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Identification of the Hepatitis C Virus E2 Glycoprotein Binding Site on the Large Extracellular Loop of CD81
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The binding of hepatitis C virus glycoprotein E2 to the large extracellular loop (LEL) of CD81 has been shown to modulate human T-cell and NK cell activity in vitro. Using random mutagenesis of a chimera of maltose-binding protein and LEL residues 113 to 201, we have determined that the E2-binding site on CD81 comprises residues Ile182, Phe186, Asn184, and Leu162. These findings reveal an E2-binding surface of approximately 806 Å² and potential target sites for the development of small-molecule inhibitors of E2 binding.

CD81 is a member of the tetraspanin membrane protein superfamily, characterized by the presence of four transmembrane domains, three intracellular loops, and two extracellular domains, which in the case of CD81 are referred to as the small extracellular loop and the large extracellular loop (LEL). Tetraspanins appear to have an essential function in organizing signaling molecules at the cell surface by associating with other tetraspanins, lineage-specific proteins, integrins, major histocompatibility complex molecules, and signaling proteins in a cell type-dependent manner (for a review, see reference 8). The recent elucidation of the CD81 LEL crystal structure revealed that it is a homodimer, with each monomer being composed of five α-helices (A to E) arranged in a head subdomain (consisting of the last two turns of the A helix, the B, C, and D helices, and their interconnecting loops) atop a stalk subdomain (comprising antiparallel A and E helices) (6). Four cysteine residues, conserved in all tetraspanin sequences, participate in the formation of two disulfide bonds, Cys156-Cys190 and Cys157-Cys175, which stabilize the head subdomain.

Hepatitis C virus (HCV) encodes a ~3,300-amino-acid polyprotein from which the E1 (polypeptide residues 191 to 383) and E2 (residues 384 to 746) glycoproteins are cleaved cotranslationally. The mature forms of E1 and E2 are noncovalently associated, and each contains an N-terminal ectodomain and a C-terminal transmembrane domain. Recently, it was shown that recombinant soluble E2, E1-E2 complex, and HCV-like particles (12, 14) as well as HCV particles from infectious plasma (14) bind to the CD81 LEL with nanomole-level affinity (13, 14). Furthermore, the LEL can inhibit the binding of E2 to liver sections (13, 14) and of HCV-like particles to MOLT-4 T cells (21). The available evidence suggests that CD81 is unlikely to play a role in HCV entry (1, 4, 13, 16, 17, 21, 22). However, E2-CD81 ligation was recently found to induce in naïve and antigen-experienced T cells in vitro a costimulatory signal leading to the production of the proinflammatory cytokine gamma interferon (20). As HCV-associated liver damage is primarily due to a massive infiltration by activated proinflammatory lymphocytes (for a review, see reference 19), these findings raise the possibility that the CD81-E2 interaction plays a role in T-cell-mediated liver inflammation and pathology.

To further characterize the E2-binding residues on the LEL of CD81, we produced a protein chimera consisting of maltose-binding protein (MBP) fused to the N terminus of LEL residues 113 to 201 (MBP-LEL113-201) via a trialanine linker. MBP is monomeric and serves as an ideal scaffold for studying oligomerization determinants of heterologous proteins (2, 7). Superdex 200 gel filtration chromatography of amylose-agarose-purified MBP-LEL113-201 reveals two major species, corresponding approximately in molecular mass to the dimer (eluting at 92 min) and the monomer (100 min) (Fig. 1A). Nonreducing sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) of the MBP-LEL113-201 dimer and monomer peak fractions revealed an ~48-kDa major species and a trace amount (<5%) of an ~97-kDa band (Fig. 1A, inset), consistent with the virtual absence of disulfide-linked dimers. The ~97-kDa trace species is likely to be an artifact induced by boiling of samples in the absence of β-mercaptoethanol, as it was obtained from both the dimer and monomer peaks and was not detected by electrospray mass spectrometry (see below).

Sedimentation equilibrium analysis confirmed the molecular mass of the MBP-LEL113-201 putative dimer. The experimental data, plotted as the normal log of the optical density at 280 nm (OD280) versus the squared radius, results in a straight line that is consistent with the presence of a stable single solute (data not shown). The slope of the line and the observed molecular mass (± standard deviation) of 96,300 ± 4,500 Da closely approximate those of the dimer, whose theoretical mass is 100,518 Da. Next, the disulfide bond statuses of the MBP-LEL113-201 dimer and monomer were examined by treatment with the alkylating agent 4-vinylpyridine (4-VP) followed by mass spectrometry to detect the covalent modification of free sulphydryl groups (2). The molecular masses of the 4-VP-treated MBP-LEL113-201 dimer (50,264 Da) and monomer (50,260 Da) were unchanged relative to those of their untreated counterparts (50,265 and 50,262 Da, respectively), confirming that two intramolecular disulfide bonds had formed.

To examine the ability of the MBP-LEL113-201 dimer and monomer to bind HCV E2, we developed a solid-phase en-
zyme immunoassay (EIA) in which plate-bound MBP-LEL chimeras capture E2\textsuperscript{661}myc secreted from transfected 293T cells. Binding is detected by using the anti-myc monoclonal antibody (MAb) 9E10 and peroxidase-conjugated anti-mouse immunoglobulin G. Figure 1B shows that E2\textsuperscript{661}myc binds optimally to the MBP-LEL\textsuperscript{113-201} dimer; a substantially lower level of binding is observed for the MBP-LEL\textsuperscript{113-201} monomer. The dimeric form of MBP-LEL\textsuperscript{113-201} also captured noncovalently associated complexes of E1-E2 from transfected-cell lysates, as determined with the conformation-dependent MAb H53 (data not shown). The ability of the chimeras to inhibit the binding of solution-phase E2\textsuperscript{661}myc to plate-bound MBP-LEL\textsuperscript{113-201} dimer was used to rule out the possibility that the attachment of MBP chimeras to the solid phase had induced the formation of an E2\textsuperscript{661}myc binding site (Fig. 1C). The 50% inhibitory concentration (IC\textsubscript{50}) for the MBP-LEL\textsuperscript{113-201} dimer is approximately 10-fold lower than the IC\textsubscript{50} for monomeric MBP-LEL\textsuperscript{113-201} (30 \pm 10 nM versus 283 \pm 29 nM), suggesting that dimerization of the LEL enhances binding to E2 by approximately 10-fold.

A random mutagenesis approach was used to identify amino acid changes that resulted in the loss of E2-binding function in MBP-LEL\textsuperscript{113-201} as determined by the method of Lin-Goerke et al. (9). We used the EIA-based E2\textsuperscript{661}myc-binding assay to screen 300 transformants and identified four point mutants (N184Y, L162P, I182F, and F186S) that retained the ability to form dimers (Fig. 2A) but exhibited a decrease in (N184Y) or a complete loss of (L162P, I182F, and F186S) E2\textsuperscript{661}myc-binding function (Fig. 2B). In agreement with the solid-phase assay data, solution-phase N184Y dimer exhibited a lower affinity for E2\textsuperscript{661}myc than its wild-type counterpart, having an IC\textsubscript{50} of 766 \pm 251 nM, approximately 20-fold higher than the IC\textsubscript{50} for the wild-type dimer. Examination of these mutants by 4-VP mass spectrometry confirmed that these mutants had formed two intramolecular disulfide bonds, indicating that the L162P, I182F, N184Y, and F186S mutations are accommodated in the LEL fold (data not shown). Therefore, the ability of these mutations to inhibit LEL E2-binding function implies that Leu\textsuperscript{162}, Ile\textsuperscript{182}, Asn\textsuperscript{184}, and Phe\textsuperscript{186} are E2 contact residues. We next confirmed that these four mutations had similar effects on E2-binding function in the context of the entire tetraspanin molecule. We engineered a cytomegalovirus promoter-based expression vector (CD81C8), containing the CD81 open reading frame linked to a C-terminal MAb C8 epitope tag, for transient-transfection experiments. By using the anti-CD81 MAb 1.3.3.22 in flow cytometry, we first confirmed that the C8 epitope tag did not affect the cell surface expression of CD81 (data not shown). Mutations were introduced into CD81\textsubscript{C8}, and the expression and stability of the mutants in transfected HEK 293T cells were compared with those of the wild type by Western blotting with MAb C8. Figure 2C shows
FIG. 2. (A) Superdex 200 gel filtration of F186S, L162P, I182F, and N184Y mutants. Dotted lines represent elution times for the MBP-LEL\textsuperscript{113-201} monomer (98 min), dimer (92 min), and putative trimer (86 min). Symbols above peaks have been used to identify corresponding MBP-LEL species in panel B. V\textsubscript{v}, void volume. (B) EIA of E2\textsuperscript{661}myc binding to solid-phase peak Superdex 200 fractions collected in the experiment whose results are shown in panel A. Binding curves for the MBP-LEL\textsuperscript{113-201} dimer (solid line) and monomer (dashed line) are included. (C) Synthesis and stability of CD81\textsubscript{C8} mutants. Lysates of CD81\textsubscript{C8}-expressing 293T cells were subjected to reducing SDS-PAGE in 12 to 17% polyacrylamide gradient gels followed by Western blotting with MAb C8, directed to the C-terminal epitope tag. (D) Binding of E2\textsuperscript{661}myc to cell surface-expressed CD81\textsubscript{C8} mutants. Transfected CHO-K1 cells were incubated with serial dilutions of concentrated E2\textsuperscript{661}myc tissue culture fluid, washed, and then incubated with radioiodinated MAb 9E10. After being subjected to further washing, the cells were lysed and their radioactivity was quantitated in a Packard Autogamma counter. Relative E2\textsuperscript{661}myc binding is expressed as follows: (c.p.m. bound to cells expressing mutant CD81\textsubscript{C8})/c.p.m. bound to cells expressing wild-type CD81\textsubscript{C8}) × 100. The results are representative of two independent transfections. The inset shows the relative cell surface-expression of CD81\textsubscript{C8} mutants. Intact CD81\textsubscript{C8}-expressing CHO-K1 cells were incubated with \textsuperscript{125}I-labeled MAb 1.3.3.22 for 1 h on ice and then washed prior to quantitation of radioactivity. Relative \textsuperscript{125}I-labeled MAb 1.3.3.22 binding is expressed as follows: (c.p.m. bound to cells expressing mutant CD81\textsubscript{C8})/c.p.m. bound to cells expressing wild-type CD81\textsubscript{C8}) × 100. The means ± standard deviations of data from three independent transfections are shown.
that the mutants and the modified wild-type CD81cs were expressed in transfected-cell lysates at similar levels, migrating to a position corresponding to a molecular mass of 26 kDa.

The abilities of wild-type and mutated CD81 molecules to bind secreted E2<sup>661</sup>myc were compared by using a surface to a position corresponding to a molecular mass of 26 kDa. expressed in transfected-cell lysates at similar levels, migrating T163A, at an adjacent residue of the African green monkey CD81 has been shown to enhance E2 binding when placed in the human CD81 sequence (5), suggesting that the Leu<sup>162</sup> region modulates E2 binding. The involvement of Leu<sup>162</sup> in E2 binding extends the molecular surface of the binding site from 591 Å<sup>2</sup> (Thr<sup>166</sup>, Ile<sup>181</sup>, Ile<sup>182</sup>, Asn<sup>184</sup>, Leu<sup>185</sup>, and Phe<sup>186</sup>) to 805 Å<sup>2</sup>.

An inspection of E2-binding sites in the context of the human CD81-LEL dimeric structure reveals that one site is rotated 90° relative to the other. The orientation of the LEL dimer such that an E2-binding site projects out from the plane of the membrane (Fig. 3; see also Fig. 4 in reference 6) indicates that the E2-binding site of the partner monomer faces the membrane, making it unavailable for E2 ligation. Such a relationship between binding sites in native CD81 would preclude bivalent binding of CD81 to E2, particularly if virion-associated E2 molecules are related by dyad symmetry, as is the case with the flavivirus counterpart, the E glycoprotein (15, 23). However, such a relationship between binding sites may allow asymmetric binding of E2 and other members of the tetraspanin web to LE/L.

The results of recent in vitro studies indicate that the E2-CD81 interaction has immunomodulatory implications including enhanced production of the proinflammatory cytokines gamma interferon and interleukin-4 (20), down-regulation of T-cell receptors (20), and suppression of NK cell activity (3, 18). These findings raise the possibility that E2-CD81 ligation contributes to an HCV-induced liver immunopathology that is characterized by large numbers of liver-infiltrating activated T cells. Our identification of the E2-binding footprint in the context of the human CD81-LEL crystal structure provides new targets for the design of small-molecule inhibitors of the E2-CD81 interaction.

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