after 6-7 days.

To assess spontaneous ACPA production, PBMC were cultured at a density of 2\(^{\times}\)10\(^{5}\) cells/well in IMDM supplemented with 10% heat-inactivated FCS, penicillin/streptomycin (100U/ml), and 2mM Glutamax with no additional stimuli. FACS-sorted B cell subsets were cultured under similar conditions, with only variation in cell density (2\(^{\times}\)10\(^{4}\) cells/well for memory/naïve populations; 1500–8000 cells/well for PB/PC) and culture plate (96-well round-bottom plates). Supernatants were collected at day 6 and 13.

Measurement of IgG-ACPA and total IgG

Serum and culture supernatants were assessed for the presence of IgG-ACPA by ELISA based on reactivity against the CCP2-peptide (Euro Diagnostica). Serum samples were tested at a 1:50 dilution or higher according to the manufacturer’s instructions, culture supernatants were tested undiluted. The commercial standard was diluted to assess the lowest concentration at which the standard curve was consistently linear (0.125U/ml (supplementary figure 2.S1)). This value takes into account that the concentration of the commercial standard relates to a 1:50 dilution of serum samples, requiring a division by 50 of the manufacturer’s units. A representative set of patients (25%) was tested, and found negative, for reactivity against the arginine containing control peptide.

The presence of total IgG in culture supernatants was assessed by standard ELISA (Bethyl Laboratories).
Figure 2.1: ACPA production upon in vitro B cell stimulation. CD19-positive B cells were isolated and cultured for 6-7 days. Supernatants were assessed for the presence of IgG-ACPA by ELISA. Each dot represents a culture well, results are grouped per patient. Values below linearity of the standard (0.125U/ml) were calculated by extrapolating the standard curve linear through zero (supplementary figure 2.S1). Displayed are results from (A) ACPA-positive RA patients. On average, we obtained 38 (6-170) wells per patient, with a mean ACPA titre of 0.45U/ml (0.0-3.1)) at the end of the cultures, (B) ACPA-negative RA patients, n=3; average of 45 (16-96) wells per experiment, (C) Healthy donors, n=5; average of 51 (32-73) wells per experiment. (D) Shows the correlation between ACPA serum titres and mean ACPA titres produced in culture.
Statistical analysis
Analysis was performed using GraphPad Prism 5.01. Correlations were assessed as non-parametric correlation. p Values <0.05 were considered significant.

Results

ACPA production upon in vitro B cell stimulation
To assess the general presence of ACPA-producing B cells in peripheral blood, we stimulated freshly isolated CD19+ B cells of ACPA-positive RA patients (n=30) as described previously.[16, 17] After 6 to 7 days, ACPA were detected in 80% of the cultures and, in some patients, in up to 100% of the culture wells (figure 2.1A). No ACPA were detected using B cells isolated from healthy individuals and ACPA-negative RA patients (figure 2.1B-C), although total IgG production was comparable (not shown). Of interest, mean ACPA titres obtained in the cultures correlated with ACPA serum titres (figure 2.1D). Based on the number of B cells per culture well, we estimate that, in some patients, at least one in every $2\times10^4$ B cells can produce ACPA.

Spontaneous ACPA production
If the ACPA-response would result from an ongoing immune response, we reasoned that naïve or memory B cells should be continuously activated resulting in the presence of IgG-ACPA producing PB/PC in peripheral blood. Therefore, we analysed whether B cells spontaneously producing ACPA could be observed. To this end, peripheral blood mononuclear (PBMC) cells were cultured without exogenous stimuli. Interestingly, comparable amounts of ACPA-positive culture wells were detected in most patients when comparing stimulated versus

Figure 2.2: Spontaneous versus stimulated ACPA production. (A) PBMC of ACPA-positive RA patients were split in two and cultured either with or without stimulation at a density of $2\times10^5$ cells/well. Supernatants were assessed for the content of IgG-ACPA at day 13 of culture. (B) Shows the correlation between ACPA serum titres and mean ACPA titres spontaneously produced in culture.
non-stimulated PBMC per patient (figure 2.2A), whereas non-specific IgG production was strongly increased upon stimulation (up to 100 fold, data not shown). Importantly, also mean titres of spontaneous ex-vivo ACPA production correlated with ACPA serum titres (figure 2.2B). This suggests that ACPA detected in the culture wells originate primarily from circulating PB/PC.

Figure 2.3: Circulating plasmablasts/cells as primary source of ex-vivo ACPA production. CD19-positive B cell subpopulations were FACS-sorted based on the expression of surface markers CD20 and CD27 into CD20^+CD27^- naïve B cells, CD20^-CD27^+ memory B cells and CD20^+CD27^+ plasmablasts/cells (PB/PC) (A). In addition, the markers were used to deplete PBMC of the respective B cell subset. Sorted B cell populations and PBMC depleted of subsets were cultured without stimulation (B). Representative data of one patient. Each dot corresponds to one culture well. (C) Combined data of 5 patients. Each dot represents one donor. The lowest point of linearity of the standard was used as cut-off for ACPA positivity. (D) Naïve and memory B cells were FACS-sorted as described above for four patients. Both populations were stimulated as described in the Methods section. Supernatants were harvested after 6 days of culture and assessed for ACPA by ELISA. Data are depicted as units corrected for culture volume.
Circulating plasmablasts/cells as primary source of ACPA ex vivo

To confirm the presence of ACPA-producing PB/PC in peripheral blood, and to exclude the possibility of endogenous stimulation of memory B cells by antigen presenting cells within the PBMC population, we next isolated PB/PC by FACS sort (see figure 2.3A for gating strategy). ACPA production by purified PB/PC was compared to ACPA production of total PBMC, PBMC depleted of PB/PC, and to PBMC depleted of the CD20 positive B cell fraction (figure 2.3B-C). In line with our hypothesis, isolated PB/PC readily produced ACPA ex-vivo. The total PBMC fraction and PBMC depleted of CD20-positive cells both showed comparable ACPA production, whereas ACPA production by PBMC depleted of PB/PC was considerably diminished.

ACPA were not produced by unstimulated, FACS sorted memory B cells (not shown), but were detectable upon stimulation. This indicates the additional presence of CD20-positive, ACPA-specific memory cells in the circulation (figure 2.3D). ACPA were hardly detected in wells containing naïve B cells, irrespective of stimulation.

Together, these data highlight the presence of circulating, ACPA-producing PB/PC in peripheral blood of RA patients and indicate that depletion of the CD20-positive B cell compartment does not eliminate these cells.

Discussion

The immune response against citrullinated antigens in RA is incompletely understood. Specifically, characteristics of ACPA-producing B cells have scarcely been studied. Here, we show that ACPA-producing PB/PC circulate in peripheral blood of patients with RA.

In the steady state, most circulating PB/PC in healthy individuals are thought to originate from mucosal immune responses.[11] Upon vaccination, antigen-specific PB appear in the circulation only transiently. Therefore, our findings provide evidence for the recent and continuous activation of ACPA-producing B cells in RA patients. This observation is relevant, as recently activated PB/PC are phenotypically different from fully differentiated, long-lived plasma cells or memory B cells. Indeed, depletion of the CD20-positive B cell compartment did not eliminate the spontaneous ACPA production observed.

Next to circulating PB/PC, we identified ACPA-specific memory B cells that produced ACPA upon stimulation. The latter observation could be taken as a surprise as it is conceivable that citrullinated protein-directed memory B cells are constantly driven to the PB/PC population as a consequence of the persistent and systemic presence of citrullinated antigens. Novel therapeutic approaches aimed at inhibiting ACPA production should thus target the CD20-positive B cell subset, but need to also include the PB/PC compartment.

Finally, we found an association between ACPA production in culture and ACPA serum titres. This suggests that ACPA serum titres originate partly from tissue resident, long-lived plasma cells producing rather stable ACPA “background” titres, and partly from recently generated, circulating plasmablasts/cells. The latter reflect the active part of the immune response and are thus more variable in their contribution to the ACPA serum titre, as suggested by our study.
In line with this, intensive treatment was previously shown to be able to decrease ACPA serum titres by up to 50%. [18-20], suggesting indeed a fluctuation in ACPA titres due to an ongoing immune response.

In conclusion, we here identify ACPA-specific plasmablasts/cells circulating in peripheral blood of ACPA positive RA patients. This population, which is not targeted by CD20 depletion, reflects the active part of the ACPA-specific B cell response and could represent a relevant target for therapeutic intervention.

References


Supplementary figures

Figure 2.S1: Standard curve determining the linearity of the IgG-ACPA ELISA. To determine the linearity in the lower range of the IgG-ACPA ELISA the standard was extended using a 1:1 dilution series. The concentration of 0.125U/ml IgG-ACPA was identified as the lowest point that is consistently within the linear part of the standard curve.