Kinetic Analysis of the Interactions of Complement Receptor 2 (CR2, CD21) with Its Ligands C3d, iC3b, and the EBV Glycoprotein gp350/220

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The molecular mechanisms involved in the interaction of complement receptor 2 (CR2) with its natural ligands iC3b and C3d are still not well understood. In addition, studies regarding the binding site(s) of the receptor on C3 as well as the affinities of the C3 fragments for CR2 have produced contradictory results. In the present study, we have used surface plasmon resonance technology to study the interaction of CR2 with its ligands C3d, iC3b, and the EBV surface glycoprotein gp350/220. We measured the kinetics of binding of the receptor to its ligands, examined the influence of ionic contacts on these interactions, and assessed whether immobilized and soluble iC3b bound with similar kinetics to CR2. Our results indicate that 1) gp350 binding to CR2 follows a simple 1:1 interaction, whereas that of the C3 fragments is more complex and involves more than one intramolecular component; 2) kinetic differences exist between the binding of C3d and iC3b to CR2, which may be due to an additional binding site found on the C3c region of iC3b; and 3) iC3b binds to CR2 with different kinetics, depending on whether the iC3b is in solution or immobilized on the surface. These findings suggest that binding of CR2 to iC3b and C3d is more complex than previously thought. The Journal of Immunology, 2001, 167: 1490–1499.
coupled to anti-idiotypic Ab, it induced a strong idiotypic and Ag-specific response in mice (27). This peptide is homologous in sequence to the CR2-binding region on gp350/220. The observation that synthetic peptides representing aa 295–306 (β-chain) and 744–755 (α-chain) showed similarity to the above peptide of C3 that binds to CR2 (25) suggested that additional residues in C3 might participate in the interaction with CR2. These peptides were identified based on their sequence homology with the C3d residues 1201–1214, and the binding was corroborated by the ability of peptides and C3c immobilized to microspheres to bind to Raji cells through CR2 (25).

Kalvi and coworkers (28) have shown that soluble C3d and iC3b inhibit soluble CR2 binding to iC3b-coated erythrocytes to a similar extent. The authors thus concluded that the interaction between CR2 and C3d is delimited to the C3d region. In view of this conclusion and in an effort to map the CR2 binding sites on C3, mutagenic scan studies have been performed in which erythrocytes coated with mutated iC3b were tested for their ability to bind to soluble CR2 and induce the rosetting of CR2-bearing Raji cells. In these studies, mutagenesis of residues 1199–1210 (EDPFKLYNVEA) was shown to reduce the binding of iC3b to CR2 by only 20% (29). The recent availability of the x-ray crystallographic structure of C3d has indicated the presence of a negatively charged pocket on the concave surface of the molecule (30). The authors of this x-ray crystallographic study proposed this pocket as a candidate site for CR2 interaction. Replacement of two clusters of negatively charged C3d residues (E379–383/E390–394) and E1660–1613/D1613–1614/E1664–1672, C3d numbering, mature C3 numbering) present on the opposite side of the acidic pocket led to severe inhibition of the interaction (31). Because the results of the localization of the C3 binding sites for CR2 are apparently contradictory, it is not clear whether the interaction between CR2 and C3 is mediated only through the C3d region or other regions in C3 are involved in the interaction. In addition, it is not clear whether C3d interacts with CR2 through single or multiple sites and whether differences in experimental settings may affect these interactions. In the present study, we have analyzed the kinetics of the interaction of CR2 with C3d and iC3b in real time using surface plasmon resonance (SPR) and have asked whether these interactions are similar in nature. Because gp350 also serves as a ligand for CR2, we have analyzed its interaction with CR2 and have compared this interaction with those of CR2 with C3d and iC3b. Our results indicate that the binding of CR2 to viral gp350 follows a simple 1:1 binding model, whereas its binding to the C3 fragments is more complex. We observed direct binding of CR2 to a synthetic peptide with a sequence comprising C3 amino acids K1187-A1214, suggesting that this region in C3d is involved in its interaction with the receptor. We have also detected kinetic differences between the binding of iC3b and C3d to CR2. Finally, we observed direct binding of CR2 to the C3c fragment, which suggests that the regions of iC3b and C3d involved in their interaction with CR2, in addition to common contact sites, may involve regions that differ from each other.

Materials and Methods

Recombinant proteins

In this study we have expressed a truncated form of EBV gp350/220 (gp350/470t), the ectodomain of CR2 (soluble CR2 containing SCR 1–15), and soluble CR2 in fusion with biotinylation tag (CR2-BCCP) using the baculovirus system. The cloning procedure for the expression of soluble CR2 in the baculovirus system has been described elsewhere (32). This clone was a gift from Michael Holers (Department of Medicine and Immunology, University of Colorado Health Science Center, Denver, CO). The EBV gp350 (470t) and CR2-BCCP expression constructs were cloned as follows: to obtain gp350, the gp350 (470t) cDNA was amplified by the PCR from the pGEM-gp350 clone (33) using the primers 5’-TCCG GA TCCA ATG GAG GCA GCC GTG ATG TCT AGT C-3’ (forward) and reverse. The resulting product was cloned into the pVT-Bac vector (Life Technologies, Gaithersburg, MD) grown in suspension cultures. The biotinylated biotin carboxyl carrier protein (BCCP) from Escherichia coli (35). The construction of this clone involved several steps. First, total DNA was extracted from an overnight culture of E. coli DH5a cells. The cDNA encoding the BCCP protein was then amplified by PCR using the primers 5’-AAC TGA CCT GAT GGA AGC GCC AGC ACG A-3’ (forward) and 3’-ATT AGC TAG CCT ACT CGA TGA CGA CCA GCG GC-5’ (reverse) and cloned into the pVT-Bac vector (pVT-Bac-BCCP). The CR2 cDNA was then PCR-amplified from the pGEM-CR2 clone (32) with primers 5’-AAT TCC CGG GAT GGG CCC GCG GGC GCC CCT G-3’ and 5’-CAC GAG CTC TGA ACG GGA TTC TCA AAC-3’ and cloned into the pVT-Bac-BCCP vector, in-frame with the BCCP protein. The resulting gp350 (470t) and CR2-BCCP constructs were recombinant baculovirus (Autographa californica nucleopolyhedrovirus) by cotransfection with Baculogold DNA (PharMingen, San Diego, CA). Recombinant plaques were isolated and amplified, and the resulting culture supernatants were screened for protein expression by SDS-PAGE and Western blotting. Baculovirus recombinants expressing the protein of interest were subjected to two additional rounds of plaque purification. For production of larger quantities of protein, Spodoptera frugiperda S9 cells (Life Technologies, Gaithersburg, MD) grown in suspension cultures were infected at a multiplicity of infection of 4. Forty-eight to 72 h after infection the supernatants were cleared by centrifugation, concentrated, and then dialyzed against PBS.

The baculovirus-expressed recombinant proteins were purified by using Ni-NTA agarose or mAb HBs affinity columns. The His-tagged protein gp350 (470t) was affinity-purified over an Ni-NTA-agar column. The bound protein was eluted with increasing concentrations of imidazole (0.01–0.25 M) in 0.02 M phosphate buffer (pH 7.5) with 0.5 M NaCl. The other two recombinant proteins (CR2 and CR2-BCCP) were affinity-purified over a Sepharose column covalently coupled to mAb HBs specific for CR2 (American Type Culture Collection, Manassas, VA). Soluble CR2 was further purified over a protein G-Sepharose column to remove traces of HBs Ag that might have leaked from the column. The eluates from all purifications were dialyzed against PBS and concentrated using the Amicon (Beverly, MA) ultrafiltration system.

The biotin holoenzyme synthetase protein, BirA, catalyzes the incorporation of biotin into BCCP. This protein was expressed in E. coli using a cloning method that was provided by Dr. D. Beckett (Department of Chemistry and Biochemistry, University of Maryland, College Park, MD) (36). The expressed protein was affinity-purified over a NicNTA agar column by elution with increasing concentrations of imidazole (0.01–0.25 M) in 0.02 M phosphate buffer (pH 7.5) with 0.5 M NaCl. The eluted protein was dialyzed overnight into a buffer solution containing 200 mM KCl with 50 mM Tris and 5% glycerol (pH 7.5).

Serum proteins

C3 (37) and factors I (38) and H (39) were purified from human serum as previously described. To obtain C3b or C3c, C3d was cleaved at 37°C with 1% w/w trypsin for 10 min, or 5% w/w trypsin for 1 h, respectively (40). The reactions were stopped by adding a 3-fold (w/w) excess of soybean trypsin inhibitor over the amount of trypsin used. To obtain iC3b, 500 μg of C3b was incubated with 150 μg of factor H and 64 μg of factor I for 4 h at 37°C. C3b, C3c, and iC3b were purified on a Mono Q column and gel filtration (Pharmacia, Piscataway, NJ) as previously described (41, 42).

Pepitide synthesis, purification, and characterization

The peptide with the C3 sequence K1187-A1214 was expressed by using the C3 synthetic peptide (mole 4313) using Fmoc amide resin (4'-2'-dimethoxyphenyl-Fmoc-aminomethoxy)-phe- nyl resin. The side chain-protecting groups were Asp (tBu), Arg (Pmc), Thr (tBu), Gln (Trt), and Trp (Boc). Biotin (1 mmol) was dissolved in equal
volumes of DMSO and N-methylpyrrolidone, placed in an amino acid cartridge, and activated according to Applied Biosystems User Bulletin no. 35 before carrying on the peptide synthesis. The peptides were cleaved from the resin by incubation for 3 h at 22°C in a solution containing 5% phenol, 5% thioanisole, 5% water, 2.5% ethanediol, and 82.5% trifluoroacetic acid (TFA). The reaction mixture was filtered through a fritted funnel, precipitated with cold ether, dissolved in 50% acetonitrile containing 0.1% TFA, and lyophilized. The crude peptides obtained after cleavage were dissolved in 10% acetonitrile containing 0.1% TFA and purified using a reversed-phase C-18 column (Waters, Milford, MA). The purity of all peptides was monitored by analytical chromatography on a reversed-phase C-18 column and by laser desorption mass spectrometry (43).

Protein sequencing and mass spectrometry

To obtain the NH2-terminal sequence of gp350 (470t), the protein was subjected to electrophoresis and electroblotted onto a Problot membrane (PerkinElmer-Applied Biosystems, Foster City, CA). After the bands were excised from the membrane, automated Edman degradation was performed with the PerkinElmer-Applied Biosystems model 473A Protein Sequencer equipped with on-line phenylthiohydantoin-amino acid analysis, using programmed chemistry cycles and HPLC operation programs provided by the manufacturer. For mass spectrometric analysis, gp550 (470t) and soluble CR2 were desalted using a matrix para-crystalline method previously described (43) and then subjected to matrix-assisted laser desorption ionization-mass spectrometry analysis using a Micromass (Beverly, MA) ToySpec 2E time-of-flight mass spectrometer (1.0 m flight tube) equipped with a nitrogen laser (337 nm).

Site-specific biotinylation of CR2-BCCP and C3

To incorporate biotin at a single residue on CR2-BCCP and C3, the following strategies were used: 1) CR2-BCCP labeling was achieved in the presence of the BirA protein, which catalyzed the reaction; and 2) the labeling of C3 and its fragments was performed using the sulphhydryl-reactive EZ-Link polyethylene oxide (PEO)-maleimide-activated biotin (Pierce, Rockford, IL).

For labeling, CR2-BCCP was first dialyzed using 40 mM Tris-Cl, 5.5 mM MgCl2, and 100 mM KCl (pH 7.3) at 4°C and then added to the C3 protein suspension at a 5:1 molar ratio. The reaction mixture was incubated for 30 min at room temperature, and then the sample was desalted and exchanged into PBS twice over a PD-10 column (Amersham Pharmacia Biotech, Uppsala, Sweden) gel filtration chromatography and extensive dialysis against PBS. The specificity of this reaction was determined by using soluble CR2 as a control.

Human C3 and C3 fragments were biotinylated as follows. The thioester bond of native C3 was hydrolyzed by incubating 7.8 mg of C3 (43 μg) with an equal volume of a 0.2 M solution of methanethiol (pH 7.3) over night at 37°C (44–46). The buffer was then exchanged into a solution consisting of 0.1 M NaH2PO4 and 5 mM EDTA (pH 6.3) by using a PD-10 column (Amerham Pharmacia Biotech). EZ-Link PEO-maleimide-activated biotin (Pierce) was dissolved in PBS at a concentration of 10 nM and then added to the C3 protein suspension at a 5:1 molar ratio. The reaction mixture was incubated for 30 min at room temperature, and then the sample was desalted and exchanged into PBS twice over a PD-10 column to remove the free biotin. The C3 fragments C3b, iC3b, and C3d were obtained by limited tryptic cleavage of the same stock of biotinylated C3 protein as described above. Monitoring of the biotinylation reactions, cleavages, and purifications was performed with SDS-PAGE and Western blotting.

Western blot assays

Proteins were separated on SDS-PAGE and electrotransferred to a nitrocellulose membrane. Nonspecific binding to the membrane was prevented by incubation in blocking buffer (PBS containing 10% milk). For CR2 detection, the membranes were incubated with a monoclonal anti-CR2 Ab (HB5, 2 μg/ml); gp350 (470t) was detected with a 1/5000 dilution of an affinity-purified goat C-terminus C3d Ab (Invitrogen, Carlsbad, CA) or a 1/500 dilution of a polyclonal anti-EBV Ab (provided by Dr. Michael Holers). Bound Abs were detected with a peroxidase-labeled goat anti-mouse IgG Ab (Bio-Rad, Richmond, CA), and bound polyclonal Abs were detected with a peroxidase-labeled mouse anti-rabbit Ab (Bio-Rad). The incorporation of biotin was verified by reactivity with peroxidase-labeled streptavidin (Life Technologies). All of these incubations steps were performed at 1 h at 22°C, and the membranes were washed with PBS containing Tween 20 (0.05%) between each incubation step. The proteins were detected using the ECL kit (Amersham Pharmacia Biotech).

**SPR measurements**

The kinetics of CR2 binding to its ligands C3d, iC3b, gp350, C31187–1214, and C3c were determined on the SPR-based biosensor BIAcore X (BIACore, Piscataway, NJ). All the experiments were performed at 25°C in phosphate buffer containing 0.05% Tween 20 (pH 7.4), either at physiologic ionic strength (150 mM NaCl), or in 75 mM NaCl. To orient C3d, C3c, iC3b, and C31187–1214 peptide onto the chip and thus obtain a homogeneous ligand surface, the proteins were biotinylated at a specific residue as previously described and then immobilized on a streptavidin chip (Sensor Chip SA; BIACore). C3c was immobilized on a CM5 chip by using amine-coupling chemistry (Sensor Chip SA). One hundred forty resonance units (RU) of CR2, 250 RU of C3d, 350 RU of iC3b, 2800 RU of C3c, or 300 RU of C31187–1214 peptide were immobilized on flow cell two (Fc2). Flow cell one (Fc1) was used as a blank control, except for the binding experiments involving the C31187–1214 peptide, in which a biotinylated control peptide was immobilized on Fc1. Binding was measured at 50 μl/min to avoid mass transport effects. At this flow the initial on-rate was maximum. Flow was allowed to occur for several seconds to establish a baseline, and then various concentrations of analyte were injected and the association was followed for 120 s. At that time, sample was replaced with running buffer, and the dissociation of the complex was monitored for 240 s. The sensor chip was regenerated with brief pulses of 0.2 M sodium carbonate (pH 9.5). The activity of the immobilized protein was monitored by observing the binding response of a single concentration of analyte throughout the experiment. Biosensor data for the control Fc1 were subtracted from those obtained for the immobilized proteins. Each binding experiment was performed at least twice, and the responses were averaged. Several buffer injections were also performed, averaged, and subtracted from Fc2 signal. BIAevaluation 3.0 (BIACore) software was used to analyze the binding data, using global fitting. Linear analysis of the binding data was obtained as previously described (46–48) by plotting DR/dt vs RU, where RU is the relative response of the biosensor at time t. The apparent equilibrium dissociation constant (Kd) was calculated from the equation Kd = koff/kon.

**Results**

**Characterization of the recombinant proteins**

To analyze the interaction of CR2 with its ligands, we expressed both the receptor and its ligand gp350 (470t) using the baculovirus system. Because previous studies had shown that EBV gp350/220, when truncated at residue 470, can inhibit EBV binding to CR2 (23), we chose to express this gp350 (470t) construct for use in the interaction analysis. A C-terminal His tag was added to this protein to aid in its purification, gp350 (470t) does not have a well-defined signal peptide. To verify whether its putative signal peptide had been processed, we sequenced its N terminus. Edman degradation yielded the sequence DPMEEAALIV, with DP being the amino acids added in the cloning process and MEAALIV the putative signal peptide. Thus, its putative signal peptide had not undergone cleavage upon secretion of the protein. The molecular size of gp350 (470t) as predicted from the amino acid sequence of the protein was 50 kDa. However, analysis by SDS-PAGE and Western blot revealed a band of 65–70 kDa (Fig. 1A). Mass spectrometric analysis showed a single homogeneous species and indicated that the gp350 (470t) had a total mass of 72 kDa (Fig. 1A). Because this protein contains 18 potential N-glycosylation sites, we conclude that the gp350 (470t) had a total mass of 72 kDa (Fig. 1).

Expression of soluble recombinant CR2 in the baculovirus system has been previously reported (32). Purified soluble CR2 was analyzed by SDS-PAGE, Western blotting, and mass spectrometry. The recombinant soluble CR2 had a molecular size of 116 kDa, (Fig. 1B) which does not differ substantially from the size deduced from its amino acid composition (111 kDa). It is possible that associated carbohydrates added 5 kDa of mass because CR2 contains eleven putative sites of N-glycosylation. The recombinant CR2-BCCP was analyzed by SDS-PAGE and Western blotting.
Purified CR2-BCCP migrated as a single band, as visualized by its reactivity with mAb HB5 (Fig. 1).

Generation and characterization of site-specific biotinylated proteins

To measure the affinity and binding kinetics of the interaction between CR2 and its ligands, we used SPR technology. In such studies, immobilization of the ligand is generally achieved by the use of amine-coupling chemistry. However, this approach produces heterogeneous surface ligands and precludes the measurement of homogeneous binding kinetics. To overcome this problem, we designed an enzymatic and a chemical strategy to incorporate biotin at a specific residue on CR2 and C3, respectively (Fig. 2). This strategy provided a means of orienting these proteins on a...
streptavidin sensor chip surface in a way mimicking their physiological orientation.

To allow for site-specific biotinylation on the CR2 molecule, we cloned and expressed a CR2 protein in fusion with aa 70–156 of the BCCP from E. coli (35) at its C terminus. BCCP is a subunit of the E. coli enzyme acetyl-CoA carboxylase and contains a single, specific K residue that acts as a biotin acceptor (35, 36, 51). Therefore, this construct allowed us to incorporate biotin at a single residue on the C-terminal tag of CR2. The CR2-BCCP was labeled with biotin on its C-terminal BCCP tag, in the presence of the catalyzing enzyme BirA and ATP. To verify that the reaction was specific, we used recombinant CR2 as a negative control. We then analyzed biotin incorporation into both molecules by Western blotting. As shown in Fig. 3, only CR2-BCCP reacted with peroxidase-labeled streptavidin, indicating that only the BCCP-tagged protein had incorporated biotin.

To orient the C3 fragments C3d and iC3b on a BLAcore chip, they were biotinylated at Cys988, which participates in the formation of the thioester bond. Biotin was incorporated into the thiol group after hydrolysis of the thioester bond with methanolic amine. Following biotin incorporation, the biotinylated C3 was cleaved into its degradation fragments iC3b and C3d. The iC3b and C3d were generated from the same stock of biotinylated C3 to ensure that any differences in our BLAcore data that we obtained for the two molecules were not related to heterogeneity at the biotinylation stage. To verify that only the -SH group of Cys988 had incorporated biotin, we analyzed the reactivity of C3d and iC3b with peroxidase-labeled streptavidin in a Western blot (Fig. 3). Biotinylated iC3b and C3d. Arrow indicates C3d position. B, Coomassie blue staining of biotinylated iC3b and C3d. Arrow indicates C3d position. C, Western blot of the biotinylated C3 fragments, probed with peroxidase-labeled streptavidin. The proteins were analyzed under reducing conditions.

**Materials and Methods**

iC3b and C3d (see **SDS-PAGE and Western blotting of the biotinylated CR2-BCCP and C3 fragments.** A, Biotin was incorporated into CR2-BCCP in a reaction catalyzed by the enzyme BirA. Lane 1, Western blot of biotinylated CR2-BCCP, detected with peroxidase-labeled streptavidin. Lane 2, Nonbiotinylated CR2 control. B and C, PEO-maleimide biotin bound to Cys988 at the C3 thioester site. Biotinylated C3 was thereafter cleaved into iC3b and C3d (see Materials and Methods). B, Coomassie blue staining of biotinylated iC3b and C3d. Arrow indicates C3d position. C, Western blot of the biotinylated C3 fragments, probed with peroxidase-labeled streptavidin. The proteins were analyzed under reducing conditions. (e.g., when C3d/iC3b are attached to complement-activating particles) can vary as a result of differential exposure of interacting residues. Thus we tested these possibilities by measuring the binding of iC3b and gp350 to CR2 immobilized on the sensor chip as well as the interaction of soluble CR2 with iC3b and C3d immobilized on the sensor chip.

When we examined the binding of gp350 (470t) to CR2 immobilized on the sensor chip, we found that the binding reaction was dose-dependent and saturable. Analysis of the binding data by global fitting analysis (BLAevaluation software 3.0) showed a close fit to 1:1 Langmuir binding model (A + B ↔ AB; Fig. 4A). A random distribution of residuals and a χ² value for this interaction of 1.8 indicated that this model describes well the experimental data.

Next, we evaluated the interaction of CR2 with its natural ligands, the C3 fragments C3d and iC3b. Experiments were performed in which soluble C3d and iC3b were allowed to bind to CR2 immobilized on the sensor chip. Only the kinetics of iC3b binding was further analyzed. The data corresponding to soluble C3d binding to immobilized CR2 were not reproducible and were not included in the present study. We also examined the converse arrangement, in which soluble CR2 was injected onto a chip containing bound C3d or iC3b. When we analyzed these binding reactions using the 1:1 model the global fit of the data was not possible. Although the residual plot for the fitted data and the χ² values were <2, the values obtained for the maximum response (Rmax) were lower than the predicted values, and even lower than the observed response values. Eliminating the refractive index (RI) parameter that ascribes a bulk to the interaction resulted in a χ² value >120. Linear transformation of the binding data showed nonlinear plots (Fig. 4, B–D), indicating that the binding follows complex models.

Data from our laboratory and others have indicated that two regions of CR2 (52, 53) and more than one region of C3d/iC3b (24, 25, 29, 31, 54) may be involved in C3d/iC3b-CR2 interactions. Thus, it is possible that two regions of C3d/iC3b may interact with two regions of CR2. To test this possibility, we tried to fit our data to a bivalent analyte model: AA + B ↔ AB; AAB + B ↔ AABB (Fig. 4). These data include binding of CR2 to immobilized iC3b and C3d and binding of iC3b to immobilized CR2. The latter experiment was performed in the presence of 75 mM NaCl in the binding buffer, which allowed us to observe better binding. For these analyses, the residual plot for the fitted data and the χ² values ranged from 1 to 3 and from 1.5 to 2.1, respectively, and the residuals were randomly distributed around the fit. Besides, the Rmax values were close to the predicted maximum response. Other models, such as a conformational change model, did not describe the data obtained. A two-step binding model describes well the binding data where two or more binding steps are affecting the binding equilibrium between CR2 and C3d or between CR2 and iC3b. The overlapping data, with the result of the global fitting analysis for a bivalent model for iC3b binding to immobilized CR2, are shown in Fig. 4B, and those for the binding of soluble CR2 to immobilized C3d or iC3b are shown in Fig. 4, C and D, respectively. The kinetic constants for the interaction of CR2 with its ligands that we obtained from the global fitting analysis are summarized in Table I.

**Binding of CR2 to a peptide corresponding to C3 amino acids K1187-A1214**

In light of our observation that C3d interacts with CR2 in a complex manner and the controversy regarding the involvement of the C3 region comprising aa 1187–1214 in CR2 binding, we analyzed the binding of CR2 to a synthetic peptide carrying this amino acid sequence.
sequence (24). The C3 1187–1214 peptide (24) was biotinylated on its N terminus and immobilized on the Fc2 of a streptavidin sensor chip. As control we used an irrelevant biotinylated peptide, which was immobilized on Fc1. The experiment was performed in the presence of 75 mM NaCl in the running buffer to allow for better detection. Binding of CR2 to the C31187–1214 peptide was dose-dependent (Fig. 5). Furthermore, the interaction was inhibited by soluble C3d, but not by a control protein (BSA) (data not shown). Although global fitting of the data was not possible, these data indicate that the C3 region, 1187–1214, binds to CR2.

Effect of NaCl on the kinetics of CR2-ligand interaction

Recent availability of the x-ray structure of human C3d (30) and mutation analysis of charged residues of C3d (31) have highlighted the importance of ionic interactions in C3d-CR2 interactions. To understand the influence of ionic contacts on the kinetics of CR2-ligand interactions, we have studied these interactions at various salt concentrations using SPR technology.

In the present study we measured the binding of CR2 to its ligands at physiologic (150 mM) and half-physiologic (75 mM) ionic strength buffers. In all the interactions analyzed in this study, decreasing the amount of NaCl in the binding buffer lowered by ~4- or 5-fold the quantity of C3 fragments or gp350, respectively, that was required to obtain a similar response in PBS (data not shown). Therefore, a decrease in the salt concentration apparently facilitates the binding of CR2 to its ligands, by increasing the affinity of these interactions.

We then attempted to fit these binding data to a kinetic model. The binding of gp350 to immobilized CR2 and that of CR2 to immobilized C3d and iC3b fit well with a 1:1 and a bivalent interaction model, respectively (data not shown). Therefore, these data suggest that the mechanism of binding of CR2 to C3d, iC3b, and gp350 is not affected by the ionic strength of the buffer. The K_D values obtained from these analyses are presented in Table I.

To better understand the significance of these values for the binding of CR2 to C3d, we have overlaid the sensograms for the interaction of soluble CR2 with C3d in the presence of either 75 or 150 mM NaCl. As shown in Fig. 6, the total binding of 475 nM CR2 was increased by lowering the salt concentration. We then split the total binding curves into the components of a bivalent model. This strategy allowed us to analyze the parameter T_50, which is the time in the association phase at which the encounter and final complexes are equimolar, as defined by Lipschultz et al. (55). T_50 at 75 mM NaCl was 40 s whereas at 150 mM NaCl T_50...
Data were collected with 75 mM NaCl present in the binding buffer. The concentration (nanomoles) of injected CR2, is indicated at right.

Binding of CR2 to C3c

The existence of kinetic differences between the binding of C3d and iC3b to CR2 led us to test the hypothesis that additional binding sites for CR2 may exist on C3c. C3c was immobilized on a sensor chip by amine-coupling chemistry, and soluble CR2 was allowed to interact with it. The experiment was performed under half ionic strength (75 mM NaCl) buffer conditions to allow better detection. Binding of CR2 to C3c was dose-dependent (Fig. 7A) and was inhibited by the presence of low molar excess of soluble C3c but not C3d (data not shown). The presence of traces of C3d on the C3c sample was examined with the reactivity of an anti-C3d monoclonal Ab (C3-19) in a Western blot assay (data not shown). The detection limit of this assay was 0.8 pmol of C3d, which on the C3c sample was examined with the reactivity of an anti-C3d Abs (C3-9, 133-H11) (Fig. 7B). All these data argue strongly against the possibility that the observed CR2 binding was due to the presence of traces of C3d in the C3c sample.

Effect of assay orientation on the iC3b-CR2 binding kinetics

To assess whether protein orientation had any effect on the kinetics of CR2-iC3b complex formation, we compared the binding of soluble CR2 to iC3b and vice versa, in the presence of 75 mM NaCl (Fig. 8). The initial association and dissociation phases occurred more rapidly when soluble CR2 bound to iC3b immobilized on the chip (Fig. 8A) than in the converse situation (Fig. 8B). This observation suggests that, indeed, experimental design may affect the interaction between CR2 and its ligands as a function of time.

Discussion

In the present study, we have used SPR technology to analyze the interaction of CR2 with its ligands, EBV gp350/220, C3d, and iC3b. Although the importance of these interactions is well established, the detailed nature of the molecular mechanisms involved is still unclear. Here, we have demonstrated that binding mechanism of CR2 varies with its ligands: its binding to gp350 follows a simple 1:1 model, whereas its binding to C3d and iC3b is more complex and suggests involvement of more than one site in these interactions. Furthermore, our data show kinetic differences between the binding of iC3b and C3d to CR2, which suggest that the regions of iC3b and C3d that are involved in the interaction with CR2 may not be the same or may be differently exposed in the two molecules.

![FIGURE 5. Binding of CR2 to C31187–1214 peptide. Overlay plot showing binding of soluble CR2 to immobilized C31187–1214 peptide. Several concentrations of CR2 were injected over a streptavidin sensor chip containing a biotinylated control peptide immobilized on Fc1, and biotinylated C31187–1214 peptide on Fc2. Data were collected with 75 mM NaCl present in the binding buffer. The concentration (nanomoles) of injected CR2, is indicated at right.](image1)

![FIGURE 6. Effect of NaCl concentration on the binding of CR2 to immobilized C3d as analyzed by SPR. Overlay plot of the binding of 475 nM CR2 to immobilized C3d in the presence of 75 or 150 mM NaCl in the binding buffer. BIAevaluation software was used to view the two binding components, $K_{D1}$ and $K_{D2}$ of these interactions. Thick lines represent the total binding. The thin solid lines represent the $K_{D1}$ and $K_{D2}$ at 150 mM NaCl; the thin dashed lines represent $K_{D1}$ and $K_{D2}$ at 75 mM NaCl.](image2)
In an attempt to mimic the actual in vivo orientation of these proteins, we have biotinylated them at key residues, i.e., the Cys 988 residue that participates in the thioester bond formation in C3 (45, 56), and the C-terminal end of CR2. The immobilization of biotinylated C3d/iC3b on a streptavidin surface (i.e., the BIAcore chip) was intended to simulate the deposition of C3 onto activating particles. Likewise, immobilization of a CR2 molecule containing biotin at its C terminus mimicked anchoring on the cell surface. Orientation of proteins in this way provided an advantage over the random amine-coupling method generally used in SPR-based assays, because it produced a homogeneous ligand surface that facilitated the measurement of homogeneous binding constants.

To study the interaction of CR2 with EBV gp350, we expressed a truncated form of gp350/220, consisting of residues 1–470. This truncated molecule had previously been shown to inhibit EBV binding to CR2 (23). The SPR data obtained for this CR2-gp350 (470t) interaction showed a close fit to 1:1 Langmuir binding model (Fig. 4). The apparent $K_D$ value obtained for this interaction was 45 nM (Table I). Affinities for the interaction of gp350/220 with CR2 have been previously determined by equilibrium binding in two different experimental settings. Tanner and coworkers (23) examined the ability of gp350/220 to inhibit the binding of 125I-labeled gp350/220 to CR2-expressing cells (Raji), and Moore and coworkers (57) studied the binding of soluble CR2 to gp350/220 by measuring the changes in mobility of the CR2-gp350/220 complexes in the ultracentrifuge as a function of ligand concentration. The $K_D$ values obtained by the two groups were 12 and 3.2 nM, respectively. The differences observed between these studies could reflect different affinities of CR2 for gp350/220 when present on the cell surface or in solution. In our case, the differences in $K_D$ values observed between the present study (CR2 anchored to the sensor chip; $K_D = 45$ nM) and the previous study (23) (CR2 present on cell surface; $K_D = 12$ nM) could be due to truncation of the protein used in our study. We used a protein truncated at residue 470, whereas the previous studies used the entire molecule. Therefore, these data suggest that other regions in gp350 may be directly involved or influencing its binding to CR2.

Our analysis of the binding between CR2 and its natural ligands C3d and iC3b revealed that these interactions do not follow a simple 1:1 binding model (Fig. 4). We eliminated surface heterogeneity by implementing the directed orientation of the immobilized ligands. Additionally, the fact that the gp350 binding data fit a 1:1 model both argue strongly against the influence of experimental artifacts in these data. Linear transformation of the association and dissociation data showed nonlinear plots (Fig. 4). These nonlinear binding curves could reflect multiple binding sites with different affinities, cooperativity, or more complex models. Global fitting analysis of the binding data suggested that the interaction between CR2 and C3d/iC3b follows a bivalent model (Fig. 4). It is possible that two regions of the C3d/iC3b molecule actually interact with two regions of the CR2 molecule; in this case the stoichiometry of the complex would be 1:1, which is in accordance with the previous findings of Moore and coworkers (57).

Several lines of evidence support a multiple site interaction model for C3d/iC3b binding to CR2. First, it has been shown that two different regions within SCR 1 and 2 of CR2 are involved in its interaction with C3 (22, 53). Furthermore, data from our laboratory and others have indicated that at least two regions of human C3d are involved in its interaction with CR2 (24, 29, 31, 54). However, involvement of the region in C3 comprising aa 1199–1210 in binding to CR2 has been disputed (29, 31). To reassess this interaction using SPR technology, we performed a direct binding assay. We observed that CR2 binds to a synthetic peptide with the C3 sequence $\text{KFLTTAKDLNRWEDPGKQLY}$.
The binding data could not be fit to any kinetic model, but it allowed us to observe that the dissociation of CR2 from C3\(^{1187-1214}\) appeared to occur much faster than that from C3d (in the same buffer conditions). This suggests that other regions in C3d may be involved in its interaction with the receptor, possibly stabilizing the complex. In fact, when we attempted to competitively inhibit the binding of soluble CR2 to immobilized C3d with the C3\(^{1187-1214}\) peptide, no inhibition of the CR2-C3d interaction could be observed (data not shown), suggesting that other regions on C3d also interact with the receptor. Alternatively, correct orientation of the peptide may be necessary for it to bind to CR2.

The data obtained in this study (Fig. 6) and in previous studies (31, 57) clearly indicate that the binding of C3d to CR2 is highly dependent on ionic interactions. SPR data obtained in the present study indicate the involvement of two components in the C3d-CR2 interaction (Fig. 4, right panel). To elucidate the influence of ionic interactions on these components we obtained \(T_{50}\) values (the time required to achieve an equimolar distribution of initial and final complexes) for binding of C3d to CR2 at physiologic and half-physiologic ionic strengths. We observed that lowering the amount of salt reduced the time required for the onset of secondary contacts in the interaction (Fig. 6). This finding suggests that some or all of the charged cluster of amino acids on C3d (E37, E39, E160, D163, and E166) that are known to influence CR2 binding when absent (31) may participate in the second component of the interaction. In the absence of these residues, binding of CR2 to C3d may have such fast association and dissociation rates that it may not be detected in assays that do not involve chemical cross-linking or SPR. In fact, our SPR data revealed that dissociation of CR2 from the C3\(^{1187-1214}\) peptide occurs much faster than its dissociation from C3d (Figs. 5 and 6). Alternatively, the charged cluster of residues in C3d may cooperatively contribute to the interaction.

A careful analysis of our SPR data indicates that the interactions of C3d and iC3b with CR2 are not identical. Although both interactions seemed to follow a complex model, there appear to be significant differences between the binding of soluble CR2 to immobilized C3d and iC3b. For example, the on-rates of both the first and second components of C3d and iC3b differed from each other by \(\pm 2\)-fold (Table I). Our results indicate that the interactions of C3d and iC3b with CR2, although similar, appear to involve some notable differences. This may be due to the participation of the CR2-interacting iC3b residues located outside the C3d region (25). We observed that CR2 interacted with C3c when the latter was immobilized in a sensor chip (Fig. 7). This result points toward the presence of CR2-interacting residues on the C3c fragment that may explain the differences observed between C3d and iC3b binding to CR2. Interestingly, this interaction could be inhibited by soluble C3c (data not shown), which suggests that the CR2 binding residues in C3c are exposed when in soluble C3c, i.e., immobilization of C3c did not significantly alter the conformation of these residues. High amounts (2800 RU) of C3c were immobilized in the chip in a random orientation, two factors that may induce surface heterogeneity. Therefore, we did not attempt to fit the CR2-C3c binding data to any kinetic model.

In addition to the presence of contact sites on C3c, other factors may contribute to differential binding of iC3b and C3d to CR2. Several residues in the C3d region of iC3b may be less accessible to CR2 as compared with those of C3d. Evidence suggesting that such steric hindrance may exist was presented in a study that used mAb 130 (58, 59). This Ab, which recognizes a neoantigenic determinant that is expressed when C3b is cleaved to iC3b and inhibits CR2 binding (24), was found to bind C3d better than iC3b (59).

Although it is clear from our data that C3d/iC3b-CR2 interactions do not follow simple 1:1 interactions and that the data show two components (Fig. 4, right panel) and fit well with a bivalent interaction model (Fig. 4, left panel), the complexity of the model precludes us from concluding that these are bivalent interactions, because increasing the complexity of a model increases the potential for multiple local minima in the \(\chi^2\) function that decrease the stability of the fitting procedure. However, we have shown in this study that multiple regions in C3 interact directly with the receptor. It is also conceivable that CR2 has different affinities for soluble or surface-bound C3 fragments (i.e., those bound to the activating particle at their thioester site). In this study we have examined the effect of ligand orientation on the binding kinetics of the iC3b-CR2 interaction. We observed differences in the on- and off-rates of iC3b, depending on whether it was oriented on the surface or present in solution. Complex formation and dissociation was initially faster when iC3b was immobilized than when it was presented in soluble form. Immobilization of iC3b through Cys\(^{988}\) which is involved in the thioester bond formation, may render the CR2-interacting residues more accessible. Similarly, differential expression of various epitopes on fluid-phase vs surface-bound C3 has already been proposed. Nilsson and coworkers have found that the reactivity of several mAbs recognizing the C3 \(\alpha\)-chain differ when the C3 fragments are in fluid phase or surface-bound (60).

In summary, we have used SPR technology to gain insight into the interaction between CR2 and its ligands. We have found that in contrast to gp350 (470t), C3d and iC3b appear to bind to CR2 in a complex manner.

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**References**