Characterisation of the rat oesophagus epithelium antigens defined by the so-called ‘antikeratin antibodies’, specific for rheumatoid arthritis*

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Abstract

Objectives—An attempt was made to characterise the antigens recognised by serum IgG antibodies directed to the stratum corneum of rat oesophagus epithelium, the so-called ‘antikeratin antibodies’, which were shown to be highly specific for rheumatoid arthritis (RA) and thus to have an actual diagnostic value.

Methods—Immunoblotting was performed with RA serum samples on different extracts of rat oesophagus epithelium separated by various monodimensional and two dimensional electrophoreses.

Results—Three low-salt-soluble antigens sensitive to proteinase K and, therefore, of protein nature were identified. Two proteins, with apparent molecular masses of 210 and 120–90 kilodaltons, shared isoelectric points ranging from 5-8 to 8-5; the third protein exhibited isoelectric points from 4-5 to 7-2 while its molecular mass ranged from 130 to 60 kilodaltons. Immunoadsorption of RA serum samples onto cyto-keratins extracted from the stratum corneum of rat oesophagus epithelium did not change their immunoreactivity towards the three antigens. As well as de-glycosylation and dephosphorylation methods failed to modify either the electrophoretic migration of the proteins or their immunoreactivity with RA serum samples.

Conclusion—The so-called ‘antikeratin antibodies’ do not react with cyto-keratins. They specifically recognise three late epithelial differentiation proteins which had not been previously described. These proteins may be related to (pro)filaggrin.


Serum samples from patients with rheumatoid arthritis (RA) show a wide variety of circulating autoantibodies, such as rheumatoid factors,1 2 antinuclear antibodies,3 4 and autoantibodies to tissue antigens, such as perinuclear granules5 6 or collagen.7 8 This study deals with the circulating IgG antibodies, described in 1979 by Young et al,9 which label the stratum corneum (SC) of rat oesophagus epithelium when tested by indirect immunofluorescence (IF). Over the past few years numerous authors have shown that these antibodies are specific for RA and thus useful for its diagnosis.10–19 Previously we used semi-quantitative IIF, on a large series of serum samples, to determine the diagnostic specificity and sensitivity of detection of these antibodies (99% and 41-5%, respectively).18 Their presence in the RA-serum samples was found to be independent of age, sex, and disease duration but to be related significantly to disease severity and activity (see 18 for cross references). Moreover, they were recently shown to be present in early RA and even before the onset of clinical symptoms.20 21 Today, these antibodies are the most reliable diagnostic and prognostic serological marker for RA.18 Lastly, they have also been detected in rheumatoid synovial fluids19 22 23 with higher concentrations than in the related serum samples,23 suggesting that they might be locally synthesised and possibly play a part in the pathogenesis of RA synovitis.

Probably because cyto-keratins constitute the major proteins of the SC, these antibodies, which produce a linear laminated labelling restricted to this epithelial layer only, were called ‘antikeratin’ antibodies without any previous biochemical characterisation of the antigens. Confirming data obtained on short series by other authors,10 22 24 we have shown that these antibodies also label the SC of human epidermis and are thus genuine autoantibodies.17 On the other hand, naturally occurring IgM and IgG autoantibodies directed to epidermal cyto-keratins have been described in all normal human serum samples, as well as in serum from patients with various diseases, including RA.17 25 26 The simultaneous investigation, on a large series of RA serum samples, of these natural IgG autoantibodies to epidermal cyto-keratins and of the antibodies to SC specific for RA, allowed us to show that they were directed to different epitopes, probably borne by different molecules.17

In this study we present the first isolation, physicochemical and biochemical characterisation of the rat oesophagus epithelial antigens recognised by the antibodies to SC specific for RA, the so-called ‘antikeratin antibodies’.
Materials and methods
HUMAN SERUM SAMPLES
Serum samples were obtained from healthy donors and from patients with RA according to the clinical, radiographic, and biological criteria of the American Rheumatism Association.27 The samples were assayed by semi-quantitative IIF for the presence of IgG antibodies to the SC of rat oesophagus epithelium, as previously described.18 Briefly, the titre-like value was estimated by eye on a semi-quantitative scale from 0 to 4 (0-25 unit steps) independently by two readers who had not been informed of the clinical context. The results were summed giving titre-like values ranging from 0 to 8.

PREPARATION OF ANTIGENS FROM RAT OESOPHAGUS EPITHELIUM
Specimens of frozen oesophagus from 6-week-old male Wistar rats were obtained from Iffa-Credo (Lyon, France) and stored at –80°C until use. After removing the outside muscularis by gentle scraping the oesophagus was incubated at 55°C for five minutes in phosphate buffered saline, pH 7.4, containing 5 mM EDTA and 0.4 mM phenylmethylsulphonyl fluoride, then chilled for five minutes in the same ice cold buffer without EDTA. The epithelium was mechanically separated from the underlying connective tissue then pooled and ground at 4°C with a tissue homogeniser in a low salt, 40 mM TRIS-HCl buffer, pH 7.5 containing 150 mM NaCl, 10 mM EDTA, 0.1% sodium azide, and 0.1 mM phenylmethylsulphonyl fluoride (0-25 ml buffer per epithelium). The insoluble material was removed by centrifugation at 25 000 g for 15 minutes and the supernatant stored at –30°C. The protein concentration of the epithelial extract was estimated using chicken egg albumin as standard by either absorbance measurement at 280 nm or by the Bio-Rad Protein Assay (Bio-Rad Laboratories, Richmond, CA).

PREPARATION OF CYTOKERATINS FROM RAT OESOPHAGUS EPITHELIUM
Sequential extraction of epithelial proteins was performed according to Sun and Green.28 The epithelial sheets were extracted three times in 20 mM TRIS-HCl, pH 7.4, then the insoluble pellet was extracted three times with the same buffer containing 8 M urea (solubilisation of the cytokeratins from the viable layers). Lastly, the pellet was extracted three times with the same buffer containing 8 M urea plus 0-1 M β-mercaptoethanol (solubilisation of the cross linked cytokeratins from the SC). The TRIS-urea-β-mercaptoethanol-soluble fractions were pooled, dialysed against water, and lyophilised.

ELECTROPHORESIS
Polyacrylamide gel electrophoresis (PAGE) was performed with a PhastSystem (Pharmacia-LKB, Uppsala, Sweden) using precast gels and following the procedures suggested by the manufacturer. Sodium dodecyl sulphate (SDS)-PAGE was performed either with 7.5% homogeneous or with 8–25% gradient PhastGels, non-denaturing PAGE (native-PAGE) with 8–25% gradient PhastGels, and isoelectrofocusing (IEF) using PhastGels with ampholytes generating either a 3–9 or a 5–8 pH gradient. The high molecular weight standard protein kit from Biorad was used in SDS-PAGE and the broad range isoelectric calibration kit from Pharmacia was used for IEF. For native-PAGE the following proteins (Pharmacia-LKB) were used as markers: horse spleen ferritin (440 kilodaltons), beef liver catalase (232 kilodaltons), beef heart lactate dehydrogenase (140 kilodaltons), and bovine serum albumin (67 kilodaltons). Epithelial proteins were concentrated by lyophilisation or precipitation either with four volumes of ethanol for 30 minutes at –30°C or with 10% trichloroacetic acid for 15 minutes at 0°C. For SDS-PAGE the precipitate was dissolved in 10 mM TRIS-HCl, pH 7.4, 2% SDS, 0.01% bromophenol blue, and 1% β-mercaptoethanol (SDS sample buffer), and boiled for five minutes. The lyophilised SC cytokeratins were dissolved and boiled in 20 mM TRIS-HCl, pH 7.4, containing 6 M urea, 4% SDS, 1% β-mercaptoethanol, and 0.01% bromophenol blue. For native-PAGE and IEF the precipitate was dissolved in distilled water with 0.01% bromophenol blue. For two dimensional electrophoresis one lane of the first dimensional gel (IEF or native-PAGE) was equilibrated either in 112 mM TRIS-HCl, pH 6.5, containing 112 mM acetic acid, 2.5% SDS, 1% β-mercaptoethanol, and 0.01% bromophenol blue, when the second dimension was SDS-PAGE, or in 200 mM TRIS-HCl, pH 8.8, with 0.01% bromophenol blue, when the second dimension was native-PAGE. Then, the lane was placed horizontally onto the second dimension stacking gel and electrophoresis was run as above.

IMMUNOBLOTTING
After electrophoretic separation the epithelial extracts were electrotransferred to nitrocellulose membranes (Bio-Rad) according to the method of Towbin et al.29 The membranes were blocked for 30 minutes with 10 mM TRIS-HCl, pH 8.0, 150 mM NaCl, containing 0.05% Tween-20 and 0.5% skimmed dry milk (working buffer). Then the membranes were incubated overnight at 4°C in working buffer containing human serum samples diluted to 1:100. After three washing steps in working buffer the membranes were treated for 90 minutes at room temperature with peroxidase conjugated goat antibodies to the human γ heavy chain of immunoglobulins (Southern Biotech Inc, Birmingham, AL) diluted 1:400 in working buffer. After one wash in working buffer and two washes in 10 mM TRIS-HCl, pH 8.0, 150 mM NaCl, the colour reaction was developed by incubation with peroxidase substrates: 0.5 mg/ml 4-chloro-1-naphthol...
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(Bio-Rad) in 0.025% (v/v) hydrogen peroxide.

In some cases the staining intensity of the strips was estimated semiquantitatively exactly as described above for IIF reactivity and a titre-like value was attributed to each serum within a scale from 0 to 8 with steps of 0.25.

IMMUNOADSORPTION
Nitrocellulose membranes were electroblotted either with the low-salt-soluble epithelial extract separated by native-PAGE or with the TRIS-urea-β-mercaptoethanol-soluble SC cytokeratins separated by SDS-PAGE. An RA serum, diluted 1:200 in working buffer, was immunoadsorbed by five successive incubations at 4°C (three times for three hours then twice for 12 hours) with the two types of membranes previously saturated. Unblotted nitrocellulose membranes were used as control.

CHROMATOGRAPHIC TECHNIQUES
Chromatography was performed on prepacked columns using an FPLC System (Pharmacia-LKB). The low-salt-soluble epithelial extract was first applied to a gel filtration column (Superose 12, 30 × 14 cm) and the proteins were eluted in 50 mM sodium phosphate, pH 7.2, 150 mM NaCl, at 0.3 ml/min. Each collected fraction was analysed by native-PAGE and immunoblotting. The fractions of interest were pooled and 1.7 M ammonium sulphate added for further hydrophobic interaction chromatography on a phenyl-Sepereose column (5 × 0.5 cm) previously equilibrated with 50 mM sodium phosphate, pH 7.2, containing 1.7 M ammonium sulphate. Elution was performed at 0.5 ml/min by decreasing the ammonium sulphate concentration in the equilibrium buffer; it was led stepwise to 0.765 M then progressively decreased to 0.36 M before final washing with sodium phosphate buffer. Absorbance was monitored at 280 nm and the fractions were concentrated, then analysed by native-PAGE and immunoblotting.

ENZYMATIC AND CHEMICAL TREATMENT OF THE ANTIGENIC EPITHELIAL EXTRACT

Protease K digestion
The low-salt-soluble epithelial extract (6-5 mg/ml) was precipitated with ethanol and solubilised, concentrated 20-fold, in 50 mM TRIS-HCl, pH 8.0 and 5 mM CaCl2. Aliquots were incubated at 37°C for one hour with either water or an aqueous solution of Trichinella spiralis proteinase K (Boehringer Mannheim, Germany) at a final concentration of 2 U/ml.

Alkaline β elimination
Several conditions of alkaline β elimination of the O-linked carbohydrates were tested using NaOH concentrations from 0.1 to 0.5 mol/l applied during times varying from 0.5 to 20 hours at 20 or 37°C. The lyophilised epithelial extract was incubated either with NaOH or in water. After neutralisation the samples were precipitated with trichloroacetic acid.

N-glycosidase treatment
The lyophilised epithelial extract was treated with N-glycosidase (Genzyme, Boston, MA); samples (8 mg/ml final concentration) were boiled for five minutes in one volume of 2% SDS, 50 mM β-mercaptoethanol, then incubated for 24 hours at 37°C after adding two volumes of a 250 mM potassium phosphate buffer, pH 8.0, containing 15 mM 1,10-phenanthroline, 15% (v/v) methanol, and 1% (v/v) Nonidet P-40 with or without N-glycosidase at a final concentration of 0.03 U/ml. The samples were concentrated by trichloroacetic acid precipitation. The N-glycosidase activity was tested on fetuin calf serum (Sigma, St Louis, MO) using the same conditions.31

Neuraminidase treatment
The lyophilised epithelial extract (2 mg) was incubated for 120 minutes at 37°C with or without neuraminidase from Vibrio cholerae (Boehringer Mannheim) at a final concentration of 1 U/ml in 200 μl of 50 mM sodium acetate buffer, pH 5.5, containing 150 mM NaCl, 9 mM CaCl2, 0.05% sodium azide, and 25 mg/ml human serum albumin. Then the samples were precipitated with ethanol. Human α1 acid glycoprotein (Sigma), digested in the same conditions, allowed the neuraminidase activity to be checked.32

Alkaline phosphatase treatment
The paradoxically acid conditions determined by Harada et al33 for the phosphoprotein phosphatase activity of the bovine intestinal alkaline phosphatase were used. The lyophilised epithelial extract (1.5 mg) was incubated for two hours at 37°C with or without calf intestine alkaline phosphatase (Boehringer Mannheim) at a final concentration of 0.024 U/μg of treated material in 750 μl of 25 mM sodium acetate buffer, pH 5.6, containing 280 mM NaCl, 3 mM MgCl2, 9.5 μM ZnCl2, and 2.8 mM triethanolamine. Then the samples were dialysed and lyophilised. The phosphoprotein phosphatase activity was checked on hen yolk phosphvitin (Sigma) and the method of Cutting and Roth for specific staining of phosphopeptides after SDS-PAGE34 was used to evaluate dephosphorylation.

After digestion with enzymes or chemical treatment the samples of epithelial extract were analysed by monodimensional or two dimensional electrophoresis and by immunoblotting with RA serum samples, as described above.

STATISTICAL ANALYSES
Correlations were sought using Spearman rank correlation coefficients.
Results
IDENTIFICATION OF THE SC ANTIGENS
Various conditions for extraction of the rat oesophagus epithelium antigens were compared using several low salt buffers with or without addition of denaturing or reducing agents (urea, SDS, β-mercaptoethanol) and detergents (Nonidet P-40, Triton X-100). No significant difference in the efficiency of antigen extraction was found between the various conditions; urea and detergents only inducing the additional solubilisation of non-antigenic epithelial proteins. Thus a simple low ionic strength buffer was used for the immunoblotting analysis of a series of RA serum samples (fig 1). After separation of the extract by SDS-PAGE, 16 of the 18 RA serum samples chosen with IIF titre-like values for antibodies to SC ranging from 0-25 to 7 (scale 0 to 8) were immunoreactive with a smear of 90–130 kilodaltons apparent molecular mass, and 14 of the 16 serum samples also reacted with a sharper band of about 210 kilodaltons. With the high titred serum samples, the reactivity spread between these two regions and on down to 60 kilodaltons. After separation of the extract by non-denaturing electrophoresis (native-PAGE), the RA serum samples with high titre-like values of antibodies to SC recognised three antigenic components: the first migrated like the 440 kilodalton protein marker and was referred to as the A antigen, a second migrated like the 232 kilodalton marker (B antigen), and a third migrated between the 140 and 67 kilodalton markers (C antigen). The major reactivity was directed to the B antigen with weaker staining of the A antigen. All serum samples with an IIF titre-like value greater than 2-00 recognised the A antigen and all those with an IIF titre-like value greater than 0-25 recognised the B antigen. The C antigen was detected by some RA serum samples with IIF titre-like values higher than 0-75 and by all RA serum samples with IIF titre-like values higher than 3-00. The staining intensity of the C antigen was not always related to that of the two other antigens.

Despite the difference in the patterns of reactivity obtained after SDS-PAGE and after native-PAGE separation of the extract, both the immunoblotting analyses showed a striking specificity for RA, as healthy donor serum samples showed either no or weak reactivity with the antigenic components (not shown).

For the whole series of RA serum samples a highly significant correlation was observed between the IIF titre-like value of antibodies to SC and the staining intensity of the blots semi-quantitatively estimated ($r=0.96$, $p<10^{-4}$; antigens separated by SDS-PAGE; $r=0.94$, $p<10^{-4}$; antigens separated by native-PAGE). In the same way, the titre-like values of the serum samples in the two immunoblotting assays were highly significantly correlated ($r=0.98$, $p<10^{-4}$).

Although the solubility and the apparent molecular masses of the antigens specific for RA were different from those of the rat oesophagus epithelium cytokeratins, we performed immunoadsorption experiments to assess the antigenic independence of the two types of epithelial molecules. Certain RA serum samples, highly reactive with the three, A, B, and C, epithelial antigens and also reactive with the rat oesophagus SC cytokeratins when assayed by immunoblotting, were immunoadsorbed either onto epithelial antigens or onto cytokeratins, then analysed by immunoblotting. The immunoadsorption of one serum onto SC cytokeratins (fig 2) did not change its reactivity to antigen A, B, or C, whereas it completely abolised the reactivity to the cytokeratins. Immunoadsorption of the serum onto antigens A, B, and C did not modify its reactivity to the cytokeratins, whereas it substantially reduced its reactivity to all three antigens. When unbblotted nitrocellulose sheets were used as immunosorbent the reactivity of the serum samples remained largely unmodified.

Figure 1 Immunoblotting analysis of low salt extract of oesophagus epithelia with RA serum samples. A rat oesophagus epithelium homogenate was obtained in a low salt buffer. The soluble fraction was precipitated by ethanol and solubilised either in sodium dodecyl sulphate (SDS) sample buffer then separated by 7-5% SDS-PAGE (A), or in water then separated by 8–25% native-PAGE (B). Electroblotting was performed on nitrocellulose membranes which were cut into vertical strips and incubated with human serum samples according to 'Materials and methods'. Lane 0=Coomassie blue staining of the related gel; lane 1=control strip incubated with peroxidase labelled antihuman IgG; lane 2–19=immunodetection using RA serum samples with decreasing IIF titre-like values of antibodies to SC ranging from 7 to 0-25. Migration of protein markers in kilodaltons (kDa) is shown on the left.

Figure 2 Immunoblotting analysis using an RA serum previously immunoadsorbed either on stratum corneum (SC) cytokeratins or on the low-salt-soluble epithelial antigens. The low-salt-soluble antigens, separated by native-PAGE (A), and SC cytokeratins, extracted with TRIS-urea-β-mercaptoethanol and separated by SDS-PAGE (B), were either stained by Coomassie blue (lane 0) or electrobotted onto nitrocellulose membranes. Immunoblotting analysis was performed with an RA serum (lane 1) and with the same serum previously immunoadsorbed either onto unblotted control membranes (lane 2), or onto membranes blotted with SC cytokeratins (lane 3) or with low-salt-soluble antigens (lane 4).
Two dimensional IEF/native-PAGE was used to determine the isoelectric characteristics of the three antigens identified by one dimensional native-PAGE. The molecules with the higher molecular masses, referred to as the A and B antigens, showed isoelectric points ranging roughly from 6 to 7-2 (5-8 to 8-5 with high tittered serum samples, like the 210 and the 120–90 kilodalton antigens separated by IEF/PAGE-SDS). The C antigen had the same isoelectric point distribution as the 130–60 kilodalton comma shaped antigen.

Lastly, two dimensional electrophoresis combining native-PAGE and SDS-PAGE was used to confirm the relations between the antigens defined by native-PAGE and by IEF/PAGE-SDS. The A, B, and C antigens were identified as the 210, the 120–90, and the 130–60 kilodalton antigens respectively. Their large heterogeneity both in apparent molecular mass and in isoelectric point made purification of the A, B, and C antigens somewhat difficult. Gel filtration, anionic exchange, and hydrophobic interaction, used either alone or sequentially, did not allow a complete purification of the molecules (data not shown). During hydrophobic interaction chromatography (fig 5), however, the antigens were sequentially eluted: the C antigen was identified in early fractions, while decreasing salt concentrations induced the desorption of the B and A antigens, which were identified in later fractions. This showed the lower hydrophobicity of the C antigen compared with the A and B antigens, which were similar to each other.

**BIOCHEMICAL CHARACTERISATION OF THE SC ANTIGENS (fig 6)**

Several enzymatic and chemical treatments were performed for the biochemical characterisation of the antigenic components and to determine the biochemical nature of the epitope(s) recognised by the antibodies to SC.

Treatment of low salt extracts with proteinase K resulted in a complete loss of immunoblotting reactivity with the RA serum samples, demonstrating the involvement of protein moliecties in the formation of the three
antigenic molecules. The marked isoelectric and molecular mass heterogeneity of the three antigens suggested the existence of extensive substitutions, such as glycosylation or phosphorylation of the proteins. Deglycosylation of the antigenic proteins was attempted by various methods classically used to remove the most common carbohydrate moieties. Whatever the conditions of treatment with sodium hydroxide, used for the β elimination of O-linked sugars, no modifications of the electrophoretic pattern were obtained before degradation of the peptide chains as seen by Coomassie blue staining. Treatment by N-glycanase, despite its broad specificity for N-deglycosylation, did not produce any substantial modifications of the electrophoretic migration of the antigens, and the intensity of reactivity of the RA serum samples remained unchanged.

The charge heterogeneity of the antigens might be due to a variable quantity of terminally linked sialic acid residues. This hypothesis was explored using Vibrio cholerae neuraminidase, which hydrolyses terminal N- and O-aclyneuraminic acids. This enzymatic treatment did not modify either the migration of the antigens or the reactivity of the serum samples. Varying degrees of phosphorylation of amino acids might also account for the charge heterogeneity. Calf intestine alkaline phosphatase, used under conditions that induced an almost complete dephosphorylation of the phosphoprotein phosvitin, induced no modifications in either the migration or the immunoreactivity of the treated antigens.

Discussion
Since Young et al., 10 nine different groups have used IEF to study the serum IgG antibodies labelling the SC of rat esophagus epithelium in large series of patients with rheumatic diseases and have confirmed their high diagnostic specificity for RA (review in 18). This study provides the first isolation and detailed characterisation of the rat esophagus epithelium antigens targeted by these antibodies. The highly significant

![Figure 5](Image)

Figure 5 Immunoblotting analysis of the hydrophobic interaction chromatography fractions of the low-salt-soluble antigens previously purified by gel filtration chromatography. Gel filtration of the low salt extract was performed as described in 'Materials and methods'. The eluted fractions containing the antigens were pooled and injected into a phenyl-sepharose column (5×0.5 cm) previously equilibrated in 50 mM sodium phosphate, pH 7.2, containing 1.7 M ammonium sulphate. Elution was performed at 0.5 ml/min, by decreasing ammonium sulphate concentration (---) and absorbance was monitored at 280 nm (→). The low-salt-soluble extract (lane 0) and the phenyl-sepharose fractions (lanes 1–10) were concentrated by precipitation with trichloroacetic acid, solubilised in water, separated by 8–25% native-PAGE, and analysed by immunoblotting with an RA serum.

![Figure 6](Image)

Figure 6 Immunoblotting analysis of the low-salt-soluble antigens after enzymatic and chemical treatments. Various enzymatic or chemical treatments, including proteinase K (A) or N-glycanase (B) digestions, mild alkaline β-elimination (C), and neuraminidase (D) or alkaline phosphatase (E) digestion, were applied to the low salt extract of rat esophagus epithelia. Treated samples (*) and untreated samples (–) were separated by one dimensional 7.5% SDS-PAGE (A, B) or two dimensional IEF/SDS-PAGE (that is, 3–9 (C) or 5–8 (D, E) IEF followed by 7.5% (D) or 8–25% (C, E) SDS-PAGE) then simultaneously analysed by immunoblotting using RA serum samples. In the sample treated with alkaline phosphatase, serum reactivity to the enzyme produced additional staining.
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leads to the conclusion that α-2–3, α-2–6, or α-2–8 linked N-acetyl or N-glycosyl neuraminic acids are not extensively represented in the antigens. As varying degrees of phosphorylation might also be a source of charge heterogeneity, we attempted to dephosphorylate the antigens. The absence of effect produced on the antigens shows that phosphorylation is probably not responsible for charge heterogeneity. Taken together, these results suggest that the molecular mass and isoelectric heterogeneity have to be explained in other ways, such as uncommon glycosylation, other classes of sialic acids, phosphate groups with molecular environments different from those of the control phosphoprotein we used, or even other types of modification.

The persistence of protein antigenicity after all the treatments targeting the sugar moieties shows that these sugars are not the major antigenic determinants. Moreover, periodate oxidation of blotted antigenic proteins did not modify their reactivity with RA serum samples (data not shown), indicating that sugar residues with vicinal groups, such as hydroxyl or mono- or disubstituted amines, are not involved in epitope formation. Together with the protein nature of the A, B, and C antigens, and our previous demonstration of predominant distribution in the IgG1 subclass of the antigens, we have already indicated that the absence of effect produced on the antigens shows that phosphorylation is probably not responsible for charge heterogeneity.

Until now, the antibodies to SC which label the rat oesophagus epithelium have been called 'antikeratin' antibodies, probably in reference to the term 'keratin layer' formerly used for the SC and because cytokeratins are the most abundant proteins in this layer. It was shown, however, that labelling of the rat oesophagus SC by an RA serum was not abolished either by a preabsorption of the serum with phosphorylated human epidermal cytokeratins or by the 

By immunoblotting analyses we found, in the low salt extracts of rat oesophagus epithelium, three different antigenic molecules referred to as A, B, and C antigens, separable by two dimensional electrophoresis and identified as proteins by their susceptibility to proteinase K. The absence of modification of the electrophoretic patterns of the antigens after denaturing and reducing treatments showed that the three proteins are not constituted by natural or artefactual associations (neither disulphide nor weak bonds) between each other and confirmed that the antigenic system is actually composed of three distinct proteins. The A and B proteins appear to be strikingly related, however. Firstly, they share physicochemical properties such as isoelectric point distribution and hydrophobicity and, secondly, their immunoreactivity with RA serum samples was found to be closely related. This latter point was statistically confirmed as the staining intensities of the two molecules, analysed with a large panel of RA serum samples, were found to be highly significantly correlated (Gomes-Daudrix V, Sebbag M, et al, in preparation).

These physicochemical and immunological analogies might be explained by a precursor-product relation between the two proteins, the A protein being cleaved to generate the B protein. In contrast, the C protein did not seem to be recognised by all the antibodies which target the A and B proteins. As above, on a large series, its immunoreactivity was also significantly correlated to that of the two other proteins but slightly less so (Gomes-Daudrix V, Sebbag M, et al, in preparation). Nevertheless, these correlations indicate that the existence of epitopes shared by the A, B, and C proteins is highly probable.

To explain both the extensive charge heterogeneity and the poor resolution of the antigens after SDS-PAGE the existence of post-translational modifications of the protein moieties was investigated. Protein glycosylation has been frequently shown to have a microheterogeneous character and charged sugar residues, such as sialic acids or sulphated monosaccharides, could modify the net charge of the protein bearing them. Thus classical and widely used enzymatic and chemical methods were chosen to demonstrate glycosylation of the antigens. The immunoblotting analysis of samples treated by the various methods used never showed any significant change in the electrophoretic pattern of the A, B, or C antigens. The results suggest that the targeted substituting groups—that is, the best characterised types of asparagine linked glycoconjugates and the 'mucin-type' O-linked saccharides—are not the major groups involved in post-translational modifications of the antigens. In the same way the absence of susceptibility to neuraminidase

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38. The molecular masses, the isoelectric points, and the high solubility of the A, B, and C antigens are clearly different from those of rat oesophagus epithelium cytokeratins.
43. The unmodified reactivity of RA serum samples to the A, B, and C antigens after adsorption on the SC cytokeratins extracted from the same tissue clearly indicated that the major epitope(s) recognised on the antigenic proteins are not present on the SC cytokeratins, at least in the form and the conformational state they present on nitrocellulose sheets. Considered as a whole, these results show that the A, B, and C rat oesophagus epitelial antigens recognised by RA serum samples are not cytokeratins, and that the recognised epitope(s) are absent from these molecules in our experimental conditions.
The IIF labelling by RA serum samples of the cornified layer of the rat oesophagus epithelium indicates that the recognised proteins correspond to molecules whose synthesis or processing generates epitopes during the late stages of keratinocyte differentiation. The physicochemical and biochemical characteristics of the A, B, and C antigens differ from those of all the previously described late differentiation proteins of rat squamous epithelia. We recently identified in human epidermis, however, the targets of the auto-

antibodies to SC to the basic filaggrin and to a neutral to acidic variant of this protein. Filaggrin is an intermediate filament aggregating, histidine rich protein which has been characterised in the epidermis of a variety of species, including murine animals and humans. In rat epidermis its synthesis occurs in the keratinocytes of the granular layer as a heavily phosphorylated 60 kDa kilodalton precursor (profilaggrin), which accumulates in the cytoplasmic keratohyalin granules. This precursor is formed by the tandem repetition of about 20 filaggrin units separated by short linker peptides. During the transition from the granular layer to the SC the serine residues are dephosphorylated and profilaggrin is proteolytically cleaved to produce the basic 45 kilodalton filaggrin with heterogeneous iso-electric points ranging from 9 to 10.7 8 When we applied the various published methods for extraction and purification of epidermal filaggrin to rat oesophageal epithelium, we were unable to detect any protein with physicochemical characteristics identical to those of the mature basic form of rat epidermal filaggrin. The absence of this form of filaggrin has also been reported in the rat palate epithelium, another cornified squamous epithelium, whereas immunohistological analyses allowed (pro)filaggrin epitopes to be detected in the keratohyalin granules and in the SC of this epithelium. Moreover, profilaggrin mRNA was detected in rat oesophageal epithelium in situ hybridisation. Therefore, it may be speculated that profilaggrin synthesis actually does occur in rat oesophageal epithelium but that the differentiation pathway in this epithelium differs from that observed in epidermis, and induces specific profilaggrin maturation processes. Therefore, the A, B, and C antigens, immunologically related to human epidermal filaggrin, and different from rat epidermal filaggrin and profilaggrin, might correspond to forms of (pro)filaggrin specific for the rat oesophageal epithelium.

In conclusion, we characterised the proteins targeted by the antibodies to SC, so-called ‘antikeratin’ antibodies, which have been found to be highly specific for RA and to be related to its most severe and active forms, to be present at its early stages, and even to precede its clinical onset. Identification of the epitope(s) defined by these antibodies on the three rat oesophageus antigens and the search for these epitope(s) molecules of the human synovial membrane may provide new insights into the pathogenesis of RA.

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