Cross-Link Formation between Mutant Galectins of Caenorhabditis elegans with a Substituted Cysteine Residue and Asialofetuin via a Photoactivatable Bifunctional Reagent

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LEC-1 is the first tandem repeat-type galectin isolated from an animal system; this galectin has two carbohydrate recognition domains in a single polypeptide chain. Because its two lectin domains have different sugar-binding profiles, these domains are thought to interact with different carbohydrate ligands. In our previous study, we showed that a mutant of LEC-1 in which a cysteine residue was introduced at a unique position in the N-terminal lectin domain (Nh) can be cross-linked with a model glycoprotein ligand, bovine asialofetuin, by using a bifunctional photoactivatable cross-linking reagent, benzophenone-4-maleimide. In the present work, we applied the same procedure to the C-terminal lectin domain (Ch) of LEC-1. Cross-linked products were formed in the cases of two mutants in which a cysteine residue was introduced at Lys177 and Ser268, respectively. This method is very useful for capturing and assigning endogenous ligand glycoconjugates with relatively low affinities to each carbohydrate recognition domain of the whole tandem repeat-type galectin molecule.

Key words  galectin; LEC-1; ligand; cross-link; maleimide; benzophenone

Galecins are a group of animal lectins characterized by their specificity for β-galactosides and an evolutionarily conserved sequence motif in the carbohydrate-binding site. Galectins are involved in a wide variety of biological phenomena, including development, cell differentiation, tumor metastasis, apoptosis, RNA splicing, and regulation of immune functioning. Galectins can be classified into three types on the basis of their molecular architecture: proto, chimera, and tandem-repeat types. LEC-1, the 32-kDa galectin of the nematode, Caenorhabditis elegans, was the first example of a tandem repeat-type galectin composed of two homologous regions.7,8)

A variety of pyridylaminated oligosaccharides were used as analytes in frontal affinity chromatography,9) and the result showed that the two lectin domains of LEC-1 have different sugar-binding profiles.10) This finding suggests that the endogenous ligand glycoconjugates for the two domains of LEC-1 differ. However, it is difficult to confirm this finding, especially when the binding ability of each domain is not adequately strong. We developed a new approach to overcome this problem. We prepared LEC-1 mutants in which a cysteine residue was introduced using site-directed mutagenesis of the LEC-1 cDNA (LEC-1 contains no cysteine residues).8) The following oligonucleotide primers were used for mutagenesis (substitution sites are underlined): E153C, 5'-ACCCGGTACCATACGAGCCGACTGCGCAACAGGAC-3'; K177C, 5'-GTTGAGAAGAAGGCATGCGCTTCCACGTCAACCTTCTCCGC-3'; D199C, 5'-TCCACCTTACCCGCGGTCTTGTAGAAGCAATCGTAC-3'; N211C, 5'-AACCTCTTGGCCGCATGCGAATGGGGAATATGGAGGCCTGGAGGA-3'; and S268C, 5'-ACATTCGGGCTGAGATCTGTGGAGAAATGGAGCGC-3'.

MATERIALS AND METHODS

Chemicals  Rabbit anti-bovine fetuin polyclonal antibody was purchased from Chemicon International (Temecula, CA, U.S.A.). BPM and bovine fetuin were purchased from Sigma (St. Louis, MO, U.S.A.). Sialidase and β-galactosidase were purchased from Seikagaku Co. (Tokyo, Japan).

Construction of Mutant Lectin Genes  A cysteine residue was introduced using site-directed mutagenesis of LEC-1 cDNA (LEC-1 contains no cysteine residues).11) Residues subjected to substitution were selected on the basis of the X-ray crystal structure of LEC-1 in complex with galactose.13) To facilitate cross-link formation, hydrophilic amino acid residues with side chains extending from the surface of the protein toward the bound galactose but not contacting it were selected. Site-directed mutagenesis was performed as described previously.11,12) The following oligonucleotide primers were used for mutagenesis (substitution sites are underlined): E153C, 5'-ACCCGGTACCATACGAGCCGACTGCGCAACAGGAC-3'; K177C, 5'-GTTGAGAAGAAGGCATGCGCTTCCACGTCAACCTTCTCCGC-3'; D199C, 5'-TCCACCTTACCCGCGGTCTTGTAGAAGCAATCGTAC-3'; N211C, 5'-AACCTCTTGGCCGCATGCGAATGGGGAATATGGAGGCCTGGAGGA-3'; and S268C, 5'-ACATTCGGGCTGAGATCTGTGGAGAAATGGAGCGC-3'.

Purification of Recombinant Galectins  Recombinant proteins were expressed and purified as described previously.11,12)
Preparation of Asialofetuin for Cross-Linking Bovine fetuin was treated with sialidase to remove the sialic acid attached to the non-reducing galactose end residues of the sugar chain. The resultant asialofetuin was purified as described previously \(^{11,12,14}\) and used as a model glycoprotein ligand for LEC-1.

Covalent Cross-Linking of Mutated LEC-1 Proteins to Asialofetuin Cysteine-introduced LEC-1 was cross-linked to the model glycoprotein ligand, asialofetuin, as described previously. \(^{11,12,14}\) In brief, purified recombinant LEC-1 or the single cysteine-containing mutant of LEC-1 was dialyzed to the model glycoprotein ligand, asialofetuin, as described previously. \(^{11,12,14}\) In brief, purified recombinant LEC-1 or the single cysteine-containing mutant of LEC-1 was dialyzed against phosphate-buffered saline (PBS) containing 1 mM ethylenediaminetetraacetic acid (EDTA) (pH 7.2) (EDTA-PBS). BPM was dissolved in dimethylformamide, and the solution was added to the lectin solution (final concentration, 1 \(\mu\)M BPM). For negative controls, dimethylformamide was added instead of the BPM solution. After incubation in the dark for 30 min at 23 °C, unreacted BPM was quenched by the addition of dithiothreitol (final concentration, 20 mM). Asialofetuin in EDTA-PBS was added, and the reaction mixture was irradiated with a long-wavelength ultraviolet light for 3 min at 4 °C. The cross-linked products were analyzed by western blotting using a rabbit polyclonal antibody against LEC-1 \(^{15}\) and a rabbit polyclonal antibody against fetuin.

RESULTS

Construction of LEC-1 Mutants Containing Substituted Cysteine Residues In the current study, we investigated whether the model glycoprotein ligand asialofetuin is actually recognized by the C-terminal lectin domains of LEC-1 mutants that have a substituted cysteine residue near the binding site of the C-terminal lectin domain and whether they can be cross-linked to asialofetuin via BPM.

Each of the cysteine-substituted mutants was adsorbed to the asialofetuin-Sepharose and could be eluted with 0.1 M lactose. This confirms that the mutants retained their binding ability (data not shown). The purity of each recombinant protein was checked by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE). In the absence of other bands with higher molecular weights, if any band appeared in the higher-molecular-weight region after the cross-linking reaction, it was expected that the formation of cross-linked complexes would be easily detected.

Cross-Linking of Mutant LEC-1s to Asialofetuin BPM primarily reacts with the introduced sulphydryl group in the binding site region of the mutant LEC-1 via its maleimide moiety. \(^{15}\) Ultraviolet irradiation activates the benzophenone group at the other terminal, which can form covalent bonds with nearby groups. Therefore, if lectin binds a glycoprotein molecule, a covalently linked complex that can be easily detected on SDS-PAGE can be formed. The fact that no species with higher molecular mass were observed in a control experiment wherein wild-type LEC-1 was used suggests that although asialofetuin contains several cysteine residues that can react with the maleimide group of BPM, no cross-link formation occurred with the wild-type LEC-1. Therefore, any cross-linked products observed in the experiments using mutant LEC-1 should be attributable to BPM reacting with the introduced cysteine residues.

Five LEC-1 mutants containing one cysteine residue each at Glu \(^{153}\) (E153C), Lys \(^{177}\) (K177C), Asp \(^{199}\) (D199C), Asn \(^{211}\) (N211C), or Ser \(^{268}\) (S268C) were reacted with BPM in the dark, mixed with asialofetuin, irradiated with ultraviolet light, and analyzed by western blotting. As shown in Fig. 1, the antibody against LEC-1 recognized an additional major band with a molecular mass of 90 kDa was detected with antiserum against LEC-1 in mutants K177C and S268C, but not in the remaining mutants E153C, D199C, and N211C. The 90-kDa band was also detected with anti-fetuin antiserum. The molecular masses of marker proteins are indicated on the left. The black arrowhead shows the position of the LEC-1 mutants; the positions of asialofetuin (arrow) and the expected positions of K177C/S268C–asialofetuin heterodimers (white arrowheads) are shown on the right.

Effect of Lactose on Cross-Linking As shown in Fig. 2, in the presence of 0.1 M lactose, cross-linking between asialofetuin and mutants K177C and S268C was remarkably reduced. These results suggest that cross-linking occurs only when a specific lectin–sugar interaction exists. Furthermore, no cross-linked products were observed when asialofetuin treated with Streptococcus \(664K\) \(\beta\)-galactosidase was used instead of asialofetuin (data not shown). This indicates that the removal of terminal galactose moieties from glycoproteins results in the loss of specific interaction.
We previously performed a detailed analysis with a variety of fluorescence-labeled oligosaccharides in frontal affinity chromatography, and reported that the two lectin domains of LEC-1 have different sugar-binding profiles. The method reported in this paper should enable us to capture and characterize independent endogenous ligand glycoconjugates for each binding site of a lectin with multiple binding sites, even if the interactions between the lectin and its ligand glycoconjugates are relatively low and no other procedure is applicable. LEC-1 is considered to be bound to the β-galactoside structure of the endogenous ligand existing in the insoluble fraction because LEC-1 was extracted from C. elegans after lactose extraction to remove the bound endogenous galectins. Since cross-linking of LEC-1 mutants with their ligands via BPM and isolation from other proteins is applicable even when the isolation process requires severe conditions such as detergent treatment or use of denaturing agent while the interaction may be relatively weak, this procedure should enable us to isolate and assign the captured ligand glycoproteins by using mass spectrometry. The development of such methodology would certainly contribute greatly to understanding of the significance of carbohydrate recognition in biological systems.

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REFERENCES


