



Purification of Fab using High Binding Capacity TOYOPEARL® AF-rProtein L-650F

A High Capacity Protein L Resin for the Purification of Monoclonal Antibody Fragments

Introduction

Protein L based affinity chromatography is used for the capture of antibodies and antibody fragments that do not bind to protein A. Unlike protein A and G, which bind to the Fc region of immunoglobulins (IgGs), protein L binds through interactions with the variable region of an antibody's kappa light chain. Figure 1 shows several possible binding sites of protein L to fragments (Fabs), single-chain variable fragments (scFvs) and domain antibodies (dAbs).



TOYOPEARL AF-rProtein L-650F is an affinity chromatography resin that combines a rigid polymer matrix with a recombinant ligand, which is derived from the B4 domain of native protein L from Peptostreptococcus magnus and is expressed in E.coli (Figure 2). Code optimization of the domain results in high binding capacity and an improved alkaline stability of the ligand compared to the native molecule.

STRUCTURE OF TOYOPEARL AF-rPROTEIN L-650F



Figure 2

The selected recombinant protein L ligand also has an affinity for a broad range antibody subclasses as demonstrated in Table 1. This application note demonstrates the ability of TOYOPEARL AF-rProtein L-650F resin to capture and purify Fabs with a high yield and high purity. TOYOPEARL AF-rPROTEIN L-650F LIGAND WITH A BROAD AFFINITY RANGE FOR MAB SUBCLASSES

Species	Class	Affinity
General	Kappa light chain	++
	Lambda light chain	-
	Heavy chain	-
	Fab	++
	ScFv	++
	Dab	++
Human	lgG(1-4)	+
	lgA	+
	lgD	+
	IgE	+
	lgM	+
Mouse	IgG ₁	+
	lgG ₂ a	+
	lgG ₂ b	+
	IgA	+
	lgM	+
Rat	IgG ₁	+
	lgG2a,b,c	+
	IgA	+
Hen	IgY	+
Table 1		

MATERIALS AND METHODS

DIGESTION

Humanized IgG1, 3 mg/mL, was digested using papain enzymatic protocol as described in the Pierce[™] Fab Preparation Kit (catalog number: 44985).

CHROMATOGRAPHY

Purification of Fab: 100 μL of the papain digested IgG1 was loaded onto a TOYOPEARL AF-rProtein L-650F column, 5 mm ID \times 2.5 cm (0.5 mL total volume). See chromatogram for operating conditions.

Size analysis of fractions: the collected peaks, 10 μL each, were injected onto a TSKgel® UP-SW3000, 4.6 mm ID \times 15 cm size exclusion chromatography (SEC) column. See chromatogram for operating conditions.

RESULTS AND DISCUSSIONS

PURIFICATION OF FAB

Figure 3, panel A, shows the purification result of Fab from the lgG1 papain digested sample using the TOYOPEARL AF-rProtein L-650F column. Only two peaks are observed in the chromatographic profile - the flow through peak and the eluted peak. The eluted peak is eluted with 0.1 mol/L citrate buffer, pH 2.05 at ~90% gradient. The data suggests that Fab is bound strongly onto the TOYOPEARL AF-rProtein L-650F resin. Both peaks were collected for further analysis using a TSKgel UP-SW3000, 4.6 mm ID × 15 cm SEC column. The eluted peak is expected to contain only the Fab fragment.

Figure 3, panel B, shows results of the size exclusion analysis of the digested sample, the flow through and eluted peaks. The SEC profile showed that the digested sample contained two different molecular weight products based on the estimated retention times. As mentioned above, the papain digestion dissociates Fc and Fab regions of IgG and the protein L ligand only has an affinity for the Fab fragment. Therefore, there is no binding of the Fc fragment to the protein L resin and only Fab can be bound and eluted from the protein L column. Data from Figure 3, panel B, shows that the first/flow through peak from the papain digested sample has a retention time of 17 minutes and the second/eluted peak has a retention time of ~19 minutes.

► PROCESS RESINS

The 17 minute retention time peak is suggested to be the Fc fragment because it has the same retention time with the flow through peak. And the 19 minute retention time peak is suggested to be the Fab fragment due to it having the same retention time as the eluted peak from the TOYOPEARL AF-rProtein L-650F column.

DYNAMIC BINDING CAPACITY

A purified Fab sample was loaded onto the TOYOPEARL AF-rProtein L-650F resin, 0.83 mL column volume with 10% breakthrough at 3.4 minutes and 4.0 minutes residence times to determine the dynamic binding capacity (DBC) of a Fab fragment on the resin. Table 1 shows the comparison data between TOYOPEARL AF-rProtein L-650F resin to its competitor 85 µm agarose-based particle size protein L resin at various residence times. Due to the excellent mass transfer characteristics of TOYOPEARL AF-rProtein L-650F resin, dynamic binding capacities at 1 to 3 minutes residence time excel capacities obtained with the agarose-based resin (nearly double) as shown in Table 2. As the molecular weight of fragments is much smaller compared to full length lgGs, a dynamic binding capacity of about 50 mg/mL for a Fab with a typical molecular weight of 55 kDa equals a DBC of >130 mg/L for a ~150 kDa lgG when considering molar binding capacities.

PURIFICATION OF FAB USING TOYOPEARL AF-rPROTEIN L-650F



A: Purification of Fab by Protein L resin

DYNAMIC BINDING CAPACITY OF PROTEIN L RESINS FOR FA	AB	3
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			Fab DBC (g/L-resin) Residence time			
		lgG SBC (g/L-resin)				
			3.4 min	4.0 min		
rProtein L-65	i0F	72	33	50		
Competitor p	roduct	42	19	26		
Column: Detection: Sample:	TOYOPEARL AF-rProtein L-650F/competitor resin, 4.6 mm ID × 50 mm (0.83 mL) UV @ 280 nm 2 g/L human Fab in 0.1 mol/L Na-phosphate, pH 6.5 DBC measured at 10% breakthrough					
Table	2					

CONCLUSIONS

The dynamic binding capacity of TOYOPEARL AF-rProtein L-650F resin exceeds the protein L agarose-based, 85 µm chromatography media currently available. Due to the high binding capacity, high yield and ease of use, TOYOPEARL AF-rProtein L-650F resin can considerably improve process economics of protein L capture steps. With high affinity antibody fragments, single chain variable fragments and domain antibodies, it is suited for the purification of new antibody formats that cannot be purified with protein A media.

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