Chromatographic Behavior of Proteins on Stationary Phase with Aminocarboxy Ligand

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Received November 2, 2010, Accepted December 8, 2010

An aminocarboxy aspartic acid-bonded silica (Asp-Silica) stationary phase was synthesized using L-aspartic acid as ligand and silica gel as matrix. The standard protein mixtures were separated with prepared chromatographic column. The effects of solution pH, salt concentration and metal ion on the retention of proteins were examined, and also compared with traditional iminodiacetic acid-bonded silica (IDA-Silica) column. The results show that Asp-Silica column exhibited an excellent separation performance for proteins. The retention of proteins on Asp-Silica stationary phase was consistent with electrostatic characteristic of cation-exchange. The stationary phase displayed typical metal chelate property after fixing copper ion (II) on Asp-Silica. Under competitive eluting condition, protein mixtures were effectively isolated. Asp ligand showed better ion-exchange and metal chelating properties as compared with IDA ligand.

Key Words: Cation-exchange, Metal chelate, Aminocarboxy ligand, Competitive elution, Protein

Introduction

Aminocarboxy is one of the commonly used stationary phase ligands for high performance liquid chromatography (HPLC).1,2 The characteristics of these ligands are: 1) aminocarboxy stationary phase without metal ion can be used as cation exchanger to separate oppositely charged protein.3-7 2) The stationary phase with metal ion can be used to isolate protein with selective affinity for immobilized metal ion.8,9 Hence, aminocarboxy ligand has been widely used for the separation and purification of proteins.

A series of aminocarboxy ligands were explored in order to improve the selectivity and stability of chromatographic columns. The common ligands are iminodiacetic acid (IDA), nitrolotriacetic acid (NTA), N,N,N′-tris(carboxymethyl)-ethylene-diamine (TED), carboxymethylated aspartic acid (CM-ASP), poly-aspartic acid (PAA), etc.. Among these ligands, PAA is a commonly used ligand.10 However, PAA just can be used as an ion-exchange ligand, but not as a metal chelating ligand. In this paper, the study mainly focused on examining the characteristics of aminocarboxy-silica stationary phase with dual chromatographic functions. Another most widely used ligand is IDA.11,12 Studies show that ion-exchange property of IDA is significant and stable, but metal chelating property of IDA is unstable, especially for strong affinity column. A typical strong affinity metal chelate column is obtained if IDA column is immobilized with Cu²⁺. Adsorbed proteins on the column can be eluted only by adding the competitive agent with a certain concentration.7 However, the fatal defects of this approach are ineffective separation for proteins and serious leakage of Cu²⁺. As a result, the performance and life of the column are all diminished, and the column even can not be applied.

To improve above defects, Asp-Silica stationary phase possessing ion-exchange and metal chelating properties together was synthesized in our laboratory. Based on examining ion exchange property of Asp-Silica stationary phase, a new metal chelate stationary phase-Asp-Cu(II)-Silica was prepared through fixing Cu²⁺ onto the Asp-Silica stationary phase. Ion exchange and metal chelating properties of Asp ligand were compared with those of traditional IDA ligand.

Experimental

Reagents and Chemicals. Silica gel (7 μm, 300 Å) was obtained from Lanzhou Institute of Chemical Physics of the Chinese Academy of Sciences (Lanzhou, China). γ-glycidoxypropyltrimethoxysilane (γ-GLDP) was purchased from Gaixian Chemical Engineering Institute (Liaoning, China). L-Asp was purchased from Shangpu Chemical Engineering Company (Shanghai, China). Imidazole (Imid), glycine (Gly) and the other reagents were of analytical grade and purchased from Xi’an Chemical Reagent Company (Xi’an, China). Cytochrome-C (Cyt-C), lysozyme (Lys) and ribonuclease (RNase) were purchased from Sigma Company (St. Louis. Mo., USA). Bovine serum albumin (BSA) was obtained from Biochemical Reagent Company (Shanghai, China). 2.0 mg/mL protein mixture of BSA, RNase, Cyt-C and Lys were prepared with 20 mM phosphate buffer (PB, pH 6.0).

Apparatus. ÄKTA purifier 10 (Amersham Biosciences, Sweden) was used for chromatographic experiments. Atomic absorption spectrometry TAS-986 (Puxi General Apparatus Co., Ltd., China) was used for determining binding capacities and leakages of Cu⁷⁺. Potentiometer TitraMate 20 (Metttler Toledo, Switzerland) was used for adjusting the pH of the solutions. Slurry packed apparatus 124PP (Chemico, Japan) was used for column packing.

Synthesis of Epoxy-Activated Silica. Epoxy-activated silica was synthesized in acetate-acetic acid buffer (NaAc-HAc, pH
4.0) according to the procedure in reference.12

**Synthesis of Asp-Silica Stationary Phase.** 1.5 g of L-Asp was added to 30 mL of 1 M sodium carbonate (Na₂CO₃). The pH of the solution was adjusted to 8.5 followed by the addition of 2.0 g dried epoxy silica gel. The reaction was carried out at 65 °C for 12 h with stirring. The obtained Asp-Silica was washed successively with water, 10% acetic acid and water again until neutral pH.

**Column Packing.** Synthetic stationary phase was packed into 100 × 4.6 mm I.D. stainless-steel columns under 40 MPa pressure.

**Preparation of Asp-Cu(II)-Silica Stationary Phase.** Adopting dynamic chelate method, the packed Asp-Silica column (100 × 4.6 mm I.D.) was connected to the chromatographic system, and then perfused with 50 mM copper sulfate in 10 mM PB, successively (detection with Na₂S).

**Determination of Binding Capacity for Immobilized Cu²⁺.** Copper ion (II) immobilized to metal chelate column was eluted by gradient elution with 50 mL volumetric flasks and then diluted to the scale with water. Binding capacity of Cu²⁺ in the diluted eluate was determined by atomic absorption spectrophotometry (AAS) using EDTA-2Na as a blank solution.

**Determination of Leakages for Immobilized Cu²⁺.** The packed Asp-Cu(II)-Silica column (100 × 4.6 mm I.D.) was connected to the chromatographic system. Copper ion (II) immobilized to metal chelate column was eluted by gradient elution with the selected competitive systems. The eluate was collected into 50 mL volumetric flasks and then diluted to the scale with water. Leakage of Cu²⁺ in the diluted eluate was determined by AAS using competitive eluant as a blank solution.

**Chromatographic Experiments.** Effects of the pH of solution on retention, the concentration of salt and immobilized metal ion on retention behavior of proteins, separation of protein mixtures on the columns were performed according to the chromatographic conditions given in Figures.

**Results and Discussion**

**Ion Exchange Properties of Asp-Silica Stationary Phase.**

**Effect of pH Value on the Retention of Proteins:** Effect of pH on the retention of proteins is a rather complicated issue, involving not only the electronegativity of stationary phase ligand but also the electronegativity of protein.13 To examine effect of pH on the retention of proteins, retention behaviors of three basic proteins on Asp-Silica column were investigated using gradient elution at a wide range of pH 3.0 - 8.0. The results are presented in Fig. 1. It can be seen from Fig.1 that: (1) At pH 3.0 - 4.0, retention times of RNase, Cyt-C and Lys increased with the increase of pH values; However, retention times of three proteins decreased with the increase of pH values at the range of 5.0 - 8.0. This is since L-Asp is a ternary weak acid (pI = 2.97) and exists dissociative equilibrium at different pH values. The primary dissociation style of L-Asp without charge (Asp±) is transformed into the secondary dissociation style (Asp±) (Asp-) (Asp²⁻) (Asp³⁻) in solution as follows

\[
\begin{align*}
\text{H}_2\text{N} & \quad \text{OH} \\
& \quad \text{COOH} \\
\text{pK}_1 & = 2.09 \\
\text{pK}_2 & = 3.86
\end{align*}
\]

**Figure 1.** Effect of mobile phase pH on retention of proteins Column: Asp-Silica column (4.6 mm × 100 mm I.D.); Mobile phase: A: 20 mM PB (pH 8.0, 7.0, 6.0, 5.0, 4.0, 3.0); B: 20 mM PB + 0.5 M NaCl (pH 8.0, 7.0, 6.0, 5.0, 4.0, 3.0); Gradient elution: B from 0 to 100% in 20 min. Flow rate: 1 mL/min. Detector: UV (λ = 280 nm). Size of sample: 5 μL.
If \( k' \) represents retention factor of protein on ion-exchange column, \( \lg f \) is a constant related to properties and volumes of stationary phase and mobile phase, electrostatic interaction between stationary phase ligand and protein as well as salt, then the relationship between retention of protein on ion-exchange column and salt concentration should meet stoichiometric displacement theory (SDT-R),

\[
\lg k' = \lg f + z \lg (1/[D])
\]

To demonstrate the correctness of formula (1), retention values of three proteins on Asp-Silica column were determined by isocratic elution using PB with different salt concentrations. Fig. 2 is a plot of \( \lg k' \) (logarithm of retention factor) vs. \( \lg 1/[D] \).

\[
L - P + zD = L - D^*_1 + P
\]

\( D_1 \) is the degree of dissociation of the ligand on the ion-exchange column.

Figure 2. Effect of salt concentration on retention of proteins Mobile phase: 20 mM PB + X M NaCl (pH 6.0, 0 - 0.5 M). Isocratic elution. Other conditions are the same as in Fig. 1.

The above results show that the retention of protein on Asp-Silica column follows electrostatic interaction characteristic of cation-exchange.

Separation Comparison of Proteins on Asp-Silica and IDA-Silica Columns: To examine the separation performance of Asp-Silica column, standard protein mixture was separated with synthetic column using gradient elution, and the separation property of Asp-Silica column was compared with that of traditional IDA-Silica column. The results are presented in Fig. 3. As seen in Fig. 3, four proteins were effectively isolated with prepared Asp-Silica column. Furthermore, adsorbed proteins on Asp-Silica column were eluted in the order of an increase in pI respectively, that is, acidic protein BSA (pI 4.9), basic proteins RNase (pI 8.7), Cyt-C (pI 10.1) and Lys (pI 11.0). This result once again demonstrated that synthetic Asp-Silica stationary phase had typical cation-exchange characteristics. In addition, the column efficiency and resolution of Asp-Silica were all superior to those of commonly used IDA-Silica column under the same chromatographic conditions.

Metal Chelating Properties of Asp-Cu(II)-Silica Stationary phase.

Effect of Immobilized Cu(II) on the Retention of Proteins: Adsorption of protein is occurred due to the coordination role of protein with immobilized Cu(II). When using competitive elution, chelate protein can be replaced by competitive agents such as Imid and Gly, and then adsorbed protein on metal chelating column is desorbed. Different proteins have the different affinities for Cu(II). In most cases, this difference can be used for the effective separation and purification of protein.

Fig. 4 displays chromatograms of proteins on Asp-Cu(II)-Silica and IDA-Cu(II)-Silica columns in PB-NaCl eluting system. As seen in Fig. 4, after immobilizing Cu(II) on the naked
Asp-Silica column, the eluting capacities of PB-NaCl for four proteins all significantly decreased. RNase and Cyt-C were not separated, and the eluting order of these two proteins also changed. These are the results of coordination role between immobilizing Cu²⁺ and protein. In addition, four proteins on IDA-Cu(II)-Silica column were hardly eluted. This shows that IDA-Cu(II)-Silica column exhibited stronger coordination role with proteins as compared with Asp-Cu(II)-Silica column, proteins adsorbed on it could not easily be eluted.

Competitive Elution Comparison of Proteins on Asp-Cu(II)-Silica and IDA-Cu(II)-Silica Columns: In addition to stationary phase, the choice of separation conditions is also important for metal chelate chromatography. Among these conditions, the type and concentration of competitive agents are particularly important because these conditions directly affect separation performance of chelating column and leakage of Cu²⁺ from chelating column. Fig. 5 displays chromatograms of proteins on Asp-Cu(II)-Silica and IDA-Cu(II)-Silica columns in competitive eluting system. As seen in Fig. 5, separation performance of Asp-Cu(II)-Silica column for proteins was better than that of IDA-Cu(II)-Silica column, four proteins on IDA-Cu(II)-Silica column were hardly eluted. On the other hand, although the leakage of Cu²⁺ on IDA-Cu(II)-Silica column (0.023 ± 0.02 μmol/mL) was slightly lower than that on Asp-Cu(II)-Silica column (0.028 ± 0.02 μmol/mL) at the same chromatographic conditions, the eluting strength of proteins on IDA-Cu(II)-Silica column was weak. This is because IDA is a stronger complexing agent, the stability constant of IDA with Cu²⁺ (lgK_{IDA-Cu(II)} = 10.62) is greater than that of L-Asp with Cu²⁺ (lgK_{L-Asp-Cu(II)} = 8.57). Therefore, the bonding capacity of Cu²⁺ on IDA-Cu(II) column (20.11 μmol/ggel) was also higher than that of Asp-Cu(II)-Silica column (18.93 μmol/ggel). Adsorbed proteins on IDA-Cu(II)-Silica column was difficult to be eluted in comparison with Asp-Cu(II)-Silica column. Consequently, the concentration of Imid should be increased in order to obtain efficient elution. However, the poor separation and high leakage of Cu²⁺ would be caused due to the increase in the concentration of Imid.

It follows that separating and eluting performances of metal chelate column should also be considered for the selection of ligand in addition to the stability of metal chelate column. The chelate with higher stability constant and bonding capacity with Cu²⁺ is not always the best choice of metal chelate ligand. The coordination intensity between the chosen ligand and immobilized Cu²⁺ should be moderated in order to ensure a less leakage of Cu²⁺, effective elution and separation of proteins on metal chelate Cu(II) column.

Conclusion

In summary, chosen L-Asp ligand had dual functions of cation-exchange and metal chelate properties. At same time, separation and elution performances of Asp columns were all superior
to those of traditional IDA columns, especially in the aspect of metal chelate property. These studies may provide certain reference values for the selecting and developing the aminocarboxy ligand with multifunction, improving the stability and elution performance of chromatographic column.

Acknowledgments. This work was supported by grants from Natural Science Foundation of Shaanxi Province (No. 2007 B22), Natural Science Fund of Shaanxi Provincial Education Department (No. 09JK758), Shannxi Provincial Key Discipline Program and Scientific Research Foundation for the Master Program of Northwest University (No. 09YSY24). We thank to Ms Tan Hui of Xi’AN University of Architecture and Technology for the revision of the manuscript.

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