Amylose engineering: phosphorylase-catalyzed polymerization of functional saccharide primers for glycobiomaterials

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Interest in amylose and its hybrids has grown over many decades, and a great deal of work has been devoted to developing methods for designing functional amylose hybrids. In this context, phosphorylase-catalyzed polymerization shows considerable promise as a tool for preparing diverse amylose hybrids. Recently, advances have been made in the chemoenzymatic synthesis and characterization of amylose-block-polymers, amylose-graft-polymers, amylose-modified surfaces, hetero-oligosaccharides, and cellodextrin hybrids. Many of these saccharides provide clear opportunities for advances in biomaterials because of their biocompatibility and biodegradability. Important developments in bioapplications of amylose hybrids have also been made, and such newly developed amylose hybrids will help promote the development of new generations of glyco materials.

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INTRODUCTION

Amylose is a linear polysaccharide in which a large number of glucose units are linked via α(1,4) glycosidic bonds¹ (Figure 1(a)). Amylose is present in almost every plant system as an energy source² and is therefore one of the most abundant raw materials. It is biodegradable, biocompatible, and renewable. Amylose adopts a left-handed helical form in aqueous solution³ and can entrap hydrophobic molecules in the helix, which acts as a one-dimensional supramolecular host molecule.⁴–⁶ The interest in amylose and their hybrid molecules therefore has been growing because of their potential applications.

Polysaccharides are structurally well-defined molecules. This makes it difficult to synthesize polysaccharides containing amylose. In conventional chemical synthesis, the choice of substrates with appropriate leaving groups and protecting groups, catalysts, and experimental conditions (e.g., temperature and atmosphere) are crucial for obtaining stereo- and regiocontrolled oligosaccharides.⁷,⁸ Such methods also require multiple synthetic steps and are time consuming. Chemical synthetic approaches are therefore unsuitable for obtaining structurally defined amylose.

Biocatalytic synthesis is a promising method and has several advantages over chemical synthetic approaches, such as mild reaction conditions (e.g., room temperature and aqueous solution) and high stereo- and regioselectivity of glycosidic bonds.⁹–¹¹ In plants, amylose is synthesized from adenosine diphosphate (ADP)–glucose by granule-bound starch synthase. Subsequently,
a starch-branching enzyme cleaves the resulting amylose and transfers the cleaved glucan in an α-1,6 position. Pure amylose, therefore, can be obtained by the suppression of gene-encoding starch-branching enzymes. On the other hand, linear malto-oligosaccharide is synthesized from ADP–glucose by glycogen synthase (GS) in mammalian cells. However, the substrate is fragile, and cumbersome procedures are needed. Therefore, these methods are not suitable for preparing amylose in practice. Alternatively, α-glucan phosphorylases can be used for amylose preparation. α-Glucan phosphorylases naturally catalyze the hydrolysis of glycosidic bonds in α(1,4)-glucan in the presence of inorganic phosphate and the formation of glycosidic bonds in α(1,4)-glucan in the absence of inorganic phosphate (Figure 1(b)). In glycosidic bond formation, phosphorylase catalyzes the addition of a glucose unit from glucose monophosphate to the nonreducing end of an amylose primer, with a minimum degree of polymerization (DP) of 3. The reducing ends of amylose primers can therefore be freely modified. The DP of enzymatically synthesized amylose can be controlled by changing the glucose monophosphate/amylose primer ratio. The reaction is analogous to living polymerization; therefore, the molecular weight distribution of the amylose is narrow.

This method has been used to prepare amylose and amylose hybrids, including block and graft copolymers, amylose-grafted surfaces, and heterooligosaccharides, as shown in Figure 2. We have coined the term ‘amylose engineering’ to describe the generation of amylose-functionalized materials from amylose primers. An appropriate choice of amylose and other materials enables the addition of further functionalities and tuning and control of the materials for specific applications. Another aspect of this enzymatic polymerization is the entrapment of hydrophobic polymers in the amylose helix during polymerization (known as vine-twinning polymerization). This topic, however, has been reviewed recently; therefore, we will not discuss it in detail.

In this review, we provide an overview of the chemoenzymatic synthetic routes for the preparation of amylose and its hybrids using phosphorylase. We focus on the synthesis and characterization of linear and grafted architectures in which amylose units are attached as side chains to synthetic and natural polymer backbones and surfaces (see section Enzymatic Synthesis of Amylose Hybrids by Phosphorylases).

In the section Phosphorylase-Catalyzed Synthesis of Hetero-Oligosaccharides and Cellodextrins, we focus on the synthesis of hetero-oligosaccharides and cellodextrin by phosphorylase-catalyzed polymerization. Finally, some of the challenges in the application of these new amylose hybrid materials in biotechnology and therapeutics are highlighted (see section Bioapplications of Enzymatically Synthesized Amylose Hybrids).

ENZYMATIC SYNTHESIS OF AMYLOSE HYBRIDS BY PHOSPHORYLASES

Block Copolymers

As mentioned in the Introduction, phosphorylase catalyzes the addition of glucose units from glucose
monophosphate to the nonreducing end of an amyl-
ose primer (e.g., maltopentaose) in the absence of
inorganic phosphate. The reducing end of the primer
can be functionalized, and the resulting primer can
be recognized for phosphorylase. In 1987, Ziegast
and Pfannemüller first used this method for the
enzymatic synthesis of amylose hybrids.22 In their
seminal work, they prepared two amylose-chain-
functionalized poly(ethylene glycol)s (PEG); these
were the first amylose-based copolymers. It was
found that a bifunctional primer gave amylose block
copolymers with a uniform molecular weight distri-
bution of amylose. This was ascribed by the authors
to equal recognition of phosphorylases by both ends
of the primer.

We also synthesized amylose-b-mPEG using
phosphorylase-catalyzed polymerization23,24 (Figure 3
(a)). The double helix formation is a well-known
property of amylose chains, and it leads to the forma-
tion of large aggregates and precipitation (retrograda-
tion). Such low colloidal stability is a drawback in
practical applications. However, this polymer (the
amylose DP was 73) did not form a precipitate even
after 1 day in aqueous solution. Amylose aggregation
was effectively reduced by modification with water-
soluble PEG. Interestingly, the amylose-b-mPEG
copolymer formed reverse micelles in chloroform solu-
tion, and a hydrophobic dye (methyl orange) was
entrapped in the amylose helix in the reverse micelles.

Loos and Stadler reported amphiphilic
amylose-b-poly(styrene) polymers25 (Figure 3(b)),
which were prepared by coupling maltohepta-
lactone with amine-functionalized poly(styrene) by
reductive amination, followed by potato
phosphorylase-catalyzed polymerization. Although
the primers were insoluble in aqueous solution, the
enzymatic polymerization proceeded. This can be
attributed to the partial formation of micellar aggre-
gates.26,27 Recently, the same research group
used amphiphilic maltoheptaose-b-poly(2-vinyl
pyridine) as a primer.28

Phosphorylase-catalyzed polymerization can be
used not only with bifunctional primers but also with
trifunctional primers. Rachmawati et al. prepared three-
armed poly(tetrahydrofuran) (PTHF)-b-maltoheptaose
using a combination of cationic ring-opening polymeri-
zation and reductive amination.29 The resulting poly-
cmers can be used as primers for the enzymatic synthesis
of three-armed PTHF-b-amylose.

More recently, we developed a series of
amylose-based star polymers (with one, two, four,
and eight arms) using a combination of click chemis-
try and phosphorylase-catalyzed polymerization30
(Figure 4(a)). The uniform length of the elongated
amylose chains was confirmed using iodine encapsu-
lation experiments. The resulting polymers acted as
multivalent host molecules for hydrophobic mole-
cules such as oligo(phenylene vinylene) (OPV), curcu-
min, bispyrenylpropane (BPP), and lipids. The
encapsulation experiments showed that large elon-
gated molecules (more than ca 2 nm) were essential
for inclusion because various shorter, nonelongated
molecules, e.g., pyrene, 1-anilinonaphthalene-8-
sulfonic acid (1,8-ANS) and 2-anilinonaphthalene-6-
sulfonic acid (2,6-ANS), did not form complexes
(Figure 4(b)). It is worth noting that the Hill coeffi-
cients and binding constants increased with an
increasing number of arms when the polymer encap-
sulated a cationic lipid (C16SP). We attribute these
phenomena to neighboring helix formation in highly
branched polymers.

As already mentioned, amylose can wrap vari-
ous hydrophobic polymers, e.g., PTHF, poly(capro-
lactone), poly(tetramethylene carbonate), and
poly(lactide)s, during polymerization (vine-twining
polymerization). Tanaka et al. prepared
maltoheptaose-b-poly(l-lactide) (PLLA), which is a
primer–guest molecule hybrid, via a copper-catalyzed
1,3-dipolar cycloaddition.31 The elongated amylose
chains entrap other PLLA chains in their helical cav-
ities during enzymatic polymerization, resulting in
the formation of linear supramolecular polymers.
More recently, a three-armed Glc7-PLLA2 conjugate

![FIGURE 3](attachment:image.png)

(a) Chemical structure of amylose-b-mPEG and formation of reverse micelles in chloroform (a), and chemical structure of amylose-b-poly(styrene) and scanning force microscopy image of micellar aggregates of amylose-b-poly(styrene) in aqueous solution (b). (Reprinted with permission from Ref 26. Copyright 2005 American Chemical Society)
was used as a primer for the construction of supra-molecular polymers. After enzymatic polymerization, the resulting amylose–PLLA2 formed an ionic gel in 1-butyl-3-methylimidazolium chloride. The gelation was probably caused by the formation of inclusion complexes between the branched polymers. A similar approach was used to show inclusion complex formation by an amylose–PTHF conjugate.

**Branched Polysaccharides**

Glycogen is a highly branched molecule consisting of α(1,4) glucan units and is synthesized by the action of glycogenin (primer), GS, and glycogen-branching enzymes (GBEs). Glycogenin catalyzes the addition of several glucose units from uridine diphosphate-glucose. After the addition of 8–12 glucose residues, GS catalyzes the further addition of glucose units, and then, GBE cleaves the saccharide chains and transfers them from α(1,4) positions to α(1,6) positions. Loos et al. have developed an enzymatic synthesis of branched polysaccharides based on this biologically inspired synthesis. They used glucan phosphorylase and GBE, which introduces branching points. This tandem enzymatic polymerization successfully produced highly branched polysaccharides. They also used this strategy to obtain three-arm branched polysaccharides.

**Amylose-Grafted Polymers**

Phosphorylase-catalyzed polymerization is most frequently used for the synthesis of amylose-grafted polymers. To the best of our best knowledge, the first enzymatically synthesized amylose-g-polymer was reported by Husemann and Reinhardt in 1962. They synthesized acetobromo oligosaccharide-grafted 6-trityl-2,3-dicarbanilylamyllose. After removal of the protecting group, the resulting polymer served as a primer for producing amylose-grafted amylose using potato phosphorylase-catalyzed polymerization.

**Amylose-Grafted Synthetic Polymers**

This method has been used for various primer-graft-polymers (naturally occurring polymers and synthetic polymers). The first amylose-graft-synthetic polymer was reported by Stadler et al. They prepared two macro primers, namely, maltoheptaose-graft-poly(dimethylsiloxane), via either acetal or amide linkages. Although both macro primers were recognized by potato phosphorylase, their reaction rates were much lower than that of maltopentaose. The lower reaction rates were attributed to the low solubilities of the macro primers in aqueous solutions.

Kobayashi et al. synthesized two different types of amylose-graft-poly(styrene) via homopolymerization of an amylose-substituted styrene monomer (VAA) and enzymatic synthesis from maltopentaose-carrying poly(styrene) [poly(VM5A); Figure 5], and
investigated the structures of these polymers based on amylose–iodine complexations. The DP can be estimated from the maximum absorption wavelength ($\lambda_{\text{max}}$) of an amylose–iodine complex. The $\lambda_{\text{max}}$ value of the poly(VAA–iodine complex was comparable to that of the polyelectrolyte–iodine complex. This result suggests that the DP of poly(VAA) was nearly equal to that of amylose, and most of the primers were accessible to phosphorylase, enabling the production of VAA with a uniform amylose length. In contrast, the $\lambda_{\text{max}}$ of a poly(VAA-co-VM5A–iodine complex indicated an amylose chain length much greater than that estimated using a phosphate assay. The authors assumed that the amylose chains were elongated from a few primers because of the steric configuration of the polymer. As a result, the iodine complexes showed much higher $\lambda_{\text{max}}$ values than expected.

Amylose-graft-poly(acetylene), amylose-graft-poly(vinyl alcohol), and poly(styrene-block-(styrene-graft-amyllose) were also synthesized by phosphorylase-catalyzed polymerization.

**Amylose-Graft-Naturally Occurring Polymers**

Saccharide-conjugated peptides occur widely in living systems; for example, glycoproteins, peptidoglycans, and proteoglycans are found in plasma membranes, intercellular matrixes, and connective tissue, and they play important roles in biological activities. Inspired by such biological systems, Kamiya and Kobayashi reported amylose-graft-poly(l-glutamic acid) conjugates prepared by phosphorylase-catalyzed polymerization. This polymer formed helical structures in acidic aqueous solutions because poly(l-glutamic acid) is a helix-forming polymer. The helix-to-coil transition occurred at pH 6.1, which is higher than that of native poly(l-glutamic acid) (pH 4.5). This was ascribed by the authors to electrostatic repulsion between neighboring carboxylate side chains because of the surrounding amylose chains.

We synthesized a series of amylose-primer-modified cholesteryl poly(l-lysine) materials, which were prepared by coupling maltoheptaose and cholesterol-bearing poly(l-lysine) via reductive amination (Figure 6(a)). The resulting polymer self-assembled into nanometer-sized gel particles (i.e., a nanogel) of an average diameter of 50 nm and with a positive charge (19.6 mV). The nanogel showed a structural transition from a random coil to an $\alpha$-helical structure, induced by an increase in the pH. As expected, the nanogel acted as a primer for phosphorylase. The average diameter of the nanogel after enzymatic polymerization decreased to 30 nm. Interestingly, the zeta potential of the enzymatically polymerized nanogel (DP = 23) showed a weakly negative charge (~2.1 mV). These phenomena are attributed to the shielding of the cationic-charged surfaces by the elongated amylose chains (Figure 6(b)).

Natural polysaccharides are well-known structurally diverse molecules and include linear polymers and branched hetero-oligosaccharides (e.g., gum Arabic). Researchers have developed different types of branched hetero-oligosaccharides based on the...
structures of natural polysaccharides. The first example of the chemoenzymatic synthesis of a branched hetero-oligosaccharide was reported by Kadokawa et al. in 2007. They synthesized amylose-graft-chitosan, which is a linear polysaccharide consisting of α(1,4)-linked 2-amino-2-deoxy-β-D-glucan units, by coupling maltoheptaose with chitosan, followed by phosphorylase-catalyzed polymerization. Thus far, the synthesis of starch–chitosan hybrids by direct coupling of these polysaccharides is inefficient because of steric hindrance. These results suggest that the chemoenzymatic method is an efficient tool for the preparation of branched polysaccharides. The obtained polysaccharide hybrids were hardly soluble in any solvents because of helix formation by the amylose chains. The authors exploited this characteristic to prepare thin films and hydrogels consisting of branched polysaccharides. A more recent example, reported by the same group, is based on amylose-graft-cellulose. Amino-functionalized polysaccharides are easily accessible starting materials for producing amylose-graft-polysaccharides using reductive amination. To obtain amylose-graft-cellulose, amine-functionalized cellulose was prepared by treating tosylated cellulose with sodium azide, followed by reduction with sodium borohydride. The amylose-graft-cellulose was synthesized using the chemoenzymatic method described above. The polymers also formed thin films and hydrogels by simply drying. The gel was tougher than that formed from amylose-graft-chitosan. This chemoenzymatic method was recently extended to the preparation of amylose-graft-alginate, which is a linear polysaccharide consisting of (1,4)-linked β-D-mannuronic acid and α-L-gluconic acid units. Glycogen consists of linear chains of α(1,4)-linked glucose residues, which are interlinked by α(1,6) glycosidic bonds. The resulting structure is highly branched and multivalent. Because a number of nonreducing ends of α(1,4)-glucan are located on the surface, glycogen can act as a naturally occurring primer for phosphorylase. Kadokawa et al. performed α-glucan phosphorylase-catalyzed polymerization using glycogen as a primer and confirmed the elongation of amylose chains from glycogen. It was noted that the elongated amylose chains from the nonreducing ends of glycogen formed double helices among intermolecular glycogens, which acted as cross-linking points to produce a hydrogel.

**Amylose-Grafted Surfaces**

Because of its versatility, the chemoenzymatic approach is also suited for the modification of spherical and planar surfaces. The first example of an amylose-modified silica gel was reported by
Okamoto et al. in 1996.\textsuperscript{53} The modified silica gels were prepared using two different approaches (methods 1 and 2). In method 1, 3-aminopropyltrioxyxysilane-modified maltopentaose was immobilized by reaction with silica gel. In method 2, maltoheptanolactone was treated with 3-aminopropyl-silanized silica gel. The two amylose-modified silica gels were reacted with a large excess of 3,5-dimethylphenyl isocyanate to convert the amylose hydroxy groups to the corresponding carbamate residues. The difference between the amounts of amylose immobilized on the silica surfaces using the two methods was negligible. These modified silica can be used as chiral stationary phases in high-performance liquid chromatography because of their intrinsic chirality. They displayed excellent enantioseparation of 10 different racemates. Such systems are now widely used to separate racemic compounds into their enantiomers.\textsuperscript{54} Stadler et al. used a similar strategy to produce an amylose-grafted silica gel and investigated its chiral discrimination.\textsuperscript{55}

Another application of the chemoenzymatic method is the modification of two-dimensional metal and inorganic planar surfaces. Vlist et al. synthesized amylose and branched polysaccharide brushes on silica surfaces via enzymatic polymerization.\textsuperscript{56} More recently, the same research group used this strategy to modify a gold surface.\textsuperscript{57} Such polysaccharide-grafted surfaces have potential applications as anti-fouling materials for ship hulls, underwater pipes, and biomedical implants because of the highly hydrophilic nature of the polysaccharide.

**PHOSPHORYLASE-CATALYZED SYNTHESIS OF HETERO-OLIGOSACCHARIDES AND CELLODEXTRINS**

**Hetero-Oligosaccharides**

Enzymes usually show strict specificities for substrate recognition in catalytic reactions. However, some enzymes have loose substrate specificities. This enables the synthesis of unique hetero-oligosaccharide chains. Phosphorylases frequently have this property. Evers et al. reported that potato phosphorylase catalyzes the addition of D-glycal to maltotetraose and degrades the resulting 2-deoxy-maltooligosaccharide to 2-deoxy-\(\alpha\)-D-glucopyranosyl phosphate in the presence of inorganic phosphate.\textsuperscript{58} Since this discovery, various novel polysaccharide chains have been synthesized using phosphorylases (Figure 7).

![Saccharide primer](image)

**FIGURE 7** Phosphorylase-catalyzed synthesis of various hetero-oligosaccharides.
Kadokawa et al. first established this method for the preparation of hetero-oligosaccharides. They used xylose monophosphate as a substrate and maltotetraose as a primer. Matrix-assisted laser desorption/ionization time-of-flight mass spectrometry confirmed the formation of tetrasaccharides to octasaccharides containing one xylose unit and oligosaccharides consisting of two xylose units. Glucoamylase hydrolytic studies showed that xylose was added to the nonreducing end of the obtained hetero-oligosaccharide. The authors further extended this method to the preparation of N-formyl-α-glucosaminylated malto oligosaccharide, α-glucuronylated malto oligosaccharide, α-mannosylated malto oligosaccharide, chitin and chitosan stereoisomers, and non-natural heteroaminopolysaccharides.

A further example of enzymatic synthesis of a hetero-oligosaccharide was reported by the same research group. In this case, dendritic amphoteric α-glucans were prepared by sequential glucuronolation and glucosaminylation of highly branched cyclic dextrin (a glucan dendrimer). The zeta potentials of the polymers changed from positive to negative when the pH changed from acidic to basic. Such amphoteric polysaccharides could have useful bioapplications, such as carriers in drug-delivery systems and in regenerative tissue scaffolds.

**Cellodextrins**

Cellulose is a linear polysaccharide consisting of β(1,4)-linked glucose units and is the most abundant natural resource on Earth. Cellulose is synthesized by cellulose synthases, which are β-glycosyltransferase-type enzymes, at the plasma membrane. Low-molecular-weight cellulose (cellodextrin) can be prepared using the enzyme cellodextrin phosphorylase. Cellodextrin phosphorylase also catalyzes the addition of glucose units from glucose monophosphate to the nonreducing ends of cellobiose acceptors.

Serizawa et al. used this method to prepare post-functionalizeable two-dimensional crystalline cellulose nanosheets (Figure 8). An azide-containing cellulose oligomer was synthesized by reacting β-glucosyl azide as a primer and α-glucose monophosphate as a substrate in the presence of cellodextrin phosphorylase. The resulting cellulose oligomers (average DP 10) aligned and formed nanosheets of an average thickness of 5.5 nm. Wide-angle X-ray scattering and attenuated total reflectance Fourier-transform infrared spectroscopy studies showed that...
the nanosheet formed a cellulose II allomorph, which had an antiparallel saccharide arrangement against the sheet. The azide groups of cellodextrin were therefore located on the sheet surface. Such structures offer advantages such as the possibility of nanosheet post-functionalization. As proof of concept, the nanosheet was incubated with 1-ethynylpyrene as a model substrate in the presence of CuSO₄ and sodium ascorbate; a pyrene-conjugated nanosheet was successfully obtained. This method enables the fabrication of advanced functional cellulose sheets.

More recently, the same research group reported that a cellulose nanosheet acted as an artificial hydrolytic enzyme. The nanosheet was prepared by cellodextrin phosphorylase-catalyzed polymerization, as described above. When the resulting nanosheet was incubated with p-nitrophenyl acetate (PNP) as a model substrate, there was no significant difference in the presence and absence of the nanosheet. Previously, they showed that hydrolytic activities were derived from hydroxyl groups on the nanosheet. Significantly, they showed that enzymatic treatments such as cell signaling, transportation, and adhesion.

BIOAPPLICATIONS OF ENZYMATICALLY SYNTHESIZED AMYLOSE HYBRIDS

Considering the biodegradability and biocompatibility of amylose, it is not surprising that amylose hybrids have found use in numerous biological and medical applications. In the sections below, we discuss applications of amylose hybrids in protein science, DNA manipulation, and as carriers in drug-delivery systems.

Reconstitution of Membrane Proteins

Membrane proteins participate in pivotal biological processes such as cell signaling, transportation, and adhesion. However, there have been fewer studies of membrane proteins than of water-soluble proteins. This is because of their extremely low solubility in aqueous solution. We therefore need to embed membrane proteins in hydrophobic spaces such as lipid bilayers to study their use as functional materials. A typical method for the reconstitution of membrane proteins in liposomes is to solubilize the target proteins using a surfactant–phospholipid mixture and then remove the surfactant using dialysis or chromatography. This method, however, is generally time consuming and gives low yields of reconstituted membrane proteins. Alternative methods are therefore needed.

An example of a novel membrane protein reconstitution method based on enzymatically triggered micelle-vesicle transitions was reported by us in 2006. We synthesized a series of amylose primer-modified surfactants, which were prepared by the treatment of maltopentaose lactone and alkylamines (octyl, dodecyl, and hexadecyl). The surfactants self-assembled into micelles and acted as primers for phosphorylase. On enzymatic polymerization, the micelles collapsed (Figure 9(a)). The surface tension of the reaction solution increased and reached a plateau. The surface tension value after 150 min was comparable to that of the buffer solution. This was ascribed to the increased hydrophilicity of the surfactant caused by the elongation of saccharide chains, eventually leading to micelle dissociation. We further investigated enzymatically induced micelle-to-vesicle transitions in phospholipid–primer surfactant systems. A mixture of l-a-dipalmitoyl phosphatidylcholine and a primer surfactant formed micelles of a mean hydrodynamic diameter of 21 nm. Enzymatic polymerization caused an increase in the sizes of the assemblies. The resulting reaction mixture contained liposomes (123 nm), as confirmed by transmission electron microscopy. Liposome formation was attributed to the release of the surfactant from the mixed micelles as a result of an increase in the hydrophilicity of the surfactant. We used this system to reconstitute bacteriorhodopsin (BR), which is a model membrane protein, into liposomes. First, BR was solubilized in micelles consisting of a primer-modified surfactant. Second, the resulting micelles were mixed with the phospholipid–primer surfactant micelle solution. Finally, the mixture was incubated with phosphorylase for 6 h (Figure 9(b)). BR-reconstituted liposomes were successfully formed. The liposomes showed greater light-induced H⁺ influx and light responsiveness than did the mixed-micelle system.
hydrophobicity. Polymerization increased the surfactant hydrophilicity, and the resulting surfactant was removed from the CAB/surfactant complex. As a result, the CAB gradually refolded and recovered its activity. This system was also used to fold chemically denatured proteins and showed a higher efficiency than cyclodextrin as a surfactant-stripping agent.

More recently, other polysaccharide materials based on enzymatically synthesized glycogen (ESG)-bearing hydrophobic groups have been developed for artificial chaperone systems.\textsuperscript{72,73} The chaperone-like activity was greater than that of a previously reported system based on cholesteryl—group-modified pullulan. This is attributed to the unique architecture of hyperbranched ESGs, which provide sufficient space for protein refolding but not for irreversible aggregation of denatured proteins.

**DNA Chaperone System**

DNA strand exchange between double-stranded DNA (dsDNA) and single-stranded DNA (ssDNA) is an important biological process for genetic maintenance and DNA repair. Such processes are catalyzed by recombinases (e.g., Eukaryotic Rad 51, \textit{E. coli} RecA proteins). These enzymes, called DNA chaperones, bind with DNAs and form intermediate complexes with both dsDNA and ssDNA, resulting in the exchange of DNA strands. Artificial compounds can also catalyze DNA strand exchange.\textsuperscript{74} Maruyama et al. reported that poly-l-lysine-\textit{graft}-dextran significantly accelerates strand exchange reactions when used as an artificial DNA chaperone.\textsuperscript{74} The DNA chaperone activities of self-assembled cationic nanoparticles were also reported by our research group.\textsuperscript{75} These polymers have highly localized positive charges, which allow them to bind electrostatically with DNA strands, resulting in DNA strand concentration. Based on this work, we prepared a series of spermine-functionalized amylose star copolymers and investigated their artificial DNA chaperone activities.\textsuperscript{75} Strand exchange was accelerated in the presence of the polymers, and the rate of strand exchange clearly increased as the number of amylose arms increased. The increased rate is attributed to a decrease in the dissociation constant of DNA. This decrease is probably caused by an increase in the apparent cation concentration, resulting in enhanced DNA binding.

**Nucleic Acid-Delivery Systems**

Research on nucleic acid delivery has recently expanded because of its potential applications in next-generation therapeutic strategies for treating inheritable or acquired diseases. However, nucleic acids are easily digested by nucleases. In addition, nucleic acids are large anionic molecules, and they do not cross cell membrane. Consequently, the development of safe and efficient gene carriers is needed.
Among the materials under development for use as gene carriers, polysaccharides are one of the most promising because of their low toxicity, biocompatibility, and biodegradability.76–81 Our group has developed a gene-delivery system based on enzymatically synthesized polysaccharides. The system consists of cycloamylose (CA) functionalized with cationic groups. CA, which is a macrocyclic polysaccharide consisting of \( \alpha(1, 4) \)-glucose units, is produced through the treatment of linear amylose with \( 4\alpha \)-glucanotransferase. We reported a series of cycloamylose derivatives that effectively delivered plasmid DNA (pDNA),82 siRNA,83 and CpG DNA84 in vitro and in vivo.85 We also developed a novel gene-delivery system using hexadecyl-group-bearing cationic CA and a phospholipid-degrading enzyme.86

More recent examples of enzymatically synthesized polysaccharide-based gene carriers are amylose star copolymers.87 The molecular weight of siRNA is much smaller than that of pDNA, and a strong interaction is needed to form stable nanoparticle complexes. We therefore took advantage of the multivalency of star polymers. We prepared spermine-functionalized single- and eight-armed amylose-based star polymers (Figure 10(a)) and compared their siRNA carriers’ activities. As expected, the eight-armed polymer formed more compact complexes with siRNA than did the single-armed polymer. The siRNA/eight-armed polymer complexes were effectively internalized into Renca cells and suppressed mRNA levels by 65% (Figure 10(b)). This is probably because of the formation of smaller complexes between siRNA and the eight-armed polymer, which might enhance cell internalization, leading to higher gene-silencing efficiency.

**CONCLUSION**

Amylose synthesis by phosphorylase-catalyzed polymerization has been studied for more than three decades. Over the last 10 years, significant improvements not only in phosphorylase-catalyzed polymerization but also in coupling methods for primers have enabled the synthesis of a variety of amylose hybrids with different compositions, structures, dimensions, and chemical functionalization. Amylose hybrids with unique properties have been developed. However, there are few reports of bioapplications of amylose hybrids. This means that amylose-based materials are still a developing field. However, given the biocompatibility and biodegradability of amylose hybrids, and the possibility of additional functionalization, it is anticipated that amylose hybrids will find multiple applications in biomedicine in the near future.
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