MAPPING SURFACE ACCESSIBILITY OF THE C1r/C1s TETRAMER BY CHEMICAL MODIFICATION AND MASS SPECTROMETRY PROVIDES NEW INSIGHTS INTO ASSEMBLY OF THE HUMAN C1 COMPLEX

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Running title: Lysine mapping of C1 interactions by nano-LC-MS/MS analysis

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C1, the complex that triggers the classical pathway of complement, is a 790-kDa assembly resulting from association of a recognition protein C1q with a Ca\(^{2+}\)-dependent tetramer comprising two copies of the proteases C1r and C1s. Early structural investigations have shown that the extended C1r-C1r-C1s-C1s tetramer folds into a compact conformation in C1. Recent site-directed mutagenesis studies have identified the C1q-binding sites in C1r and C1s and led to a three-dimensional model of the C1 complex (Bally, I., Rossi, V., Lunardi, T., Thielens, N. M., Gaboriaud, C., and Arlaud, G. J. (2009) J. Biol. Chem. 284, 19340-19348). In this study, we have used a mass spectrometry-based strategy involving a label-free semi-quantitative analysis of protein samples to gain new structural insights into C1 assembly. Using a stable chemical modification, we have compared the accessibility of the lysine residues in the isolated tetramer and in C1. The labelling data account for 51 of the 73 lysine residues of C1r and C1s. They strongly support the hypothesis that both C1s CUB\(_1\)-EGF-CUB\(_2\) interaction domains, which are distant in the free tetramer, associate with each other in the C1 complex. This analysis also provides the first experimental evidence that, in the proenzyme form of C1, the C1s serine protease domain is partly positioned inside the C1q cone, and yields precise information about its orientation in the complex. These results provide further structural insights into the architecture of the C1 complex, allowing significant improvement of our current C1 model.

Complement is an essential component of innate immunity due to its ability to recognize pathogens and to limit infection in the vertebrate host. In addition, activation of the complement system enhances the migration of phagocytic cells to infected areas and stimulates the adaptive immune response (1,2). The initial steps of the complement cascade involve modular proteases that are activated in a sequential manner via one of three pathways: the classical, lectin and alternative pathways. The classical pathway is triggered by C1, a 790-kDa Ca\(^{2+}\)-dependent complex resulting from the association of a recognition protein C1q and a tetramer comprising two copies of the serine proteases C1r and C1s (3-6). Recognition of targets such as pathogens or immune complexes by the C1q moiety of C1 elicits self-activation of C1r, which in turn converts C1s into its active form. Once activated, C1s specifically cleaves C4 and C2, thereby initiating a series of sequential and highly specific proteolytic reactions leading to the formation of the membrane-attack complex and the elimination of the target. The classical pathway of complement is also involved in immune tolerance due to the ability of C1 to recognize and induce clearance of apoptotic cells, and plays a major role in xenograft rejection (7). The uncontrolled activation of the complement system, however, can result in...
self-tissue damages and pathologic inflammation.

During the last years, the 3-D structure of several fragments of C1r, C1s and C1q has been solved by X-ray crystallography and other biophysical methods (6,8-11). C1r and C1s, and the mannan-binding lectin (MBL)-associated serine proteases of the lectin complement pathway, share the same type of modular organisation (12) with, starting from the N-terminal end, a C1r, C1s, Uegf, bone morphogenetic protein (CUB) module, an epidermal growth factor (EGF)-like module, a second CUB module, two successive complement control protein (CCP) modules, and a chymotrypsin-like serine protease (SP) domain. Whereas the CCP1-CCP2-SP regions of C1r and C1s mediate their enzymatic properties, their N-terminal CUB 1-EGF segments are involved in the Ca\(^{2+}\)-dependent C1r-C1s interactions required for assembly of the C1s-C1r-C1r-C1s tetramer. Available structural data have led to low-resolution models of the C1 complex in which C1s-C1r-C1r-C1s (hereafter named the tetramer) adopts a compact conformation when bound to C1q (13-15). The main ionic interactions between the C1q collagen stems and the tetramer were initially supposed to be mediated by an acidic cluster located in the C1r EGF module (16). Mutagenesis experiments have recently ruled out this hypothesis (17) and led to a refined 3-D model of the C1 complex in which C1s-C1r-C1r-C1s (hereafter named the tetramer) adopts a compact conformation when bound to C1q (13-15). The main ionic interactions between the C1q collagen stems and the tetramer were initially supposed to be mediated by an acidic cluster located in the C1r EGF module (16). Mutagenesis experiments have recently ruled out this hypothesis (17) and led to a refined 3-D model of the C1 complex in which acidic residues involved in the Ca\(^{2+}\)-binding sites of the C1r CUB\(_{1}\) and CUB\(_{2}\) and C1s CUB\(_{1}\) modules interact with the C1q stems. Given the location of these sites, the CUB\(_{2}\)-EGF-CUB\(_{2}\) interaction domains of C1r and C1s are now proposed to be located entirely inside the cone delimited by the six C1q stems, in sharp contrast with the original model.

To gain further information about the assembly and structure of human C1, we have used stable chemical modifications associated with a mass spectrometry-based strategy and a label-free semi-quantitative approach to investigate the changes of surface accessibility taking place in the tetramer upon C1 assembly. Lysine acetylation is one of the most common chemical modifications used to analyse protein complexes (18-23). Since these residues are charged, they are likely to occupy solvent-exposed regions of proteins, which makes them excellent candidates to identify protein/protein interactions. In addition, the relatively large number of lysines (146 in total) present in the tetramer and their distribution provide the opportunity to investigate the effects of C1q binding on the whole tetramer structure. Our data are consistent with the hypothesis that the C1s interaction domains interact with each other in C1, and provide experimental evidence that the C1s catalytic domains are partly located inside the C1q cone, yielding further insights into C1 architecture.

**EXPERIMENTAL PROCEDURES**

**Materials**- All chemicals were of analytical grade. Slide-A-Lyzer 10K molecular weight cut-off dialysis cassettes and sulfo-N-hydroxysuccinimide (sulfo-NHS) acetate were purchased from Pierce. Fibrinogen-binding inhibitor peptide (FBIP, fragment 400-411), Tris(2-carboxyethyl) phosphine hydrochloride (TCEP-HCl), α-cyano-4-hydroxycinnamic acid, sinapinic acid, trifluoroacetic acid, formic acid and porcine pepsin were all obtained from Sigma. Acetonitrile was purchased from VWR. Coated silica PicoTip emitters for the nano-ESI\(^{2}\) source were obtained from New Objective. ZipTip C4 and C18 tips were purchased from Millipore. Ultrapure water was obtained from a Milli-Q System (Millipore).

**Proteins**- C1q and the proenzyme form of the tetramer were isolated from human plasma as described previously (24,25). Prior to use, C1q was dialysed at 4°C against 250 mM NaCl, 4 mM CaCl\(_{2}\), 250 mM Hepes, pH 7.3 using a membrane of 10K molecular weight cut-off (final C1q concentration: 1.6 \(\mu\)M).

**Chemical modification of the lysine residues of C1s-C1r-C1r-C1s**- The overall procedure is outlined in Figure 1. The free tetramer and the C1 complex were labelled in parallel as follows. Solutions containing 15 pmoles of the tetramer were prepared in 16 \(\mu\)L of 250 mM NaCl, 4 mM CaCl\(_{2}\), 250 mM Hepes, pH 7.3 using a membrane of 10K molecular weight cut-off (final C1q concentration: 1.6 \(\mu\)M).
respectively. The number of free lysine residues taken into account was 146 for the tetramer (36 in C1r and 37 in C1s) and 271 for C1, assuming 126 unmodified lysines per C1q molecule (26). The reaction was quenched by decreasing the pH to 3.0 using 4 µL of 1 M TCEP-HCl, pH 2.5, and samples were incubated for a further 10 min at room temperature to reduce disulfide bonds. Prior to protein digestion, each sample was placed on ice for 2 min. Then, 2 µL of a pre-cooled porcine pepsin solution prepared in 100 mM H3PO4, pH 1.8, was added to achieve a protease/protein ratio of 1:1 (w/w). Proteolysis was performed at pH 3.0 for 5 min at 0°C, and stopped by adding 1.5 µL of an 8 M NaOH solution (final pH 8.0). After 1 min, samples were acidified to pH 2.5 with 3 µL of a 50% trifluoroacetic solution, flash-frozen in liquid nitrogen and stored at -80°C. Unlabelled controls were performed in parallel by replacing the sulfo-NHS acetate reagent by water. All experiments were repeated 15 times to ensure reproducibility and reliability and to generate sufficient data for statistical analysis. For nano-LC-MS/MS analysis, samples were rapidly defrosted on ice and diluted 4-fold in H2O/acetonitrile (98/2, v/v) containing 0.1% formic acid to achieve final tetramer and C1q concentrations of 136 and 143 nM, respectively. Samples were injected randomly to ensure reliability of semi-quantitative measurements. 

Validation of the lysine accessibility mapping procedure- Two critical steps of our experimental approach needed to be validated, namely the quenching of lysine labelling and the quenching of pepsin digestion. This was performed using FBIP (fragment 400-411) as a test sample; its mass increase due to labelling with sulfo-NHS acetate was monitored by MALDI-TOF MS analysis (see the MALDI-TOF-MS section below). FBIP (12.6 pmol) was prepared in 20 µL of 250 mM NaCl, 4 mM CaCl2, 250 mM Hepes, pH 7.3. One µL of a freshly prepared 3.1 mM sulfo-NHS acetate solution was then added to reach a ratio of 250 mol reagent/mol lysine residues and the mixture was incubated for 5 min at 21°C. The reaction was stopped by decreasing the pH to 3.0 using 1 µL of a 50% trifluoroacetic solution. Upon modification at neutral pH, about 50% of FBIP was acetylated, resulting in an increase in molecular weight of 42 Da (supplementary Figs. S1A & S1B). To evaluate the efficiency of the quenching by TCEP-HCl, the FBIP solution was acidified to pH 3.0 prior to the addition of sulfo-NHS acetate. In addition to decreasing the pH value, TCEP-HCl reduces protein disulfide bonds within a few minutes at room temperature. Labelling was allowed to proceed for 5 or 30 min. In contrast to the positive control (supplementary Fig. S1A), the labelling reaction was completely inhibited when the pH was decreased to 3.0. The acetylated form of FBIP was no longer observed on the MALDI mass spectra, indicating that the quenching procedure was highly effective (supplementary Fig. S1C).

In the procedure used (Fig. 1), following sample reduction, labelled and unlabelled samples are subjected to pepsin digestion on ice. After protein hydrolysis, pepsin is irreversibly denatured by increasing the pH from 3.0 to 8.0 for 1 min. Because the acetylation reaction is pH-sensitive, this increase may re-activate the labelling. To minimize as far as possible unwanted lysine modifications on the generated peptic fragments, pepsin inactivation was performed at 0°C. To check that lysine acetylation was inhibited at this temperature, FBIP was used as a control sample. Briefly, the FBIP solution was prepared and acidified as described above, then 1 µL of a 3.1 mM sulfo-NHS acetate solution was added and the mixture left on ice for 7 min. The pH was then raised to 8.0 by adding 1.5 µL of an 8 M NaOH solution and the mixture was left at 0°C for 1 more min prior to re-acidification to pH 2.5 with a 50% trifluoroacetic acid solution. The extent of acetylation observed by MALDI-TOF MS analysis was very low (supplementary Fig. S1D), confirming that the structural information obtained on the samples would be retained upon pepsin inactivation.

Control of the tetramer activation state in C1- The activation state of the tetramer upon interaction with C1q was checked by MALDI-TOF-MS analysis. For this purpose, after incubation for 30 min at 21°C, the protein solution was acidified to pH 3.0 using 110 mM TCEP-HCl and incubated for a further 10 min to reduce disulfide bonds. Acidified samples were then prepared for MALDI-TOF-MS analysis in the linear mode.
Biosystems, Boston, MA). For analysis of intact proteins, each acidified sample (10 µL) was desalted using a ZipTip C4 tip and eluted with 5 µL of H2O/acetonitrile (40/60, v/v) containing 1% formic acid. The same procedure was used for peptides, using a ZipTip C18 tip. All desalted samples were mixed in a 1:1 (v/v) ratio with a matrix solution. For analysis of proteins in the linear mode, the matrix consisted of a saturated sinapinic acid solution prepared in H2O/acetonitrile (70/30, v/v), 1% formic acid, whereas saturated α-cyano-4-hydroxycinnamnic acid in H2O/acetonitrile (1/1, v/v), 1% formic acid was used for peptide analysis in the reflectron mode. One µL of each sample-matrix mixture was spotted on the target plate and air-dried.

In the linear mode, data were acquired with an accelerating voltage of 25 kV, a 75% grid voltage and a 445 ns extraction delay time. In the reflectron mode, these values were set to 20 kV, 70% and 225 ns. Spectra were produced by signal accumulation obtained from 100 consecutive laser shots. All data were reprocessed using the Applied Biosystems Data Explorer 4.0 software. The mass scale was calibrated externally using either the TIS test mixture (Sequazyme™ Peptide Mass Standards kit, Applied Biosystems) for the reflectron mode (m/z 500 to 2500) or a bovine serum albumin solution (m/z 4500 to 120000) for the linear mode.

Nano-LC-ESI-MS/MS experiments were performed with an Ultimate 3000 nano-LC system (Dionex) connected to a LTQ Orbitrap XL mass spectrometer (ThermoFisher Scientific) equipped with a nanospray ion source. Unlabelled and labelled samples (205 fmol of tetramer and 214 fmol of C1q), were loaded onto a C18 µ-precolumn cartridge (Acclaim PepMap100, 5 mm × 300 µm i.d., 5 µm, 100 Å, Dionex) and desalted at 30°C for 5 min at a flow rate of 20 µL/min with 100% buffer A (H2O/ acetonitrile (98/2, v/v) containing 0.1% formic acid). Peptides were then separated on an analytical capillary C18 column (Acclaim PepMap100, 15 cm × 75 µm i.d., 3 µm, 100 Å, LC-Packings) by means of a 2-80% acetonitrile gradient over 50 min at 30°C and a flow rate of 0.3 µL/min. The nano-LC eluate was directly interfaced to the nanospray ion source using a coated silica tip (10.5 cm length, 20 µm i.d with 10 µm i.d at the tip extremity). The ion spray voltage was 1.6 kV with a transfer capillary temperature of 200°C.

For MS/MS experiments, the LTQ Orbitrap XL was operated in the data-dependent mode. Each scan cycle comprised one MS scan in the Orbitrap analyzer (m/z range 400-2000, scan resolution of 30000 and MS target value of 3 × 10^5 with a maximum injection time of 500 ms) followed by five sequential data-dependent MS2 scans performed in the linear ion trap to fragment the 5 most intense precursors found in the preceding MS spectrum (MS² target value of 1 × 10^4 with a maximum injection time of 100 ms). For collision-induced dissociation, the normalized collision energy was 35%, the activation time was 40 ms, the isolation width was 3.0 u, the activation q was 0.25 and the minimal signal required was 500. To ensure mass accuracy, the Orbitrap analyzer was calibrated in the positive mode according to the manufacturer’s instructions. For all MS/MS experiments, the monoisotopic precursor selection mode was selected and the dynamic exclusion was used with one repeat count, a 30 s repeat duration and an exclusion duration of 90 s. Mass spectra were processed using Bioworks 3.3.1 (ThermoFisher Scientific) to generate Mascot-compatible MGF file.

**MS/MS data processing – Procedure of label-free relative quantification**

Peptide identification was carried out using either the SEQUEST algorithm (ThermoFisher Scientific) or the locally installed Mascot search engine (version 2.2.1., Matrix Science) using an in-house database containing the amino acid sequences of C1r and C1s. The mass tolerance was set to 10 ppm and 0.8 Da for MS and MS/MS analyses, respectively. Given the unspecificity of pepsin, no enzyme was selected for peptide assignment. Acetylations on lysine, serine and tyrosine residues were set as variable modifications. Possible oxidation of methionine residues was also taken into account.

To assess the effects of C1q binding on the solvent accessible surface of the tetramer, the extracted ion currents (XIC) of the precursor ions were utilized for relative quantification. Quantification was performed either manually or by using the SuperHirn software (http://prottools.ethz.ch/muellelu/web/SuperHi...
SuperHirn was used to rapidly identify peptides exhibiting a modified acetylation pattern in the presence of C1q. To construct the MasterMaps, raw data were first converted to mzXML files with ReAdW (http://tools.proteomecenter.org) and pepXML files were generated with the Mascot search engine. Manual quantification was performed as follows: for each identified peptide containing lysine residues, the XIC of the precursor ion was extracted from the total ion current and the area-under-peak (AUP) was calculated using QualBrowser 2.0.7 (ThermoFisher Scientific). The unmodified fraction (UF) of each peptide was then estimated using the following formula (28):

$$UF = \left( \frac{\text{AUP peptide Labeling Assay}}{\text{AUP peptide Unlabelled Control}} \right) \times 100\%$$

The UF corresponds to the fraction of an unlabelled tetramer peptide remaining after acetylation in the presence or absence of C1q.

**Statistical analysis** - For each peptide, the UFs calculated from the 15 replicate samples were pooled and organized into two independent groups containing all the values obtained in the absence (group A) or in the presence (group B) of C1q. A non-parametric Mann-Whitney U-test was then applied to test the null hypothesis (H₀). H₀ is defined as the hypothesis of “no difference”, meaning no solvent accessibility difference between groups A and B. A two-sided p-value was used to perform the U-test. The p-value corresponds to the estimated probability of rejecting the null hypothesis and was set to 1% (less than 1 in 100 chances of being wrong). An excel sheet containing the Mann-Whitney U-test was directly downloaded from http://anastats.com.

UF values obtained for each peptide were graphically depicted using a box-and-whisker plot to compare the statistical dispersion between groups A and B. Box-plots were constructed as follows. Data were first ordered from the smallest to the highest UF values. Then, the lower (Q1), median (Q2) and upper (Q3) quartiles were calculated. Quartiles divide the data set into four equal parts, so that each represents one fourth of the ordered set of UF values. Q2 corresponds to the exact middle of the ordered set of all UF values (central tendency) whereas Q1 and Q3 represent the exact middle numbers of the lower and upper half of the UF data set, respectively. Then, the inter-quartile range (IQR) was calculated by subtracting the lower quartile from the upper quartile. The inter-quartile range corresponds to a measurement of the statistical dispersion of the population of each group. The minimum (Min) and maximum (Max) non-outlier values were calculated using the following equations:

$$\text{Min} = Q_1 - (1.5 \times IQR)$$
$$\text{Max} = Q_3 + (1.5 \times IQR)$$

Any data lying below the Min or above the Max values was considered to be an outlier.

**RESULTS**

Most of our current knowledge about the overall architecture of C1 arises from low-resolution studies by electron microscopy and neutron scattering as well as from the crystal structure of C1r and C1s fragments and of the C1q globular heads. The main barriers to solve the whole structure of C1 and decipher the mechanism of its assembly lie in the size of the complex and the fact that it involves non-covalent interactions. To gain access to new structural information about this complex, we used mass spectrometry in association with a stable chemical modification of lysine residues. We chose to investigate the solvent accessibility of lysines in the free and complexed forms of the tetramer for several reasons: (i) these residues are abundant and evenly distributed in the tetramer; (ii) lysine residues are mostly located on the surface of proteins and are therefore ideal candidates for probing protein-protein interfaces; (iii) unlike C1q, the tetramer does not have an oligomeric organisation, which is expected to facilitate identification of the areas of C1r and C1s involved in conformational changes and/or in binding to C1q. In addition, previous studies have shown that chemical modifications of the lysine residues of the tetramer do not prevent assembly of C1 (29).

The solvent accessibility of lysine residues was probed by stable chemical modification with the primary amine-specific reagent sulfo-NHS acetate. The free tetramer and the reconstituted C1 complex were both exposed to an excess of reagent, and unlabelled controls were prepared in parallel to
calculate the unmodified fraction of each lysine-containing peptide (see below). The labelling was then quenched by decreasing the pH to 3.0 using a TCEP-HCl solution. Following reduction of the disulfide bridges, labelled and unlabelled protein samples were both subjected to proteolysis by pepsin, an acid protease that retains its enzymatic activity under the quenching conditions i.e. at low pH, low temperature and high concentration of reducing agent (30,31). After proteolysis, pepsin was irreversibly denatured by increasing the pH to 8.0.

**Effects of C1q binding on the lysine acetylation pattern of C1r-** To validate our approach, it was essential to ensure that the acetylation reaction was carried out under conditions where the integrity of the C1 complex was fully preserved. First, labelling was performed at neutral pH so that the native conformation of C1 was retained. In addition, the activation state of C1 was checked throughout the labelling procedure, considering that C1 is known to undergo spontaneous activation *in vitro* even without binding to a target (32). Activation of the complex induces structural changes within the tetramer (9,33) and leads to the hydrolysis of the Arg-Ile bond located in the SP domain of each protease. As a result, both activated C1r and C1s proteins comprise two chains linked by a disulfide bridge (Fig. 2A). To evaluate the extent of C1 activation under the experimental conditions used, the complex was incubated at room temperature for 30 min at pH 7.3, this incubation time corresponding to the duration of the whole procedure, except pepsin digestion. As shown in Figure 2B, a minor fraction of the tetramer was found to be activated in the absence of C1q. This activation mainly occurred during the multiple purification steps required to isolate the tetramer from human plasma. However, the activated fraction of the tetramer did not increase in the C1 complex after incubation for 30 min at room temperature (Fig. 2C), indicating that, under the experimental conditions used, the structural integrity of C1 was preserved throughout the labelling procedure.

Thirty-six peptides containing lysine residues were assigned to C1r by LC-MS/MS analysis of the pepsin digests, thus accounting for 25 of the 36 lysines of this protein (Fig. 3A). Residues that were not recovered were mainly located at the C-terminal end of the CCP2 module (K\(^{419}\), K\(^{423}\), K\(^{426}\) and K\(^{436}\)) and in the SP domain (K\(^{490}\), K\(^{514}\), K\(^{516}\)). To assess the effects of C1q binding on the accessibility of each lysine residue, a statistical Mann and Whitney U-test with a two-sided \(p\)-value of 1% was applied on the 15 independent MS data sets collected. As listed in Table 1, the solvent accessibility of most of the residues located in modules CUB\(_1\) and CUB\(_2\) remained unchanged upon C1q binding. Interestingly, a single lysine located in the N-terminal part of CUB\(_1\) (Fig. 3A) behaved differently. The box-and-whisker diagram of fragment 2-8 reveals that K\(^7\) is highly exposed in the free tetramer, as only 3% remains unmodified after acetylation (Fig. 3B). This value shifts to \(~23%\) in the presence of C1q, indicating that K\(^7\) is partially protected from chemical modification within C1.

C1r K\(^{280}\) and K\(^{296}\) are also fully exposed to the solvent in the free tetramer, with solvent accessibility surface area (SASA) values of 81.2 and 159.0 Å\(^2\), respectively. These two residues are located in the N-terminal part of the CCP\(_1\) module and are covered by two distinct overlapping peptides 289-300 and 289-301 (Figs. 3A & 3C). Surprisingly, the overall acetylation extent of K\(^{280}\) and K\(^{296}\) in the free tetramer was strikingly different in these peptides: about 3% of peptide 289-300 remained unacetylated whereas this value increased to 58% for peptide 289-301 (Figure 3B). This difference was observed consistently in all experiments, suggesting the presence of two distinct conformations in solution. In addition, the chemical reactivity of both peptides towards the acetylating agent was also significantly different in the presence of C1q. Upon C1 formation, the unmodified fraction of peptide 289-301 remained unchanged whereas the accessibility of peptide 289-300 slightly decreased (Fig. 3B). Taken together, these observations suggested that the N-terminal end of the C1r CCP\(_1\) module exhibits two different conformations, both in the isolated tetramer and in C1.

Four other lysines exhibiting reduced solvent accessibility upon C1q binding were also identified in the C1r CCP\(_2\) and SP domains. Residues K\(^{382}\) and K\(^{395}\) are both located in the CCP\(_2\) region (Fig. 3C) and are fully exposed to the solvent, as judged by their calculated SASA values of 114.3 Å\(^2\) and 145.6 Å\(^2\). Upon C1q binding, both lysines became
protected from modification, with a 30% increase of their respective unmodified fraction (data not shown). Similar results were observed with K^{452} and K^{484} from the SP domain. As seen in the crystal structure of the proenzyme SP domain, these latter residues are very close to the activation segment of C1r (Figs. 3A & 3C) and display similar SASA values (Table 1). When the C1 complex is formed, both lysines become less accessible to the acetylating reagent, with an increase of 20% of their unmodified fraction (Fig. 3B). This result was surprising since the activation segment contains the susceptible Arg^{466}-Ile^{477} bond cleaved upon autolytic activation of C1r. Therefore, no major change in the solvent accessibility of this region was expected, considering that this site should remain fully accessible. A possibility is that the activation segment becomes less exposed to the solvent in the resting C1 complex in order to adopt a conformation inappropriate for C1r self-activation. However, this hypothesis does not appear consistent with previous data indicating that the C1r activation potential is prevented in the free tetramer, and restored in C1 (34).

**Effects of C1q binding on the lysine acetylation pattern of C1s-** We next investigated the effects of C1 assembly on the acetylation pattern of C1s. As summarized in Figure 4A, 30 peptic peptides were assigned to C1s by MS/MS, thus accounting for 26 of the 37 lysine residues of the protein. Most of the lysines that were not recovered are clustered in the C-terminal part of the CUB2 module (K^{240}, K^{255}, K^{266}, K^{267}) and in the SP domain (K^{560}, K^{568}, K^{579}, K^{581}). Two lysines exhibiting reduced solvent accessibility in C1 were identified in C1s CUB1 and CUB2. Residues K^{90} (CUB1) and K^{195} (CUB2) appear fully exposed in the isolated tetramer and become less accessible in the presence of C1q. This effect is particularly striking in the case of K^{90}, which shows a 40% increase of its unmodified fraction, thereby becoming virtually inaccessible in C1 (Figs. 4B & 4C). A similar overall tendency was observed for the lysine residues located in the CCP1 module, except K^{281} and K^{338} (Table 2). In contrast to the above observation, the two lysine residues located in CCP2 and the single lysine K^{420} found in the activation segment showed no change in accessibility upon assembly of the C1 complex (Table 2, supplementary Fig. S2). K^{420} is located in the activation segment of C1s that contains the Arg^{422}-Ile^{423} bond cleaved upon activation by C1r. The activation segment of C1s thus appears to remain fully accessible in C1, in contrast to our observation in C1r.

Among the 18 lysine residues located in the C1s SP domain, five (K^{432}, K^{484}, K^{486}, K^{608}, K^{614}) showed significantly reduced accessibility within C1 (Table 2; Fig. 4C). As shown in Figure 4A, K^{500} is covered by two overlapping peptides, 494-501 and 483-501. No significant modification of solvent accessibility was observed for peptide 494-501, whereas the unmodified fraction of peptide 483-501 increased by 16% (Fig. 4B), indicating that reduction of solvent accessibility was restricted to the segment containing K^{484} and K^{486}. This example illustrates the advantage of using the non-specific protease pepsin, which generates overlapping fragments, thereby allowing in some cases the accessibility of particular lysine residues to be assessed.

The accessibility of peptides 602-620 and 602-621 was also significantly reduced (about 20% in both cases) upon interaction with C1q, indicating decreased exposure of K^{608} and/or K^{614} (Fig. 4B). In contrast, two lysines showing increased solvent accessibility, K^{584} and K^{587}, were identified in the C1s SP domain (Fig. 4B). Both residues lie in a segment not defined in the C1s catalytic domain X-ray structure and are possibly located in the vicinity of the active site entrance (Fig. 4C). Binding to C1q markedly increases their reactivity towards the acetylating reagent, indicating that conformational changes occur in this segment upon C1 assembly.

**DISCUSSION**

X-ray crystallography and NMR spectroscopy have been extensively used over the past decade to gain structural information about the constituent proteins of human C1, allowing resolution of 67% and 72% of the C1r and C1s structures, respectively (6). Although there are still missing links, namely the C1r CUB1 and CUB2 modules and the C1s CUB2-CCP1 segment, these data provide an overall view of the 3-D organization of the free C1s-C1r-C1r-C1s tetramer in solution. Earlier information arising from neutron scattering (35,36) and electron microscopy analyses (37-39) had led to the concept that the
extended tetramer folds into a more compact conformation upon interaction with C1q, providing the basis for most of the low-resolution C1 models proposed originally (13-15).

The sites of C1r and C1s involved in the interactions between the tetramer and C1q have recently been delineated by site-directed mutagenesis (17), revealing that C1 assembly involves high-affinity C1q-binding sites contributed by the C1r CUB 1 and CUB 2 modules and lower affinity sites contributed by the C1s CUB 1 modules. Based on the location of these sites and available structural information, a refined three-dimensional model of C1 assembly has been proposed, where the CUB 1-EGF-CUB 2 interaction domains of both C1r and C1s are entirely located inside C1q and interact via six binding sites with reactive lysines located approximately half-way along the C1q collagen-like stems (17,26,40). Based on the use of truncated protease segments, a similar interaction model was derived by another group (41).

The present study provides for the first time a detailed comparative analysis of the accessibility of the lysine residues of C1r and C1s in the free and complexed forms of the C1s-C1r-C1r-C1s tetramer, allowing us to test whether our current concept of C1 assembly (17) is consistent with this new information. As no experimental X-ray structure is available yet for C1r CUB 1 and CUB 2 and C1s CUB 2 and CCP 1, the data relating to these modules will not be discussed in detail, considering the relative imprecision of 3-D homology models. Nevertheless, it is interesting to note that, with the exception of K7 in C1r CUB 1 and K195 in C1s CUB 2, the accessibility of the lysine residues contained in these modules remains unchanged upon C1 assembly (Tables 1 & 2). This is consistent with the fact that C1r and C1s interact with C1q through acidic residues contributed by their CUB modules (17), and with earlier studies showing that chemical modification of the lysine residues of C1s-C1r-C1r-C1s has no effect on C1 assembly (29). The fact that little structural changes are detected in the N-terminal interaction regions of C1r and C1s also likely arises for a large part from the high stability of the head-to-tail C1r/C1s CUB 1-EGF heterodimeric assemblies that connect C1r to C1s in the tetramer and are expected to retain a relatively rigid conformation in the C1 complex (16).

The labelling data relating to the C1s CUB 1 module, in contrast, can be analyzed in light of the C1s CUB 1-EGF X-ray structure (16). In this module, whereas the accessibility of K23 and K96 remains unchanged upon C1 assembly, that of K90 decreases very significantly (Table 2, Fig. 4B). As judged from the C1s CUB 1-EGF structure, all three lysine residues are expected to be accessible in the free tetramer. The new version of the C1 model is largely based on the assumption that both C1r/C1s CUB 1-EGF-CUB 2 heterodimers, which are distant in the free tetramer, become closely packed side by side through their C1s moieties to form a single compact assembly inside the C1q cone (17). That such a close packing actually occurs in C1 is fully supported by our labelling data, considering that in the resulting assembly K90 is positioned in the middle of the C1s/C1s interface, consistent with its decreased accessibility (Figs. 5A & 6A). Conversely, K23 is expected to remain accessible in C1, and the same applies to K96, despite the proximity of this latter residue with the low-affinity C1q-binding site harboured by C1s CUB 1 (17). Thus, that C1 assembly involves formation of a new interaction between both C1s CUB 1-EGF-CUB 2 domains, an essential requirement of our current C1 model, appears fully consistent with the above experimental data. Conversely, the fact that most of the lysine residues of C1r CUB 1 and CUB 2 are labelled to similar extents in the free tetramer and in C1 is also compatible with the model, considering that, except in the vicinity of the C1q-binding sites, these modules are expected to remain fully accessible in C1, given their location on the outer part of the C1r/C1s interaction domains assembly (Fig. 6A).

An intriguing finding from our study lies in the peculiar labelling data relating to the C1r segment Ile289-Phe301, suggesting that this area exhibits two different conformations in both the free and complexed forms of the tetramer (Fig. 3B). As this sequence stretch is covered by two overlapping peptides differing by a single residue (Fig. 3A), we initially suspected an artefact of the “label-free” quantification procedure. However, similar results were consistently obtained when performing 15 independent labelling experiments, hence validating the above
hypothesis. Indeed, this segment, at the interface between the CUB2 and CCP1 modules of C1r, is known to be intrinsically very flexible, as shown by its high susceptibility to cleavage by a variety of proteolytic enzymes (42). Likewise, one of the two lysines contained in this segment (K296) exhibits opposite orientations in the two X-ray structures currently available (10,11). Thus, that the two overlapping peptides are differentially labelled in the free tetramer likely reflects the fact that the corresponding sequence stretch adopts two alternative conformations. In the C1 complex, the CUB2 modules of C1r are thought to occupy the upper part of the C1q cone, the following CCP1-CCP2-SP catalytic domains being in contrast positioned in the lower part (17). Connection between these two compartments therefore likely requires significant conformational changes at the CUB2/CCP1 interface, which appears consistent with the decreased lysine accessibility observed for peptide 289-300 (Fig. 3B). Why in contrast the accessibility of the other peptide 289-301 remains unchanged in C1 is currently unknown, although it cannot be excluded that this reflects a structural asymmetry between the CUB2-CCP1 junctions of both C1r molecules, connected with the expected asymmetry of the C1r activation process itself (6). It is noteworthy that residues K293 and K295 of C1s, located in the equivalent region of module CCP1 at the interface with the preceding CUB2 module, also exhibit decreased accessibility in C1. This is consistent with the hypothesis that C1s also possesses significant flexibility at its CUB2/CCP1 interface, a feature expected to be necessary for the C1s activation process (6).

The C1r residues K155 and K157, on either side of the CCP1/CCP2 junction, are both exposed in the C1r CCP1-CCP2-SP structure (10) and remain fully accessible in the C1 model, in agreement with our labelling data (Table 1). In the same way, K395 (C1r CCP2), which is near the C1r SP domain in the CCP1-CCP2-SP structure, could move closer to this domain, by means of the limited flexibility of the CCP2/SP interface (9,10), or could become engaged in a salt bridge with the neighbouring residue E391, hence its decreased accessibility in the C1 complex. Residues K629, K672, K681, and K682 of the C1r SP domain are all exposed, to various extents, in the CCP1-CCP2-SP structure (Fig. 3C) and remain accessible in the C1 model, in agreement with the labelling data (Table 1). In contrast, the fact that residues K452 and K454 show a decreased labelling in C1 does not appear fully consistent with the model, according to which they should remain accessible despite their orientation towards a C1r CUB2 module. It should be emphasized, however, that K584 exhibits variable orientations in the four available X-ray structures (9,10,11), and lies in the vicinity of the C1r activation segment 444-450, which is very unstable in the zymogen form. Finally, according to our model, K582 (C1r CCP3) should remain accessible in the C1 complex, in contrast to the labelling data.

A major outcome of this study lies in the observed differential accessibility of the lysine residues contained in the C1s SP domain, a finding which for the first time indicates that, in the proenzyme C1 complex, this domain is at least partially inserted in the C1q cone, and yields precise information about its orientation in the complex. Thus, the lysine residues showing unmodified accessibility upon interaction of the tetramer with C1q (K500, K629, K631, K554, K662) are all located on the same side of the C1s SP domain (Fig. 5B), indicating that this part of the domain remains exposed to the solvent and therefore likely faces the outside of the complex. Consistent with this hypothesis, four residues showing decreased labelling in C1 (K432, K484, K486, K608, K614) lie on the opposite side of the C1s SP domain (Fig. 5B), providing strong indication that, conversely, this region faces the inside of the complex. Thus, it is tempting to hypothesize that K432, K484, K486 and K608 may be involved in interactions with the C1r catalytic domains positioned inside the C1q cone or with C1q itself. In contrast, the implication of K614 in such interactions appears less likely considering its particular location above the active site entrance (33). This new information allows us to propose a complete version of our C1 model (17) integrating the C1s catalytic domain (Figs. 6B & 6C). In the resulting assembly (Fig. 6), residues K584 and K587 of the C1s SP domain, which are disordered in the C1s CCP2-SP structure ((33); Fig. 4C) are located in the vicinity of a C1q collagen fibre. The C1s CCP2 module is positioned in such a way that its residues K544 and K565 remain accessible, in agreement with the labelling data (Table 2).
In summary, a strategy combining chemical modification of lysines and mass spectrometry analysis has been applied for the first time to the human C1 complex. The structural data generated are, for the most part, consistent with our recent 3-D C1 model (17). In addition, they yield the first experimental evidence that the C1s SP domain is partly positioned inside the C1q cone. The same type of analysis could be applied to probe other residues such as tyrosines and histidines and would provide further precise insights into the C1 architecture.

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FOOTNOTES

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²The abbreviations used are: ESI, electrospray ionization; nano-ESI, nanoelectrospray ionization; MALDI, matrix-assisted laser desorption/ionization; TOF, time of flight; MS, mass spectrometry; MS/MS, tandem mass spectrometry; MSⁿ, general designation of mass spectrometry to the nth degree; XIC, extracted ion chromatogram.
FIGURE LEGENDS

**Figure 1.** Overview of the experimental approach used in this study. Experimental details are described in the Materials & Methods section.

**Figure 2.** Effect of C1q binding on the activation state of the C1s-C1r-C1r-C1s tetramer. (A) Modular structure of C1r and C1s. Both proteases are activated through cleavage of an Arg-Ile bond (represented by a black arrow) located in their SP domain. The only disulfide bridge shown is the one connecting the activation peptide to the SP domain. N-linked oligosaccharides are represented by open circles. (B) MALDI-TOF mass spectra of the isolated tetramer under reducing conditions. In the absence of C1q, a small fraction of the tetramer appears activated, as indicated by the peaks corresponding to the heavy chains of C1r and C1s. (C) MALDI-TOF mass spectra of the reconstituted C1 complex under reducing conditions. The peaks at m/z 48114.6 (1), 49695.5 (2) and 51317.8 (3) correspond to random associations of the C1q chains (43).

**Figure 3.** Modification of the C1r solvent accessibility upon association of the C1s-C1r-C1r-C1s tetramer with C1q. (A) Amino acid sequence of C1r showing the 36 lysine-containing peptic fragments (in red, bold type, underlined) selected for quantitative analysis. Lysines that could not be recovered are shown in black, bold type, underlined. The catalytic residues His$^{485}$, Asp$^{540}$, Ser$^{637}$ and the Arg-Ile cleavage site are highlighted in magenta and shown in blue, respectively. C1r residues interacting with C1q (17) are highlighted in yellow. (B) Effect of C1q binding on the solvent accessibility of residues K$^{2}$ (CUB$_{1}$), K$^{291}$/K$^{296}$ (CCP$_{1}$) and K$^{452}$/K$^{454}$ (SP domain). Each box-and-whisker plot compares the statistical distribution of the unmodified fraction of a given C1r peptide in the presence (C1) or absence (tetramer) of C1q. Q1, Q2, Q3 correspond to the lower, median (red bar) and third quartile, respectively. The largest (Max) and smallest (Min) non-outlier observations are marked with a small black vertical line (whiskers). Data points lying above the upper whisker or below the lower whisker are considered as outliers and indicated by an open circle. (C) Structure of the zymogen CCP$_{1}$-CCP$_{2}$-SP C1r catalytic domain (10) showing the position of lysine residues. The catalytic triad (His$^{485}$, Asp$^{540}$, and Ser$^{637}$) is represented by 3 magenta spheres. Lysine residues are colour-coded as follows: blue, no modification of surface accessibility upon C1 assembly; red, decreased surface accessibility; yellow, decreased and/or unmodified surface accessibility; black, no data available.

**Figure 4.** Modification of the C1s solvent accessibility upon interaction of the C1s-C1r-C1r-C1s tetramer with C1q. (A) Amino acid sequence of C1s showing the 30 lysine-containing peptic fragments used for quantitative analysis. The colour coding used is the same as stated in the legend to Figure 3. (B) Effect of C1q binding on the surface accessibility of residues K$^{90}$ (CUB$_{1}$), K$^{195}$ (CUB$_{2}$) and residues K$^{484}$, K$^{486}$, K$^{500}$, K$^{558}$, K$^{587}$, K$^{608}$, K$^{614}$ of the SP domain. Each box-and-whisker plot compares the statistical distribution of the unmodified fraction of a given C1s peptide in the presence (C1) or absence (tetramer) of C1q. (C) Structures of the C1s CUB$_{1}$-EGF (16) and CCP$_{2}$-SP regions (33) showing the position of lysine residues. The Ca$^{2+}$ ions bound to CUB$_{1}$ (site I) and EGF (site II) are represented by yellow spheres, and the catalytic triad is shown as 3 magenta spheres. Orange dots correspond to residues not defined in the C1s CCP$_{2}$-SP X-ray structure. Lysine residues are colour-coded as follows: blue, no modification of solvent accessibility inside the C1 complex; red, decreased accessibility; green, increased accessibility; black, no data available.

**Figure 5.** (A) Space-filling representation of the head-to-tail C1r/C1s CUB$_{1}$-EGF heterodimer. One C1s monomer (grey) was used as a template to position and visualize the lysine residues identified in the C1r CUB$_{1}$ module, based on the sequence alignment of the CUB$_{1}$ modules of C1r and C1s (Supplementary Table S1). Lysines are colour-coded as defined in Figure 3. C1r and C1s residues interacting with C1q are coloured green (17). (B) Space-filling representation of the C1s CCP$_{2}$-SP region illustrating the position of the lysine residues (in red) showing reduced solvent accessibility upon C1q binding, except for K$^{484}$ which lies in a region not defined in the crystal structure (33). Lysines undergoing no modification of solvent accessibility within C1 are all located on the same face, opposite to the one harbouring K$^{432}$, K$^{486}$, K$^{608}$ and K$^{614}$. 

Downloaded from http://www.jbc.org/
Figure 6. (A) Space-filling representation of the assembly of the C1r/C1s CUB1-EGF-CUB2 interaction domains as proposed to occur in the C1 complex (17) (top view). C1r and C1s are shown in yellow and grey, respectively. The colour coding used is the same as stated in the legend to Figure 3. Residues interacting with C1q are coloured green. The six collagen triple helices of C1q are shown as magenta spheres. (B, C) Side and bottom views of the whole C1 complex highlighting the positioning of the C1s SP domains with respect to the remainder of the complex. Both C1r monomers are in yellow, whereas C1s molecules are shown in cyan and magenta. The colour coding used for lysine residues is the same as stated in the legend to Figure 3.
TABLE 1

Effect of C1 assembly on the solvent accessibility of the lysine residues of C1r. Residues showing significant changes in their accessibility are highlighted in grey and shown in bold type.

| Domain | Lysines | SASA (Å²) | Mann & Whitney U-test | Accessibility within C1 |
|--------|---------|-----------|----------------------|------------------------|
| CUB1   | K7      | False     |                      | Decreased              |
|        | K19     | True      |                      | Recessed               |
|        | K40     | True      |                      | Unchanged              |
|        | K60     | True      |                      | Unchanged              |
|        | K65     | True      |                      | Unchanged              |
|        | K66     | True      |                      | Unchanged              |
|        | K85     | ND (e)    |                      | ND                     |
|        | K94     | True      |                      | Unchanged              |
|        | K115    | True      |                      | Unchanged              |
| EGF    | K134    | 59.0      |                      | ND                     |
|        | K218    | True      |                      | Unchanged              |
|        | K245    | True      |                      | Unchanged              |
| CUB2   | K282    | ND        |                      | ND                     |
|        | K322    | 99.8      |                      | ND                     |
|        | K355    | 101.1     |                      | ND                     |
|        | K357    | 46.7      |                      | ND                     |
| CCP1   | K291    | 81.2      |                      | False                   |
|        | K296    | 159.0     |                      | True & False            |
|        | K322    | 99.8      |                      | ND                     |
|        | K355    | 101.1     |                      | ND                     |
|        | K382    | 114.3     |                      | False                   |
|        | K395    | 145.6     |                      | Decreased               |
| CCP2   | K419    | 64.0      |                      | False                   |
|        | K423    | 50.6      |                      | False                   |
|        | K425    | 165.6     |                      | False                   |
| a.p. (g) | K436   | 89.3      |                      | ND                     |
| SP     | K452    | 124.4     |                      | False                   |
|        | K454    | 136.1     |                      | Decreased               |
|        | K490    | 8.8       |                      | ND                     |
|        | K514    | 42.4      |                      | ND                     |
|        | K585    | 101.0     |                      | True                    |
|        | K610    | 85.3      |                      | ND                     |
|        | K629    | 115.2     |                      | True                    |
|        | K672    | 30.0      |                      | True                    |
|        | K681    | 98.9      |                      | True                    |
|        | K682    | 146.4     |                      | True                    |

(a) Solvent Accessibility Surface Area of lysine side-chains. Accessible surface areas of C1r are based on available X-ray data [pdb accession numbers: 1APQ (EGF); 1GPZ (CCP1-CCP2-SP); 1MD8 (SP)] and calculated by using the software program VADAR (44). (b) H0: hypothesis of “no difference” of solvent accessibility between the free tetramer and C1. (c) Two-sided p-value; the significant level at which H0 is rejected is set to 1%. (d) No Structure available. (e) ND: not determined. (f) Residues covered by two distinct peptic peptides with different solvent accessibility modifications upon C1q binding. (g) Activation peptide. (h) Residue undefined in the crystal structure (10).
TABLE 2

Effect of C1 assembly on the solvent accessibility of the lysine residues of C1s. Residues showing significant changes in accessibility upon C1q binding are highlighted in grey and shown in bold type.

| Domain | Lysines | SASA (Å²) | Mann & Whitney U-test | Accessibility within C1 |
|--------|---------|-----------|-----------------------|-------------------------|
| CUB1   | K23     | 93.0      | True                  | Unchanged               |
|        | K90     | 117.3     | False                 | Decreased               |
|        | K96     | 81.2      | True                  | Unchanged               |
| EGF    | K154    | 67.2      | ND (e)                | ND                      |
| CUB2   | K179    | True      | Unchanged             |
|        | K195    | False     | Decreased             |
|        | K249    | ND (d)    | ND                    |
|        | K265    | ND        | ND                    |
|        | K266    | ND        | ND                    |
|        | K269    | ND        | ND                    |
| CCP1   | K281    | ND        | ND                    |
|        | K293    | False     | Decreased             |
|        | K295    | False     | Decreased             |
|        | K331    | False     | Decreased             |
|        | K336    | False     | Decreased             |
|        | K338    | ND        | ND                    |
| CCP2   | K354    | 117.5     | True                  | Unchanged               |
|        | K405    | 122.0     | True                  | Unchanged               |
| a.p    | K420    | (f)       | True                  | Unchanged               |
|        | K432    | 66.2      | False                 | Decreased               |
|        | K484    | 87.0      | False                 | Decreased               |
|        | K500    | 112.8     | True                  | Unchanged               |
|        | K521    | 100.0     | ND                    | ND                      |
|        | K525    | 96.8      | ND                    | ND                      |
|        | K560    | 153.2     | ND                    | ND                      |
|        | K568    | 37.2      | ND                    | ND                      |
|        | K579    | 87.4      | ND                    | ND                      |
|        | K581    | 53.0      | ND                    | ND                      |
| SP     | K584    | (g)       | False                 | Increased               |
|        | K587    | (g)       | False                 | Increased               |
|        | K608    | 148.3     | False                 | Decreased               |
|        | K614    | 145.6     | False                 | Decreased               |
|        | K629    | 147.2     | True                  | Unchanged               |
|        | K631    | 54.5      | True                  | Unchanged               |
|        | K654    | 35.0      | True                  | Unchanged               |
|        | K662    | 116.3     | True                  | Unchanged               |

(a) Solvent Accessibility Surface Area of lysine side-chains. Accessible surface areas of C1s are based on available X-ray data [pdb accession numbers: 1NZI (CUB1-EGF); 1ELV (CCP2-SP)] and calculated by using the software program VADAR (44). (b) $H_0$: hypothesis of “no difference” of solvent accessibility between the free tetramer and C1. (c) Two-sided $p$-value; the significant level at which $H_0$ is rejected is set to 1%. (d) No Structure available. (e) ND: not determined. (f) Activation peptide. (g) Residue undefined in the crystal structure (33).
Figure 1

**Free protein**

Free protein complexed protein

| K | K | K | K |

Unlabelled controls

Quench labelling with TCEP, pH 3.0

Reduction of disulfide bridges

| S | H | S | H |

Pepsin digestion, pH 3.0, 0°C

| S | H | S | H |

Nano-LC-ESI-LTQ Orbitrap XL MS/MS analysis

'Label free' quantitative analysis

Unlabelled controls

Ligand Protection

Unmodified fraction 'free protein'

Unmodified fraction 'complexed protein'

Statistical analysis
Figure 2

(A) C1r

(B) C1s

(C) Mass (m/z) vs. Relative Abundance

Mass (m/z) 35000 48000 61000 74000 87000 100000

Relative Abundance

- [M+H]^+ Proenzyme
- C1r
- [M+2H]^{2+}
- Activated Heavy Chains
- C1s
- C1r
- C1q

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Figure 3
Figure 4

(A) Peptide 602-620 (K608, K614)

(B) Peptide 494-501 (K500)

(C) Peptide 483-501 (K484, 486, 500)
Figure 5

(A) 

(B)
Figure 6
Mapping surface accessibility of the C1r/C1s tetramer by chemical modification and mass spectrometry provides new insights into assembly of the human C1 complex

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