Effective Separation for New Therapeutic Modalities Utilizing Temperature-responsive Chromatography

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In recent years, drug discovery and therapeutics trends have shifted from a focus on small-molecule compounds to biopharmaceuticals, genes, cell therapy, and regenerative medicine. Therefore, new approaches and technologies must be developed to respond to these changes in medical care. To achieve this, we applied a temperature-responsive separation system to purify a variety of proteins and cells. We developed a temperature-responsive chromatography technique based on a poly(N-isopropylacrylamide) (PNIPAAm)-grafted stationary phase. This method may be applied to various types of protein and cell separation applications by optimizing the properties of the modified polymers used in this system. Therefore, the developed temperature-responsive HPLC columns and temperature-responsive solid-phase extraction (TR-SPE) columns can be an effective separation tool for new therapeutic modalities such as monoclonal antibodies, nucleic acid drugs, and cells.

Keywords Temperature-responsive chromatography, poly(N-isopropylacrylamide), smart functional surfaces, new therapeutic modality, monoclonal antibody, oligonucleotide, cell separation

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1 Introduction

In the last two decades, the development of new drugs has shifted from a focus on small-molecule compounds to large-molecule biopharmaceuticals. Medical discovery and medical care tendencies have also changed, moving from an emphasis by small-molecule medicine to biopharmaceuticals, genes, cell therapy, and regenerative medicine.1,2 The importance of biopharmaceuticals and cell therapy in medical treatments is...
increasing. Today, seven of the ten best-selling pharmaceutics in the world are monoclonal antibodies.\(^3\) Therefore, new approaches and technologies must be developed to respond to these changes in medical care. However, to achieve the purification of biopharmaceuticals, organic, low-pH, or high-salt solvents are often employed, which sometimes results in a loss of the bioactivity of protein samples.\(^4\) Biological molecules are often separated using an ion-exchange chromatography in the medicine industry and reversed-phase chromatography and affinity chromatography, or combinations of these techniques. The high salt concentrations must be taken from the final product, imposing additional processing. Organic solvents have many disadvantages in terms of their cost, toxicity, flammability and disposal costs. The development of a novel separation and purification method for biopharmaceuticals, including antibodies and oligonucleotides, under mild conditions and with sustainable associated costs, is greatly required. Moreover, cells have recently been used in drug discovery, drug development, biomedical research, and biological technology and medical applications. Recently, various types of cell separation methods

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have been developed. Especially, a cell separation-method with maintaining cell activity is strongly required for cell transplantation.

We have been investigating a temperature-responsive chromatography system based on a poly(N-isopropylacrylamide) (PNIPAAm)-grafted stationary phase. PNIPAAm is a thermosensitive polymer that exhibits a lower critical solution temperature (LCST) at 32°C (Fig. 1). This system is advantageous because analytes can be separated using only an aqueous mobile phase under mild conditions since the surface properties of the stationary phase are controlled by only changing the column temperature. Despite a recent increased focus on environmental protection, analytical methods that use only aqueous solutions without using organic solvents have not been well studied. Achieving separation without using organic solvents not only allows separation without impairing the function of bioactive substances, but it is also environmentally friendly. Therefore, we applied a temperature-responsive separation system to purify a variety of proteins and cells. In this paper, utilizing temperature-responsive chromatography (TRC), we reviewed a green analytical method for effective separation for new therapeutic modalities.

### 2 Temperature-responsive Chromatography Utilizing a Functional Polymer Surface

Stimuli-responsive polymers that respond to heat, light, and pH have found interesting applications in chemical and biomedical fields, such as fluorescent polymer probes and drug delivery systems. PNIPAAm is one of the most representative polymers that exhibits reversible soluble-insoluble changes in the vicinity of its LCST of around 32°C in aqueous solution (Fig. 1). This polymer demonstrates good solubility in aqueous solutions at low temperatures, but separates from solution when the temperature is raised above the LCST. The LCST of PNIPAAm can be controlled by the composition of the copolymer. For example, the copolymerization of PNIPAAm monomers with hydrophobic monomers, such as n-butyl methacrylate (BMA), leads to an increase in the polymer hydrophobicity and a decrease in the LCST of the copolymer. The LCST shifts from 32 to 24°C as the mole fraction of BMA increases from 0 to 5 mol% in the copolymer (Fig. 1c). By contrast, the copolymerization of PNIPAAm monomers with more hydrophilic monomers, such as N,N-dimethylaminopropyl acrylamide (DMAPAAm), increases the polymer hydrophilicity while also increasing the LCST of the copolymer (Fig. 1d). These copolymers exhibit a reversible hydrophilic/hydrophobic phase transition at their LCSTs. To prepare the functional polymer surface for temperature-responsive chromatography, we used three types of modification methods for grafting the polymers onto silica beads, as shown in Fig. 2. These polymer-grafted surfaces demonstrate temperature-responsive hydrophilic-hydrophobic surface property alterations. Yakushiji et al. reported temperature-dependent wettability changes for PNIPAAm hydrogel modified surfaces by aqueous dynamic contact-angle measurements. The graft configuration of PNIPAAm produced from different grafting methods greatly influences temperature-dependent aqueous wettability changes. Using these features, we developed temperature-sensitive stationary phases for HPLC and solid-phase extraction (SPE). Unlike in conventional HPLC systems, such as reverse-phase columns in which the partitioning is controlled by the mobile phase composition, the partitioning in temperature-responsive chromatography systems is controlled by external temperature changes, and the aqueous solution is used as the mobile phase.

We designed and synthesized multi-functional polymers based...
on PNIPAAm to optimize the separation of various samples (Table 1). By introducing cationic or anionic comonomers in polymer chains as a function of the ion-exchange group, these copolymers respond to both the temperature and pH. For example, cationic DMAPAAm (pKα: 10.4) may be protonated in an aqueous solution at pH 7.4 while providing hydrophilicity to the copolymer. The phase-transition temperature of the polymer increases with increasing DMAPAAm composition (Fig. 1d). Hoffman et al. reported that the LCST of a NIPAAm-acrylic acid (AAc) copolymer increased with an increase in the pH of the used buffer solutions. Kobayashi et al. reported that the incorporation of selected moieties next to the temperature-responsive polymer increases with increasing DMAPAAm composition. As shown in Table 1, we synthesized a thermo-responsive polymer-modified beads then packed into the HPLC-column and used in the HPLC-system (TR-HPLC). Figure 3 shows the effects of amino acid introduction to polymer chains on steroid separation. In the separation of steroids with varied hydrophobicities, more hydrophobic steroids are retained longer. This indicates that π-π interactions occurred between the steroids and the phenyl or indole moieties of phenylalanine or tryptophan, respectively. In addition, the retention of compounds with hydrogen bond acceptors were higher in a poly(NIPAAm-Trp-OMe5%)-modified column, which contained indole as a hydrogen bond donor, than in a poly(NIPAAm-AAc-IAAc-IBAAAm) column, which contained indole as a hydrogen bond donor, than in a poly(NIPAAm-AAc-IIBAAAm) column, which contained indole as a hydrogen bond donor.
Simultaneous Analysis of Multiple Cytochrome P450s Substrates

Cytochrome p450s (CYP) comprise a superfamily of heme-thiolate proteins, and are best known for their central role in phase-I drug metabolism. They are of critical importance for overcoming significant clinical pharmacological problems, including drug interactions and interindividual variability in drug metabolism. The “cocktail” approach is often used in drug development to estimate the effect of more than one single experimental CYP enzyme of a drug candidate. A simultaneous analysis of more than one CYP substrate, which may have the very different structures and physicochemical properties, generally requires organic solvents and a mobile-phase gradient method in a conventional system. In contrast, when PNIPAAm-based temperature-responsive chromatography is used, a simultaneous analysis of cocktails of multiple CYP substrates and their metabolites may be achieved with isocratic and single mobile phases. In TR-HPLC, a good separation of six CYP substrates, such as phenacetin for CYP1A2, coumarin for CYP2A6, tolbutamide for CYP2C9, S-mephenytoin for CYP2C19, chlorzoxazone for CYP2E1, and testosterone for CYP3A4 were achieved utilizing a P(NIPAAm-co-BMA)-modified silica column, as shown in Fig. 4. As shown in Fig. 5b, good separation of the CYP3A4 substrate, midazolam, and its metabolite, 1′-hydroxymidazolam, was achieved using only an aqueous mobile phase without the requirement of any complicated gradient methods for the mobile phase. Interestingly, the order of elution of midazolam and its metabolite was changed at higher column temperatures, as shown in Fig. 5a. Figure 5c shows plots of the retention factor against the column temperature, obtained from Fig. 5a. It could be interpreted that the surface property alteration of the stationary phase at higher temperatures than the LCST caused changing interactions between the polymer surface and the analytes.

Recently, Maekawa et al. reported that five CYP substrates, such as caffeine for CYP1A2, warfarin for CYP2C9, omeprazole for CYP2C19, dextromethorphan for CYP2D6, and midazolam for CYP3A4, were analyzed utilizing a P(NIPAAm-co-Trp-OMe)-modified silica column. Introducing comonomers into the polymer unit can significantly enhance molecular recognition by the column.

This method is quite useful for various purposes because molecular recognition can be expanded and/or enhanced by introducing different comonomers into the polymer unit. PNIPAAm-based temperature-responsive chromatography
requires less time to optimize the mobile phase and requires less skill compared to conventional reversed-phase liquid chromatography (RPLC) because the physical properties of the stationary phase can be easily controlled by only changing the temperature. Moreover, this chromatographic method does not require any toxic organic solvents. This is advantageous not only because it maintains biological activity, but also because of the benefits to consumer and environmental health. PNIPAAm-based temperature-responsive chromatography may be used as a simultaneous bioanalytical method for multiple CYP substrates used during drug development and may replace conventional analysis methods.

4 Oligonucleotide Separation Using a P(NIPAAm-co-BMA-co-DMAPAAm) Hydrogel-modified Column

For over 30 years, oligonucleotide drugs have been developed to treat such diseases as diabetes, cancer, and HIV/AIDS. Recently, considerable progress has been made in overcoming challenges related to their administration, biodistribution, cellular uptake, and undesired side effects, and several drugs have received regulatory approval. During oligonucleotide synthesis, the drug substance is synthesized onto a solid support. A number of different purification steps are required, especially chromatographic purification and analysis that enable the recognition of slight structural differences following synthesis.

TRC using P(NIPAAm-co-BMA-co-DMAPAAm) hydrogel-modified silica beads helped recognize differences in the lengths of single nucleotides, terminal single bases, and the number of sites (Fig. 6a). The elution behaviors of three oligonucleotides with different lengths (dpT10, dpT15, and dpT20) were observed with the P(NIPAAm-co-BMA-co-DMAPAAm) hydrogel modified TRC column (2.1 i.d. × 50.0 mm) at various column temperatures using a phosphate buffer (pH 7.0) as the mobile phase. Combinations of multiple oligonucleotides were well separated by this TRC, as shown in Figs. 6b and 6c. The retention of oligonucleotides was increased by increasing the temperature, opposite to the behavior exhibited by conventional RPLCs. The separation of single nucleobases in oligonucleotides, such as guanine (G), cytosine (C), thymine (T), or adenine (A), was also achieved using the P(NIPAAm-co-BMA-co-DMAPAAm) hydrogel-modified column.

Nucleic acids are highly susceptible to degradation by endogenous nucleases, and phosphorothioate oligonucleotides (commonly referred to as S-oligos) are often used in these systems because of their resistance to cellular nuclease degradation. In conventional RPLC, the separation of S-oligos requires an organic solvent with an ion-pairing reagent, such as tetraethylamine-acetic acid, as the mobile phase and gradient elution. In contrast, Maekawa et al. recently reported that good separation of multiple phosphorothioated oligonucleotides was obtained by TR-HPLC using a P(NIPAAm-co-BMA-co-DMAPAAm) hydrogel-modified column with an aqueous solvent and an isocratic elution without ion-pairing reagents.62 These results indicate that TR-HPLC is well-suited for ensuring the quality and safety of therapeutic oligonucleotides.

5 Temperature-responsive Solid-phase Extraction Column for Protein Purification

The biopharmaceutical market has been rapidly growing, and eight of the ten top-selling drugs in the world are currently biopharmaceuticals. In particular, antibody drugs are in high demand due to their typically small number of side effects. The market for therapeutic monoclonal antibodies (mAbs) has also grown tremendously over the last decade.

However, one issue associated with antibody-drugs is their
high price, and the high price of antibody-drugs increases the burden on patients. Because 66% of the manufacturing costs of antibodies results from the purification process, a highly efficient purification method is needed. Several antibodies may aggregate under low pH conditions during the purification process. This aggregation results in the loss of their bioactivities and a decreased the purification efficiency. Therefore, developing a novel purification method for proteins, such as antibodies, under mild conditions and with sustainable costs is greatly required.

Utilizing TRC in HPLC system, we have reported successful protein separation without the loss of bioactivity with an aqueous mobile phase without using an organic solvent. Furthermore, we developed a temperature-responsive solid-phase extraction (TR-SPE) column to purify a variety of proteins. The surfaces of aminopropyl silica beads (with an average diameter of 40 to 64 μm) were modified with hydrogels of a PNIPAAm copolymer, and then packed into a PTFE column for SPE. Retention of proteins results from the hydrophobicity of the PNIPAAm and BMA moieties, as well as the electrostatic interactions of AAc in the temperature-responsive polymer. SPE is easy to use, can be automated, and is less expensive compared to other extraction methods. It is also possible to scale-up this process for applications in the industrial purification of biopharmaceuticals.

Rituximab is a chimeric monoclonal antibody towards the antigen CD20, and it has been widely evaluated in B cell hematological malignancies. Rituximab may be purified from a hybridoma cell culture medium using the developed P(NIPAAm-co-BMA-co-DMAPAAm) hydrogel-modified column (Fig. 7). The denaturation of rituximab was not observed in the fraction eluted from the TR-SPE column (Fig. 7b). During purification, rituximab did not elute at the washing step, while most of the contaminants were eluted, as shown in Figs. 7b and 7c. The reactivity of the antibody to CD20 positive cells after purification was equivalent that of the sample before purification; thus, the maintenance of antibody activity after purification was confirmed (Fig. 7d). Figure 7e, shows that a good separation of rituximab was obtained from a mixture of MAbs with the same constant region. This indicates that recognizing differences in the variable region of MAb can be achieved by altering the temperature. These results demonstrate that the developed TR-SPE column can be applied in biomedical purification processes, which maintaining the biological activity of the proteins.

6 Temperature-modulated Cell Separation Using P(NIPAAm-co-BMA-co-DMAPAAm) Hydrogel-modified Packing Materials

In recent years, cell transplantation therapy to recover the functions of tissues and organs has attracted attention. Kymriah was recently approved by the Japanese Ministry of Health, Labour, and Welfare in 2019, becoming the first chimeric antigen receptor T-cell (CAR-T) therapy to be approved in Japan. CAR-T therapy is one of the most promising approaches
in anticancer therapy, and it has been hailed as a revolutionary, final method for people with advanced cancer following previous treatment. However, CAR-T therapies involve some of the most expensive drugs in the pharmaceutical market.

By utilizing P(NIPAAm-co-BMA-co-DMAPAAm) hydrogel modified TR-SPE system, in which silica beads with average diameters of 100 to 150 μm were used for optimizing the column towards the size of blood cells, temperature-dependent cell elution behavior was observed in the separation of HL-60 and Jurkat cells (leukemia cells) (Fig. 8). Both cell types were retained in columns containing beads with cationic properties at 37°C, whereas beads without cationic properties did not retain the cells. At 37°C, cells can be retained in the column through both electrostatic and hydrophobic interactions, and the cells are eluted at 4°C because of reduced hydrophobic interactions. The cell retention and elution behaviors were changed by varying the content of the cationic group in the copolymer. The cell retention mainly resulted from electrostatic interactions between the polymer on the beads and the cell surfaces. These results indicate that the retention and elution of cells by the prepared beads packed into the TR-SPE column can be simply modulated by changing the temperature of the system. The effects of modulating the temperature of the column separation process on the cell activity, cell morphology, and viability at each column separation step was observed. The cell morphologies of the HL-60 and Jurkat cells did not change before and after passing through the column. In addition, the passage culture of the eluted cells can be achieved without demonstrating any morphological changes. The cell viability determined via the trypan blue assay remained constant both before and after passing through the column, as well as after obtaining a passage culture of the eluted cells. These results indicate that the column separation does not affect the cell viability. The prepared column can be applied to various types of cell separation by optimizing the properties of the modified polymer on the beads.

Recently, Nagase et al. reported on the use of thermoresponsive cationic block copolymer brushes for temperature-modulated stem cell separation. They prepared a thermoresponsive cationic block copolymer brush made on a glass surface using a two-step activator regenerating by an electron transfer (ARGET)-
that the developed temperature-responsive SPE column can be applied in the purification of biomedical substances with low costs and may act as an effective cell separation tool for biomedical applications. The proposed separation system may also be applicable to various types of protein and cell separations, as well as purification in drug discovery by optimizing the properties of modified polymers. Therefore, the developed temperature-responsive column can be an effective protein and cell separation tool for applications in bioindustries.

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