Modeling of immunoglobulin uptake by N,N,N′,N′-ethylenediaminetetramethylene phosphonic acid-modified zirconia particles under static and dynamic conditions

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1. Introduction

The purification of biomolecules is an important problem in downstream bioprocessing [1–9]. Economics, efficiency and practicality are some of the constraints that dictate the search for novel chromatographic supports and methodologies that offer novel selectivity or overcome the shortcomings of existing supports. Zirconia based supports, particles with thermal and mechanical stability [6], have the potential to offer both. Our previous studies have established the usefulness of $N,N,N',N'$-ethylenediaminetetramethylene phosphonic acid-modified zirconia in the separation of immunoglobulins from complex mixtures [7]. Research-based prediction of mass transport, biological activity behavior, kinetic and thermodynamic parameters that impact protein retention and separation are essential for the integration of chromatographic-based unit-operations into the purification scheme [10–16]. A quantitative or qualitative knowledge of the parameters involved in the transport of biomolecules in a chromatographic system is needed before improvements may be designed. The determination of the rate of uptake or binding of the molecules is an essential part of the information required for the modeling of the system.

Various theories have been developed to describe the binding mechanism. The most rigorous being the general mass transfer mechanism [12, 17, 18]. Suitable mathematical models have been postulated to describe and analyze the transport of proteins and solutes in porous beads and, the protein uptake from a finite medium. The kinetic rate constant model [15, 16] and the film and
pore diffusion model and its variations [16, 19-21] were employed to approximate the protein uptake profiles in a finite medium. Dynamic breakthrough profiles were approximated by model equations as outlined elsewhere [21, 22]. The adsorption phenomena were attributed due to the combined effects of solute transport and adsorp- tion. The relevant transport equations were either analyti- cally or numerically solved after suitable approximations and assumptions were made about the rate limiting fac- tors driving the adsorption phenomena [23-25].

Application of transport-model equations require an estimate of the rate coefficients and equilibrium constants or require the determination of dimensionless parameters like the Sherwood or Peclet number. Under most circum- stances the information required for calculating such pa- rameters are not available. It is however, possible to first calculate these parameters and then proceed on with the modeling, using pulse injection techniques. Pulse tech- niques in conjunction with Laplacian transformation and statistical analysis can be used to solve the transport equa- tions [25-27] and further obtain the transport parameters.

Our goal was optimize the chromatographic perfor- mance of r_PEZ by gaining a better understanding of the solute transport under dynamic conditions and in a finite medium. In this paper, protein uptake studies by r_PEZ in a finite medium and under dynamic conditions were undertaken to better understand the interaction of human immunoglobulins (Igs) with the r_PEZ. Experimentally obtained profiles were compared to the profile predicted by the kinetic rate constant model. The dynamic break- through profiles obtained from frontal analysis were ap- proximated and compared to the profile predicted by the kinetic rate constant model [15, 16]; with the anticipation that these engineering criteria would enable us to better understand the performance of r_PEZ in bioseparations.

2. Material and methods

2.1. Reagents

All chemicals were of analytical-grade or better. So- dium chloride was purchased from Fisher Scientific (Ha- nover Park, IL, USA). N,N,N',N'-Ethylene diaminetetra- methylene phosphonic acid (EDTPA) was purchased from TCI America (Portland, OR, USA). Bovine serum albumin (BSA), pure human immunoglobulin G (HIgG), all horse- radish peroxidase conjugated anti-immunoglobulins used for ELISA were obtained from Sigma Chemical Company (St. Louis, MO, USA). All proteins were used without fur- ther purification. Human immunoglobulin A (HIgA) and human immunoglobulin M (HIgM) were purchased from Jackson Immunoresearch (West Grove, PA, USA).

Immulon II microtiter plates were purchased from Fisher Scientific (Hanover Park, IL, USA). Affinity puri- fied goat anti-mouse (whole molecule) immunoglobu- lins and goat anti-mouse immunoglobulins conjugated to horseradish peroxidase (HRP) were purchased from Sigma Chemical Company (St. Louis, MO, USA). o-Phenyl- enediame-2HCl (OPD) tablets were purchased from Abbott Laboratories (Chicago, IL, USA). Pre-cast Nu- Page 4-12% Bis-Tris gels were purchased from Invit-rogen (Carlsbad, CA, USA). A Genesys™ 5 model from Spectronic Instruments UV-vis spectrophotometer (Rochester, NY, USA) was used to record the adsorption measurements. A bench top microcentrifuge (Eppendorf Cen- trifuge 5415C) was used to sediment the r_PEZ particles for batch experiments.

2.1.1. Support matrix preparation

Colloidal zirconia was spray dried to yield zirconia particles, which were further classified, modified with EDTPA and characterized as reported elsewhere [28]. EDTPA-modified zirconia particles will be referred to as r_PEZ in this manuscript. r_PEZ particles were packed into a 0.46 cm i.d. × 5.0 cm length analytical column, and supplied by ZirChrom (Anoka, MN, USA).

2.2. Ligand binding isotherms

Batch experiments conducted to determine the equi- librium binding capacity of r_PEZ for HIgG was done as described elsewhere [28]. Briefly, microfuge tubes filled with the same and known volumes of equilibrated and wet r_PEZ beads were loaded with constant volumes of HIgG solutions with different stock concentrations. Samples were allowed to equilibrate for 24 h and the resultant supernatant concentration measured spectrophotometri- cally at 280 nm. Amount of HIgG bound was determined via mass balance.

Independent ligand binding isotherms were also deter- mined for the binding of HIgA and HIgM to r_PEZ.

2.3. Batch kinetic studies

The rate of adsorption of proteins to r_PEZ beads were determined at different protein concentration in small batch experiments. Four hundred microliters of 50% (v/v) slurry of r_PEZ beads were transferred into 3 ml plastic tubes to yield approximately 200 μl of beads. The beads were allowed to settle for at least 5 min and the liquid overlay was pipetted off after centrifuging for 5 min at 8000 rpm. Stock solutions of HIgG, HIgA and HIgM were prepared with appropriate dilutions. Prior to their use, HIgA and HIgM stock solutions obtained from suppliers were diafiltered and buffer replacement carried out with the LB, using Millipore’s Centricon YM-10 (Bedford, MA, USA). The feed concentrations investigated for HIgG were 1, 5 and 10 mg/ml. Those for HIgA and HIgM were 0.46 and 1.84 mg/ml and 0.184 and 0.92 mg/ml, respectively. Two milliliters of stock solution was introduced into the prepared r_PEZ beads at 4 °C and placed on an end-to- end rotator. Ten microliters of aliquots were drawn at 0, 0.5, 1, 5, 10, 20, 30, 45, 60, 120, 240 and 1440 min. Stock concentration was drawn for 0 min aliquot. Experiments were performed in duplicate. Protein concentration was
measured in the aliquots and the amount bound was found by mass balance. HlgG in the aliquots was measured by detecting the absorbance at 280 nm. For HlgA and HlgM their respective ELISAs were performed. Data was presented as normalized concentration, \( C/C_0 \) (aliquot/supernatant concentration against feed concentration) versus time.

2.4. Chromatography

The chromatographic system consisted of a Chrom Tech (Apple valley, MN, USA) Iso-2000 isocratic pump in conjunction with an online Model 783 Spectroflow spectrophotometer (Ramsey, NJ, USA), which was used mainly as an indicator, and an SRI (Torrance, CA, USA) PeakSimple Model 203, single channel serial port online data acquiring system. The absorbance of the fractions was then measured using the spectrophotometer.

All buffer solutions were filtered through ChromTech’s Metal-Free solvent (type A-427) 10 μm ultra high molecular weight polyethylene (UHMWPE) membrane filter at the time of use.

All column experiments were performed with a 0.46 cm × 5.0 cm (diameter × length) analytical column packed with approximately 30–100 μm diameter zirconia beads.

2.5. Dynamic studies

Zirconia packed column’s performance was evaluated by determining the breakthrough curves of HlgG at various flow rates and feed concentration. In all cases pure HlgG dissolved in Loading Buffer (4 mM EDTPA, 20 mM MES, 50 mM NaCl, pH 5.5) to obtain feed concentrations of 0.5, 2.0, 5.0 and 10.0 mg/ml. Protein solution to be used as feed was kept in a chilled reservoir and introduced to the system continuously via the multi channel valve. Linear velocities of 3.01, 6.02 and 12.04 cm/min were investigated. Aliquots of the outlet stream were collected and their protein content measured at 280 nm. In all cases the protein solution was allowed to saturate the column till the flow through protein concentration reached 75–80% of the feed concentration. At the end of the loading process the proteins were eluted from the column using Elution Buffer (4 mM EDTPA, 20 mM MES, 1 M NaCl) and protein content measured. Zero time was marked as the time when the valve was switched from the Loading Buffer to the feed solution. Data was plotted as normalized concentration, \( C/C_w \) of outlet protein concentration against the maximum protein concentration obtained in an aliquot; by normalized time, \( T/T_{max} \).

2.6. Determination of HlgG, HlgA and HlgM by ELISA

The concentrations of the IgG were determined by an ELISA procedure as outlined elsewhere [28]. Individual ELISA was carried out to determine the concentration of each species of immunoglobulins.

2.7. Modeling and simulation

Kinetic rate constant model equations [15, 16, 19] were solved using a program written in MATLAB [29]. The pore diffusion rate-limiting model was used to approximate the dynamic profiles. The parameters were optimized by least squares minimization using the constrained optimization routine LSQCURVEFIT.

3. Results

3.1. Ligand binding isotherms

The maximum binding capacity (\( Q_{max} \)) and the dissociation (\( K_d \)) constant were determined from the batch isotherm data as described earlier [28]. The \( Q_{max} \) and \( K_d \) values for HlgG were found to be 55 mg/ml and 0.7 mg/ml for r_PeZ beads. Following a similar approach, the \( Q_{max} \) and \( K_d \) values for HlgA were determined to be 18.98 mg/ml and 0.01 mg/ml, and that for HlgM was found to be 0.845 mg/ml and 2.486 mg/ml for r_PeZ beads. The inability to maintain both HlgA and HlgM at concentrations greater than 3 mg/ml, limited our construction of reliable isotherms for these molecules.

3.2. Kinetic uptake of Immunoglobulins under static conditions

Small-scale batch experiments were conducted to determine the rate of uptake of HlgG, HlgA and HlgM by r_PeZ beads from a feed solution containing IgG at various feed concentrations (\( C_0 \)). Figure 1a–c shows the rate of disappearance of HlgG, HlgA and HlgM from the solution, respectively. Analysis of HlgG was done by measuring its respective absorbance at 280 nm. The HlgA and HlgM concentration at different time points were estimated by their specific ELISA assays as reported elsewhere [26]. As all experiments were carried out in a closed system, it was assumed that the all Ig not measured in the solution had bound to the support.

Maximum HlgG retention by r_PeZ was observed at 24 h with 60–95% disappearance of protein from solution from an initial HlgG concentration of 10 mg/ml and 1 mg/ml, respectively (Figure 1a). Values did not change appreciably after the 240 min time-point. The largest drop in the percentage disappearance of HlgG, i.e. greatest adsorption rate, occurs within the first 5 min of the batch experiment. Fifty percent of the adsorption occurs roughly after 6 min from the start of the experiment for a feed concentration of 1 mg/ml. By 25 min approximately 80% of total binding has occurred. According to experimental data, 90% of the IgG has been adsorbed by the end of 50 min. A 50% adsorption (\( C/C_0 = 0.5 \)) was attained at 76 and 870 min for HlgG concentrations of 5 and 10 mg/ml, respectively.

According to experimental data, 90% of the HlgA is adsorbed by the end approximately 1400 min (data point not shown). A 50% adsorption was not attained for the
According to experimental data, 90% of the solute has been adsorbed by the end of 180 min. A 50% adsorption was attained at 200 min for HlgM concentration of 0.92 mg/ml.

The experimental data for the protein adsorption obtained under static conditions was approximated using the “kinetic rate constant model”, discussed in detail elsewhere [16, 17]. The only unknown parameter was the forward rate contact (k$_{1}$), where as the isotherm parameters K$_{d}$ and Q$_{\text{max}}$ determined from static binding experiments were used [28] and the reverse rate constant (k$_{2}$) was equated to K$_{d}$k$_{1}$. Simulations were performed with a variety of values of the unknown parameter k$_{1}$ and the value that gave the best fit of the experimental data was reported. The agreement between the experimental data and the simulation is shown in Figure 1a–c. Open circles, stars and open rectangles depict experimental data and solid lines the model prediction obtained after least squares minimization. The various values of the parameters as determined by the optimized model are as indicated in Table 1. For the uptake of HlgG by rPEZ, a k$_{1}$ value of 0.0242, 0.0025, and 0.0028 ml/mg min was obtained at a C$_{0}$(HlgG) values of 1, 5 and 10 mg/ml, respectively.
However for HIgA and HIgM, it was found after multiple attempts, that the model was unable to predict the experimentally derived profile. Hence, the procedure of unconstrained (referred further to as ‘free’) and constrained (referred further to as ‘restricted’) optimization was utilized to obtain the parameters for the best fit of the data. Free optimization was carried out on batch kinetic experiments performed for HIgG and the values obtained for $Q_{max}$ and $K_d$ were compared with the ones found experimentally from isotherm data. It was found that they did not differ significantly; this procedure was used to determine the $Q_{max}$ and $K_d$ values for HIgA and HIgM from their experimental batch kinetic data. Thereafter restricted optimization was utilized to determine the values of the respective $k_1$ and $k_2$ values.

Figure 1b and c show the best fit profiles obtained for HIgA and HIgM batch kinetic data for two different feed concentrations. The open circles indicate experimental data and the solid lines the model prediction. Constrained optimization determined the $Q_{max}$ and $K_d$ values as 8.7 and 0.29 mg/ml for HIgA and 3.8 and 0.055 mg/ml for HIgM to r_PEZ, respectively and the values are summarized in Table 1.

Table 1. Kinetic rate constant model was used to determine the lumped forward ($k_1$) and backward ($k_2$) reaction rate constant. Individual experiments were done in duplicate

<table>
<thead>
<tr>
<th>Beads</th>
<th>HIg</th>
<th>$C_0$ (mg/ml)</th>
<th>$Q_{max}$ (mg/ml)</th>
<th>$K_d$ (mg/ml)</th>
<th>$k_1$ (ml/mg min)</th>
<th>$k_2$ (min$^{-1}$)</th>
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</thead>
<tbody>
<tr>
<td>r_PEZ</td>
<td>A</td>
<td>0.46</td>
<td>8.7</td>
<td>0.29</td>
<td>0.8168</td>
<td>0.2369</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1.84</td>
<td>8.7</td>
<td>0.29</td>
<td>0.0588</td>
<td>0.0171</td>
</tr>
<tr>
<td>r_PEZ</td>
<td>G\textsuperscript{a}</td>
<td>1</td>
<td>55</td>
<td>0.7</td>
<td>0.0242</td>
<td>0.0169</td>
</tr>
<tr>
<td></td>
<td></td>
<td>5</td>
<td>55</td>
<td>0.7</td>
<td>0.0025</td>
<td>0.0018</td>
</tr>
<tr>
<td></td>
<td></td>
<td>10</td>
<td>55</td>
<td>0.7</td>
<td>0.0028</td>
<td>0.002</td>
</tr>
<tr>
<td>r_PEZ</td>
<td>M\textsuperscript{b}</td>
<td>0.184</td>
<td>3.8</td>
<td>0.055</td>
<td>0.5437</td>
<td>0.0299</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.92</td>
<td>3.8</td>
<td>0.055</td>
<td>0.0776</td>
<td>0.0043</td>
</tr>
<tr>
<td>LigoSep A</td>
<td>G\textsuperscript{a}</td>
<td>13.2</td>
<td>66.23</td>
<td>1.85</td>
<td>0.0055</td>
<td>0.0102</td>
</tr>
</tbody>
</table>

\textsuperscript{a} Concentration determined by measuring respective sample absorbance at 280 nm.
\textsuperscript{b} Concentration determined by respective ELISAs.

3.3. Frontal analysis

The dynamic binding of HIgG to r_PEZ was monitored experimentally by breakthrough analysis, at different feed concentrations and linear velocities. Figure 2a and b depict representative breakthrough profiles obtained for HIgG at a feed concentration of 2 and 5 mg/ml, respectively. Separate breakthrough profiles were generated at three different linear velocities of 3.01, 6.02 and 12.04 cm/min, respectively. For a HIgG feed concentration of 2.0 mg/ml, a 10% breakthrough was observed 17, 0.5 and 0.07 min at linear velocities of 3.01, 6.02 and 12.04 cm/min, respectively. A 80% breakthrough in column capacity was obtained at 18, 3.4 and 1.4 min, respectively at linear velocities of 3.01, 6.02 and 12.04 cm/min, respectively. We have obtained similar breakthrough profiles at other HIgG feed concentrations (data not shown).

The experimentally obtained breakthrough profiles were approximated by the various models available in the literature [21]. The kinetic rate constant model [15]
was unable to approximate the breakthrough profiles obtained in our study and was hence not pursued further. Other relevant model equations were used and the mathematical expression governing the pore diffusion model [21] gave a satisfactory fit to the experimental breakthrough profiles. The model prediction and the experimentally obtained breakthrough profiles are as Figure 3a and b. The best fit of pore diffusion model equation to the data in Figure 3a and b gave a rounded off value of $N_p$ equal to 2.

4. Discussion

The long term goal of our research effort is to better understand the rate and mechanism of solute binding and transport in r_PEZ. The objective of this study, which is the next step in achieving our long term goal, is to further understand the kinetic parameters that govern the interaction under static and dynamic conditions. We seek to put forth model equations and identify mass transfer parameters relevant for a preparative scale chromatographic separation with r_PEZ. Our previous studies have shown that the binding of Igs to r_PEZ can be modeled with a pseudo-Langmuir isotherm [28]. Additionally it has been shown that the binding is not adversely impacted by temperature. The kinetic rate constant model; which can be modified suitably to include different adsorption rate equations without making major differences to the final form, was employed to approximate the experimentally obtained protein uptake profiles. It was found that the $Q_{\text{max}}$ and $K_d$ values obtained by ‘free optimization’ process, as described earlier, was in good agreement with those determined experimentally for HIgG data. Hence, we have used this technique to approximate the protein uptake profiles for HIgA and HIgM.

The values of the forward rate constant ($k_1$) were found to decrease with increasing feed concentration as can be seen from Table 1. There is a corresponding decrease in the reverse rate constant ($k_2$) also, which is implied in the definition. For individual Igs the $k_1$ value for comparable concentrations (1 mg/ml of Ig) is largest for HIgM. HIgM is a tertiary molecule (pentamer) that is more bulky than the HIgG molecule, with multiple binding moieties. It adheres to the binding sites more strongly as a result $k_1$ values tend to be higher. It is predicted that the $k_1$ value for HIgA for a feed concentration of 1 mg/ml would be higher than that for HIgG using the same arguments. This maybe deduced from the trend in the $k_1$ values as shown in Table 1.

Higher values of $k_{1r}$ when compared to $k_{2r}$ for Ig adsorption to r_PEZ indicate that the mechanism of the adsorption of Igs are favored over desorption. This phenomenon is apparent by the presence of tailing sections in elution profiles [26]. Under dynamic loading conditions, the rate of adsorption is observed to be higher than that during desorption, as evident in Figure 2a and b. As both $k_1$ and $k_2$ are lumped coefficients it can be only inferred from the trend in their values for r_PEZ that the mechanisms responsible for mass transfer, decrease with increasing Ig concentration. This may be due to the spatial exclusion exerted by the adsorbed biomolecule and its impact on pore diffusive fluxes.

Langmuir isotherms in conjunction with the kinetic rate constant model have been reported to be able to successfully model both batch kinetic and frontal experiments [15]. However, the above-mentioned model did not provide a satisfactory approximation to the dynamic breakthrough profiles obtained in this study. The possible reason maybe that the mobility of HIgG through the pores of r_PEZ is the rate limiting process and aforementioned model does not consider it explicitly [15]. The adsorption of HIgG maybe favorable only at the outer pe-
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ripheral surface of the r_PEZ particles, making liquid film mass transfer the dominant mechanism in the initial phase of adsorption. However in the later phase, adsorption takes place in the interior of the beads owing to the unavailability of free sites at the surface, and this process maybe slower than the initial surface adsorption rate. In a parallel study, we have used FITC-labeled HlgG and its subsequent visualization by confocal laser scanning microscopy (CLSM) to visualize the distribution of binding sites through out the cross-section [30]. Our results show a uniform FITC signal throughput the cross section at HlgG loadings of 5.0 m h IgG/ml or higher (data not included).

As a next step, we have used the “pore diffusion model” to approximate and model the dynamic breakthrough profiles obtained in this study. The profiles obtained at lower feed concentration were satisfactorily approximated by the pore diffusion model equation and the parameter, N_p, that gave a satisfactory fit was found to have a rounded off value of 2. Dynamic breakthrough profiles obtained at higher feed concentrations were not amenable to approximation by the pore diffusion equation. Thus, it is conceivable that some other mechanisms in addition to pore diffusion are rate limiting. A possible explanation to this discrepancy maybe attributed to the relatively slow rate of adsorption to the matrix, as evidenced in the protein uptake profiles obtained in a finite medium.

5. Conclusion

In conclusion, the kinetics of adsorption of Igs onto r_PEZ in a finite medium can be described by the kinetic rate constant model. In the case of r_PEZ, our results suggest that the rate of adsorption of Ig to the matrix is more favorable than the rate of desorption. The mathematical equations that describe the pore diffusion model were used to model the dynamic breakthrough profiles. In the light of the analysis presented here, it appears that the mechanism of mass transfer in r_PEZ beads is limited by pore diffusion. In conjunction with the results presented in our earlier work (Subramanian and Sarkar [28, 31]), a set of engineering parameters are now available that can be used to scale up chromatographic separations based on r_PEZ. As most of the dynamic profiles obtained in this study were not satisfactorily fit using the pore and diffusion model, we will use the pulse injection techniques in conjunction with HETP equations to determine the various transport parameters relevant for scale-up, which will be a subject of a future publication.

References