The p68 autoantigen characteristic of rheumatoid arthritis is reactive with carbohydrate epitope specific autoantibodies

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Abstract

Objective—The autoantigen p68 is a target of autoantibodies as well as autoreactive T cells with a high specificity in rheumatoid arthritis (RA). The binding characteristics of the autoantibodies to their antigen were now analysed biochemically and cytologically.

Methods—Deglycosylation techniques as well as lectin and sugar competition experiments were performed to p68 to discover if the antibodies detected a glycoepitope. Its antigenicity was investigated applying anti-p68 antibodies derived from RA patients in comparison with polyclonal rabbit anti-p68 antibodies.

Results—p68 specific antibodies from RA patients did not bind to p68 that had been deglycosylated by alkaline β-elimination, O-glycosidase or periodate treatment. In contrast, binding of p68 specific antibodies raised in rabbit was unaffected by either deglycosylation protocol. Furthermore, lectins specific for the carbohydrate N-acetylgalactosamine competed with p68 specific antibodies from RA patients for antigen binding. N-acetylgalactosamine by itself also competed with patient derived anti-p68 antibodies for p68 binding. Again, rabbit anti-p68 antibodies did not elicit these competitive effects. Applying cytoimmunofluorescence, p68 was present in the cytoplasm or endoplasmic reticulum and also in low abundance on the cell surface. Under heatshock conditions, p68 was detectable in the nucleus.

Conclusions—Autoimmunity to p68 during RA is carried by anti-carbohydrate autoantibodies. The carbohydrate modification of p68 appears to be N-acetylgalactosamine, which may reflect the regulation of intracellular localisation of the antigen. It is hypothesised that a shift in glycosylation pattern accompanied by an unphysiological localisation of the antigen could trigger antigenicity of p68 during the pathogenesis of RA.

Rheumatoid arthritis (RA) is the most frequent systemic rheumatic disease. The aetiology and pathogenesis of the disease, however, are not well understood and the diagnosis, especially in the early stages of the disease, is often difficult. The rheumatoid factor (RF), by definition a human anti-IgG autoantibody, is the sole serological parameter included in a catalogue of criteria for the diagnosis of RA. It is present in approximately 70% of RA patients, but also in up to 20% of patients with other systemic disorders; it even occurs in apparently healthy people in whom the RF assay is adjusted so that up to 5% of them are positive. The RF in RA patients is often an agalactosylated IgG terminating in N-acetyl glucosamine (N-GlcNAc), rather than galactose or sialic acid. β-1,4-galactosyltransferase that catalyses the addition of galactose to the oligosaccharide chains on this molecule has been shown to undergo a variety of normal and disease associated changes. The latter may contribute to the pathological processes in RA. Monoclonal rheumatoid factors have been described that detect the agalactosylated IgG better than the normal form. For mouse anti-collagen II agalactosyl-IgG, but not for the normally glycosylated antibodies, pathogenicity in collagen induced arthritis (CIA) has been described. An IgG preparation for passive transfer of collagen induced arthritis proved more effective when first digested with β-galactosidase to increase the level of the agalactosyl IgG glycoform. It is thus evident that the glycosylation status of autoantibodies is of importance in the autoimmune response.

Previously, anti-p68 antibodies have been described by our group in 66% of RA patients, but not in healthy controls or patients with other autoimmune diseases as a possible aid for an improved diagnosis of the disease. Furthermore, the impact of these antibodies in the pathogenesis of RA could be underlined by the demonstration of p68 specific T cells. They were detected in 67% of RA patients. Family specific anti-HLA DP, DQ and DR antibodies were capable of increasing stimulation of the p68 specific response of some RA patients when added to T cell proliferation assays. This was regarded indicative of the presence of class II restricted p68 specific suppressive T cells otherwise inhibiting the proliferation of p68 specific helper T cells. The p68 antigen shows an isoelectric point of 5.1 and a relative molecular mass of 68 000. The antigen that had originally been identified in the synovium and is probably ubiquitously expressed in humans, is detectable also in mouse, but not in insect cells. To further analyse p68, affinity purified RA patient derived anti-p68 antibodies were applied for immunoscreening of various human cDNA expression vector libraries. We analysed why these autoantibodies failed to detect specific clones in these experiments. One possible explanation was that the
patients’ antibodies were directed to a carbohydrate epitope.

Various examples of anti-carbohydrate antibodies can be given, especially when occurring in certain disease states. Thus, in rats monoclonal antibodies have been described as being specific for carbohydrate epitopes on *Trichinella spiralis*, which are protective against the parasite by causing rapid expulsion. In chronic Chagas’ disease, anti-α-galactosyl antibodies are found to be lytic to *Trypanosoma cruzi*. The smallest reactive residue is O-linked Galα1–3Galβ1–4GlcNAc. The carbohydrate reagents are reactive with Chagas’, but not with healthy anti-Gal antibodies, indicating that *T. cruzi* O-linked oligosaccharides are highly immunogenic under infectious conditions. The unusual finding here is that O-linked GlcNAc is reported with further glycosidic modifications for the first time. In humans suffering from diseases like RA, Crohn’s disease, tuberculosis, sarcoidosis or erythema nodosum leprosum, autoantibodies reactive with O-GlcNAc have been described. The possibility is discussed that the anti-carbohydrate immune responses could also evoke cross reacting anti-peptide autoantibodies.

In this study we have demonstrated that the patient antibodies were directed to a carbohydrate epitope that is very probably not expressed in prokaryotic expression systems (*E. coli*). p68 was purified using patient antibodies to raise a specific antiserum in rabbits. This p68 specific rabbit antiserum is compared with the human antiserum with respect to carbohydrate epitope binding.

**Methods**

**ANTIBODIES**

Serum samples containing p68 specific antibodies were obtained from RA patients attending our rheumatology unit.

HeLa proteins were separated by SDS-PAGE as described. Immunisation was performed injecting the neutralised p68 containing polyacrylamide matrix (approximately 1 µg of p68 as estimated by Bradford quantification). For the first boost, two dimensionally purified p68 antigen contained in the neutralised and ground poly-acrylamide matrix (as described) was applied. For the second boost, ConA binding fraction was separated on SDS-PAGE, and the eluted material from the p68 band was injected. Antibodies obtained by this protocol were directed against protein epitopes and could thus function as controls in the various deglycosylation experiments described below.

The secondary antibodies were FITC conjugated IgG sheep anti-human or sheep anti-rabbit (all classes each) antibodies from Wellcome, England and used because they demonstrated the lowest number of cross reactions. Incubation was for three hours at room temperature at a 1:100 dilution as recommended. Immuno reactions were visualised at 492 nm wave length.

**ENZYMES**

All enzymes were purchased from Boehringer/Mannheim, Germany. O-glycosidase from *Diplococcus pneumoniae* is described as an O-glycopeptide-endo-D-galactose-GalNAc-hydrolase. Neuraminidase from *Arthrobacter ureafaciens* catalyses the hydrolysis of terminal N- or O-acetyl neuraminic acids in α2–3, α2–6 or α2–8 bonds in, for example, glycoproteins. N-glycosidase F from *Flavobacterium meningoseptum* is a glucan-asperaginase-amidase and cleaves off all forms of asparagine-bound N-glycans from glycolipids or N-proteins.

**PATIENTS**

Serum samples of nine RA patients fulfilling the ACR criteria were obtained after consent. All samples were anti-p68 positive, two of them were rheumatoid factor negative.

**LECTINS**

Lectins from the DIG Glycan Differentiation Kit, Boehringer/Mannheim, Germany or from Sigma. GNA (Galanthus nivalis agglutinin) were used and directed to terminal Man α(1–3)-, α(1–6)- or α(1–2)-linked to Man; SNA (Sambucus nigra agglutinin), directed to sialic acid (SA) α(2–6) linked to Gal; MAA (Maackia amurensis agglutinin), directed to α(2–3) linked to Gal; PNA (peanut agglutinin), directed to Gal β(1–3)GalNAc without further substituents; DSA (*Datura stramonium* agglutinin), directed to Gal β(1–4) linked to GlcNAc and GlcNAc. Sigma: ConA (*Concanavalin A*), directed to α-D-mannopyranoside, α-DL-glucopyranoside, GlcNAc, α/β-D-arabinofuranoside, α/β-fructofuranoside and also small hydrophobic molecules; DBA (*Dolichus biflorus* agglutinin), directed to α-GalNAc; RCA1 (*Ricinus communis* agglutinin), directed to β-linked Gal; UEA-I (*Ulex europaeus* agglutinin), directed to α-L-fucose; WGA (wheat germ agglutinin), directed to GlcNAc, β(1–4) linked oligomers of GlcNAc and SA.

Protein preparation was performed as described elsewhere.

**LECTIN BINDING AND COMPETITION EXPERIMENTS**

Lectin binding was performed applying the DIG (digoxigenin) glycan differentiation kit, which was performed as described in the product literature. Lectin competition was carried out with the same concentrations of lectin as suggested for binding studies. Lectins were then preincubated on western blots for one hour at room temperature. Subsequently, antibodies were incubated overnight. Immunoreactivity was tested with FITC conjugated secondary antibodies from Wellcome, Germany, and visualised at 492 nm.

**CARBOHYDRATE COMPETITION**

Carbohydrates were preincubated with antibodies for three hours at room temperature. Sugar concentrations were 1 M, 100 mM, and 10 mM. Carbohydrates remained present during incubation of antibodies with blotted protein material overnight.
ALKALINE β-ELIMINATION

Total HeLa protein (40 µg) or purified p68 or blot strips of SDS-PAGE separations of total proteins were incubated at 37°C for 16 hours in either 4 ml water or 5 mM or 10 mM solutions of sodium hydroxide. Samples were then adjusted to pH 7.1, precipitated and subjected to SDS-PAGE and immunoblotting. Blot strips were directly incubated with RA or control antibodies. Detection of the immunoreaction was carried out with FITC conjugated secondary antibodies.

ENZYMATIC DEGLYCOSYLATION

HeLa proteins were separated by SDS-PAGE and blotted as described.8 Blot strips were blocked in a buffer containing 1.1% NP40 and 2% BSA and incubated for 15 minutes at 75°C in 1 ml of a 1% SDS solution. Subsequently, a 10-fold excess of NP-40 was adjusted with an NP-40 buffer and again incubated for 15 minutes at 75°C. This step is mandatory for enzyme activity, which would be destroyed in the presence of a vast excess of SDS. After cooling down the solution to 37°C, 5 U O-glycosidase and 40 µU of neuraminidase or 1 U of N-glycosidase F were added and incubated for 16 hours at 37°C. Then, after a washing step, immunoreaction was performed.

PERIODATE TREATMENT

Blot strips of total HeLa protein were blocked in the above described buffer and incubated with 10 mM sodium periodate and 100 mM sodium acetate for two hours at room temperature at pH 5.2 in the dark. Controls were incubated in sodium acetate only.

CYTOMUNOFLUORESCENCE

HeLa cells were grown on microscope slides overnight at 37°C, eventually heatshocked for one hour at 42°C, and denatured in 3.5% paraformaldehyde and 0.15% NP40. Primary (anti-p68) and secondary antibodies were incubated for one hour at room temperature.

Results

DEGLYCOSYLATION EXPERIMENTS

Based on the observation that RA anti-p68 antibodies failed to detect clones in immunoscreening experiments, deglycosylation experiments were performed to consider whether these antibodies detected a glycopeptide on p68. Alkaline β-elimination specifically removes O-linked carbohydrate residues from glycoproteins.9 Either total HeLa proteins, purified p68 or blot strips of HeLa proteins separated by SDS-PAGE were subjected to mild base catalysed β-elimination. Immunoreactivity with the RA antibodies was abolished partially at sodium hydroxide concentrations of 5 mM and completely at concentrations as low as 10 mM (fig 1). Eight other anti-p68 positive RA serum samples exhibited the same pattern of immunoreactivity (data not shown). No correlation to the presence of rheumatoid factors was observed (p > 0.5; Fisher's exact test).

Rabbit control antibodies specific for proteinaceous epitopes of p68 were not affected by either treatment. This proved that incubation at mild hydroxide conditions did not degrade protein epitopes of p68. Also enzymatic cleavage of sugars was investigated applying O-glycosidase and N-glycosidase F, both of which contained no protease activity. In some O-glycosidase experiments, neuraminidase was co-incubated to potentially remove sialic acids and to improve accessibility to potential O-glycosidase cleavage in case of high sialylation. O-glycosidase treatment of blot strips with p68 or solubilised total protein containing p68 abolished immunoreactivity with the RA, but again not with rabbit anti-p68 antibodies (fig 2). Co-incubation of neuraminidase with O-glycosidase had no apparent effect on immunoreactivity (data not shown). N-glycosidase F treatment had no effect on either antibody (fig 2). No gel shift exceeding 1–2 k was visible when deglycosylated p68 protein was subjected to SDS-PAGE (10%, 12 cm length).

Figure 1 Alkaline β-elimination. HeLa protein western blot. Lanes 1 and 2 preincubated in water, 3–5 in 5 mM NaOH, 6–7 in 10 mM NaOH. Subsequent incubation with anti-p68-positive RA serum (2, 4, 6), anti-p68-negative human control serum (5), anti-p68-positive rabbit serum (1, 3, 7) and anti-p68-negative control serum. Detection with FITC conjugated secondary antibody. p68 was detected by the RA serum antibody after incubation with water (2) and with 5 mM NaOH (4), but not after incubation with 10 mM NaOH (6). p68 was detectable by the rabbit serum antibody even after 10 mM NaOH treatment (7).

Figure 2 O-glycosidase. HeLa protein western blot. Lanes 1 and 2 preincubated with water, 3+4 with N-glycosidase F, 5–6 with O-glycosidase. Subsequent incubation with anti-p68-positive RA serum (2, 4, 6) or anti-p68-positive rabbit serum. Detection with FITC conjugated secondary antibody. p68 was detectable by the RA serum antibody after incubation with water and with N-glycosidase F and not after incubation with O-glycosidase (5). p68 was detectable by the rabbit serum antibody even after treatment with O-glycosidase (6).
CARBOHYDRATE OXIDATION

General oxidation and thereby destruction of potential carbohydrate epitopes at concentrations of 10 mM periodate also abrogated p68 reactivity with RA, but not with rabbit antibodies (fig 3).

LECTIN BINDING AND COMPETITION EXPERIMENTS

To further analyse the nature of the glycosylation of p68, various lectins were tested for antigen binding. WGA, DSA, and ConA did bind to the antigen, while GNA, SNA, MAA, PNA, DBA, RCAI, and UEA-I did not. In competition experiments the first three lectins were able to compete with the antibody for antigen binding. After initial incubation of the antigen with lectin, immunoreactivity was abrogated with the human, but not with the rabbit p68-specific serum (fig 4). This did not apply to the other lectins, which again had no effect. HeLa proteins were separated by a ConA sepharose column, applying a lectin that is known to bind also N-acetylglucosamine (GlcNAc). p68 could be detected in the eluate but not in the flow through fraction (fig 5).

CARBOHYDRATE COMPETITION EXPERIMENTS

Finally, we considered whether a single sugar could compete with p68 for binding of RA antibodies. GlcNAc and (glucosamine) GlcN competed at high (1 M) but not at low (0.1 M) sugar concentrations, while all other carbohydrates tested (methyl-α-D-
of RA specific autoantibodies and T
Autoantigen p68 has been detected as a target
detectable in low amounts on the cell surface (fig 8). Furthermore, the antigen was also
42°C the antigen was detected in the nucleus
deglycosylation of p68 that its modification is
This is confirmed by the finding that only car-
binding site of an antibody (the paratope) and
Ac as well as GlcN could compete with the antigen for binding. This in
sugar is a critical determinant. This also applies
to the anti-GlcNAc antibodies described by
and generally consists of approximately 15 amino
anti-GlcNAc antibodies, for instance, is not
A protein epitope is the area of an antigen that is covered by the antigen
binding site of an antibody (the paratope) and
generally consists of approximately 15 amino acids. We thus conclude that the RA anti-p68
antibodies also recognise proteinaceous or other carbohydrate parts of p68, or both, but that the
sugar is a critical determinant. This also applies
to the anti-GlcNAc antibodies described by
Rook and Shikhman. Anti-p68 antibodies
derived from RA patients seem to recognise a
mixed carbohydrate-protein-epitope of p68.
This is confirmed by the finding that only car-
biochemical properties from 1 M on were
able of competition. These experiments
were performed on western blots and only
semi-quantitation was possible. It can be
estimated from the undetectable gel shift after
deglycosylation of p68 that its modification is
no larger than five sugar residues. The
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 carbohydrates.
The shift of compartments of the p68 antigen upon heat shock implies a regulatory role of mono-O-GlcNAc residues comparable to that seen for many proteins such as the small heat-shock protein α-B crystallin.21 Ser-mono-O-GlcNAc versus Ser-phosphate are known as a switch for various molecules with respect to activation or localisation of these proteins.21 Discussing a modification of p68 with mono-O-GlcNAc though, is in conflict with the cleavage specificity of O-glycosidase for Ser/Thr-GaINAc—GalNAc and could only be explained by a widened specificity of the O-glycosidase, which also detects and cleaves Ser/Thr-mono-O-GlcNAc bonds. To analyse the carbohydrate composition and sequence, complete sugar release of highly purified p68 antigen and subsequent HPLC analysis of the carbohydrate will be necessary.

The fact that peptides can mimic carbohydrate epitopes or vice versa, especially in autoimmune conditions,22,23 may fuel the idea of molecular mimicry in the pathogenesis of autoimmune disorders such as RA. Especially, when crossreactivity with autoantibodies occurs that are directed to cytoskeletal proteins24 and keratin.25 Interestingly, anti-keratin antibodies could be induced when immunising BALB/c mice with GlcNAc-conjugated BSA but not with BSA alone. Nevertheless, for the RA anti-p68-carbohydrate epitope antibodies no mimicking or crossreactive peptide epitope is known so far, and we were unable to detect such peptide stretches by immunoscreening with autoantibodies.

In this study it is shown that the RA patient anti-p68 antibodies are directed against a carbohydrate epitope. Although rabbits were immunised with native and thus glycosylated p68—antigen, antibodies were derived that reacted with proteinaceous parts of the antigen only. One possible explanation is that antibody production against the carbohydrate epitope of p68 may be harmful to the animal and is abrogated by B cell tolerance. The generation of anti-carbohydrate antibodies to p68 may be of significance for the pathomechanism of the disease. In this respect, variations in β-1,4 galactosyltransferase are believed to contribute to pathogenesis in rheumatic diseases and a hypothesis is suggested, whereby the disease is associated with the dysregulation of an integrated glycosylation network, comprising galactosylation and anti-carbohydrate specific antibodies.24 Agalactosylated antigens are known to be associated with autoimmune disorders: agalactosyl IgG is frequently found in RA, Crohn’s disease, tuberculosis, sarcoidosis, and erythema nodosum leprosum.25-27 Immunising mice with a defined genetic background will help in answering the question whether or not p68 can be pathogenic in an animal system. Further study is needed to consider whether anti-GlcNAc antibodies directed against p68 have any relation to those that are directed to IgG.