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DOI 10.1016/j.chroma.2022.463451

Publication date 2022 **Document Version** Final published version

Published in Journal of Chromatography A

Citation (APA)

Silva, T. C., Eppink, M., & Ottens, M. (2022). Small, smaller, smallest: Miniaturization of chromatographic process development. *Journal of Chromatography A*, *1681*, Article 463451. https://doi.org/10.1016/j.chroma.2022.463451

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Journal of Chromatography A

journal homepage: www.elsevier.com/locate/chroma

Small, smaller, smallest: Miniaturization of chromatographic process development



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ARTICLE INFO

Article history: Received 20 June 2022 Revised 12 August 2022 Accepted 24 August 2022 Available online 27 August 2022

Keywords: High-Throughput Screening Batch adsorption protein isotherms Microfluidic chromatography Liquid-Handling Stations Miniaturization

ABSTRACT

Biopharmaceuticals are becoming increasingly important in modern healthcare. Monoclonal antibodies (mAb) are one of the most widely used therapeutic proteins and are important for the treatment of cancer and autoimmune diseases, among others. After cell culture there are still large amounts of other impurities (e.g. host cell proteins) in solution. Chromatography is usually the first purification step, allowing to increase purity and reduce volume. This comes associated with high costs and chromatography accounts for a significant portion of total production costs for therapeutic proteins. Chromatographic process development may be time consuming and use large amounts of resins. Therefore, there is increased interest in finding cheaper techniques for chromatographic process development without compromising accuracy. This paper presents a highly sophisticated microfluidic chip approach for efficient adsorption isotherm determinations compared to current chromatographic process development. Implementation of an image analysis software ensures that chromatographic resin volume is accurately determined. The adsorption isotherm performance of microfluidics was compared to the robotic Liquid-handling Station (LHS) and labor intensive Eppendorf tubes. The microfluidic chip allows a 15-fold volume reduction and resin consumptions as low as 100/200 nl (200/100-fold reduction). The microfluidic chip performed comparably to the other miniaturized techniques, using less liquid and resin volume. For process development of expensive products (e.g. monoclonal antibodies), miniaturization (provided by the microfluidic chip) proved to be the most cost effective alternative whereas for less valuable products (e.g. lysozyme) automation (provided by the LHS) was the most cost effective alternative.

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1. Introduction

The past few years have seen increasing general public interest in the biopharmaceutical industry, mainly in the field of vaccine production due to the Covid pandemic. For the industry, a fast, reliable, and preferably cost-effective process development is important to respond to the market's needs. Patient accessibility to therapeutic proteins depends on several aspects, such as the cost of treatment or product availability. Although patent expiration considerably decreases the retail price of monoclonal antibodies (mAb), the price may still be prohibitive for patients [1,2]. High-Throughput Screening (HTS) is used to decrease time to market and reduce development costs, especially in early stages of process

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development. Moreover, increased use of mechanistic modeling of the processes combined with the ability to determine different parameters faster originated High-Throughput Process Development (HTPD). In the past decades HTPD has proven to be a valuable tool for faster and cheaper process development [3].

Chromatography plays a key role in the purification of biopharmaceuticals, as it usually is the first purification step in the downstream process and the one responsible for high purification factors [4]. However, this comes associated with high costs, since preparative chromatography can be very expensive. Therefore, chromatography accounts for a significant portion of the cost of producing therapeutic proteins [5]. Process development for chromatography involves the screening of different consumables (chromatographic resins, buffers) and conditions to find a suitable purification process. Equilibrium adsorption isotherms are amongst the important parameters to be determined for model based chromatographic process development [6]. Batch uptake experiments

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in 96 well-plates have also been used to calculate the partition coefficient and separation factor [7,8]. These give an indication of the purification capabilities of the tested systems. Isotherms provide insight on the thermodynamics of the studied systems, which are composed of the different buffers and the resin-protein pairs [9]. Previously, protein adsorption isotherms in batch uptake mode were determined using agitated vessels, with sample volume reaching up to hundreds of milliliters [10]. This method required large amounts of sample and resin. Current technology enabled researchers to reduce the amount of sample and materials, by using microtiter plates both with 96-well and 384-well format [11,12].

Researchers started to push for a manifold volume reduction, due to the high costs of chromatographic resin and sample waste and limited amount of sample available in early stages of process development (e.g. clinical trials). Different formats to determine adsorption isotherms emerged, like the use of plastic (Eppendorf) tubes, with volume requirements in the milliliter range [13]. Subsequently, Liquid-Handling Stations (LHS) were employed, reducing volume requirements to the sub-milliliter range. These offer a great automation to screen a plethora of consumables available for chromatographic process development [14]. Although LHS have become the status quo for both industry and academia [12,15], these platforms and consumables are expensive and still require a fair amount of product, which can be scarce in early stages of process development. It would be beneficial to reduce volume consumption in early-stage process development while maintaining accurate results. Microfluidics presents itself as a viable alternative for analytics and process development in many areas of the life sciences, often operating in the nanoliter range [16]. Although it has the obvious advantage of reducing volume (and, therefore, cost), microfluidics can also be more flexible and versatile than LHS, when the devices can be designed and produced from zero (for devices already commercialized this does not apply). Microfluidic devices have also been used for studying and screening conditions to determine protein adsorption isotherms [17], and protein purification [18]. Although these studies show the great miniaturization achieved by microfluidics, the use of a fluorescent label has proven to interfere with the protein, altering its properties [19,20]. A clear comparison of different miniaturization techniques is currently not available in the open literature.

This paper compares different miniaturization techniques for the determination of protein adsorption isotherms. It discusses the accuracy, usability, and technology readiness level (TRL) of each technique. Additionally, the cost of each screening solution is evaluated. Finally, this work shows how to use the small volume (Eppendorfs), smaller volume (LHS), and smallest volume (microfluidics) tools for chromatographic process development.

2. Materials and methods

2.1. Materials

Two different proteins were used in this study: lysozyme from chicken egg white (M_w of 14 300 Da, pl \approx 11; Sigma-Aldrich Chemie GbmH, Steinheim, Germany) and purified monoclonal antibody (M_w of 148 220 Da, pl \approx 8.6; Byondis B.V., Nijmegen, The Netherlands).

Sodium phosphate monobasic dehydrate, acetic acid (\geq 99.8%) and Tween® 20 were purchased from Sigma-Aldrich Chemie GbmH, Steinheim, Germany. Phosphoric acid (85%) and sodium hydroxide were purchased from Mallinckrodt Baker, Deventer, The Netherlands. Sodium Chloride was purchased from VWR-Chemicals, Leuven, Belgium. Di-sodium hydrogen phosphate was purchased from Merck KGaA, Darmstadt, Germany. SU-8 2100 photoresist was purchased from MicroChem, Newton, MA, USA and Developer mr-Dev 600 was purchased from micro resist technology

GmbH, Berlin, Germany. Polydimethylsiloxane (PDMS) was purchased as a Sylgard 184 elastomer kit (Dow Corning; Midland, MI, USA).

The resin used in the study was SP SepharoseTM Fast Flow (SP-Sepharose-FF) from Cytiva, Uppsala, Sweden. This is a strong cation-exchange agarose based resin and has a mean particle size of 90 μ m.

2.2. Buffers and solutions preparation

The lysozyme experiments were performed with sodium phosphate buffer containing 10 mM Na₂HPO₄ and varying NaCl concentrations adjusted to pH 6.5 with phosphoric acid. The mAb experiments were performed with buffer containing 25 mM NaOAc and 5 mM NaCl adjusted to pH 4.5 with 2 M NaOH. NaCl was added to the buffer solutions to adjust the total Na⁺ concentration of the solutions. All experiments were performed at room temperature. 0.01% w/v Tween® 20 was added to acetate buffer for the mAb experiments. The different buffers and solutions were prepared by dissolving the appropriate amount of chemical in Milli-Q water.

Lysozyme solutions were prepared by dissolving the appropriate amount of protein in the corresponding buffer. MAb solutions were used in the conditioning buffer that the protein was provided (mentioned above). All buffers and solutions were filter-sterilized using 0.20 μ m filters.

2.3. Microfluidic chip fabrication

The microfluidic chips were fabricated in two main steps: fabrication of the mastermolds and of the PDMS structures.

2.3.1. Production of mastermolds

The SU-8 mold (mastermold) was fabricated using direct write optical lithography (DWL). A SU-8 2100 layer of 100 μ m was spin coated on top of a clean Si substrate. The substrate was soft baked in a hotplate for 5 min at 65 °C and subsequently for 30 min at 95 °C. The mold was placed in the laserwriter (μ MLA Tabletop Maskless Aligner; Heidelberg Instruments Mikrotechnik, Heidelberg, Germany), where it was exposed with a 365 nm laser. The exposure energy used for this fabrication was 225 mJ/cm². After exposure, the substrate was placed in a hotplate for post exposure bake for 5 min at 65 °C and 10 min at 95 °C. Afterwards, the mastermold was developed by immersing it in mr-Dev 600 (Microresist, Berlin, Germany) and hand stirring for a minimum of 10 min. After this, it was rinsed with isopropyl alcohol (IPA) and spin-dried. Finally, the mastermold was hard baked for 15 min at 150 °C.

2.3.2. Fabrication of PDMS structures

The PDMS microchips were fabricated using standard PDMS soft lithography [21]. PDMS elastomer was prepared by mixing the elastomer base and curing agent in a (7:1) ratio. The mixture was degassed and placed on top off the mold. Simultaneously, a (20:1) mixture of elastomer base and curing agent was also prepared and degassed. This mixture was poured on top of glass slides and spin coated at 2500 rpm for 1 min. The covered mold and glass slides were baked at 80 °C for 45 min. After baking, the structure was peeled from the mold, and access holes (inlets and outlets) of 1.25 mm were punched using a hole-puncher. The structure was then placed on the PDMS-covered glass slides and sealed. The final structures were baked overnight at 80 °C.

2.4. Batch uptake adsorption isotherms

Adsorption isotherms provide information on the equilibrium concentration of a solute adsorbed to a solid phase (chromatographic resin) at different liquid concentrations. A known amount of protein is contacted with a known amount of resin and the equilibrium liquid concentrations are measured. The time to reach equilibrium varies according to the different systems. The amount of protein adsorbed to the solid phase was calculated by a mass balance, described by the following equation:

$$q_{eq} = \frac{V_l \times \left(c_{l,initial} - c_{l,eq}\right)}{V_r} \tag{1}$$

where q_{eq} is the protein adsorbed to the resin in equilibrium, V_l is the volume of liquid, V_r is the volume of resin, and $c_{l,initial}$ and $c_{l,eq}$ are the protein concentrations in the liquid phase in the beginning and after equilibrium is reached, respectively. Equilibrium adsorption isotherms were obtained with triplicate experiments.

2.4.1. LHS

Batch adsorption isotherm data were generated using a LHS (Tecan EVO Freedom 200 robotic station; Tecan, Switzerland). The LHS was equipped with an orbital mixer (Te-Shake), an automated vacuum system (Te-VacS), a multi-well plate reader (InfiniTe Pro 200), a robotic manipulator (RoMa) arm, two different liquid-handling arms (LiHa and MCA96) and a centrifuge system (Rotanta).

A known amount of resin (20.8 µL) was added to a 96-well filter plate (Pall Corporation, NY, USA) using a MediaScout® ResiQuot resin loader device from Atoll (Weingarten, Germany). To wash the resin, equilibration buffer was pipetted into the filter plate and it was shaken for 5 min at 1200 rpm, after which the solution was removed using the vacuum system. This cycle was performed 3 times in total. Protein solutions were subsequently pipetted (800 μ L) inside the well plates, which were sealed, and the plates were agitated at 1200 rpm until equilibrium was reached (2 h for lysozyme and 18 h for mAb). Once equilibrium was reached, the filter plate was placed on top of a 2 mL deep-well plate (Eppendorf AG, Hamburg, Germany) and these were centrifuged together using the centrifugation system. The supernatant was collected from the deep well plates to a UV star plate and the equilibrium concentrations were measured using the plate reader. Equilibrium concentrations were estimated using appropriate calibration curves, obtained using the LHS.

2.4.2. Eppendorfs

The batch adsorption isotherms in Eppendorf tubes were estimated using a similar method to the one described above. The studies were performed in 1.5 mL tubes. 800 μ L of protein solutions were prepared by hand into each of the tubes. The solutions were then contacted with 20.8 μ L of resin (generated using the same method as described for LHS). The tubes were rotated end-over-end at 10 rpm, until equilibrium was reached (the same was described above). After equilibrium was reached, the tubes were centrifuged (Sigma 112 from De Spatel B.V., The Netherlands), and the supernatant was hand-pipetted to a UV star plate, and the equilibrium concentrations were measured using the plate reader mentioned above. Equilibrium concentrations were estimated using appropriate calibration curves.

2.4.3. Microfluidic chip

2.4.3.1. Setup and mode of operation. The batch adsorption isotherms in the microfluidic devices were determined by recirculating protein solution through the microchip, that contained beads trapped inside. This is achieved with a closed system, where the same liquid is continuously pumped through the same resin volume, until equilibrium is reached. In the system, an inline UV–Vis detector (SPD-20AV; Shimadzu, Kyoto, Japan) with a microflow cell (0.2 μ L), allows to monitor the evolution of the adsorption over time.

Fig. 1 shows a schematic setup of the microfluidic chip. A peristaltic pump (ISM596D from Cole-Parmer, Wertheim, Germany) was used to recirculate the solution through the microchip. The pump used a PharMed® BPT tube (0.51 mm ID; Cole-Parmer, Wertheim, Germany), which was connected to PolyEtherEtherKetone (PEEK) tubes (0.18 mm ID; BGB, Harderwijk, The Netherlands). The PEEK tubes were directly connected to the microfluidic chip. The calculation of the equilibrium concentration was done using a calibration curve performed before each trial.

To perform the adsorption trials, the tube system was continuously primed with protein solutions of known concentrations, until a plateau was reached in the UV detector. After the plateau was reached, a protein solution of another concentration was successively primed until all the solutions for the calibration curve were primed. After this, the last solution to prime was the solution with the desired initial protein concentration. When the plateau of the signal was observed the pumping was stopped, and the PEEK tubes were directly connected to the microchip containing the chromatographic resin (see Fig. 1). After connecting, the pumping was resumed and the solution was recirculated until equilibrium was reached. Equilibrium was assumed to be reached when the signal stopped changing over time. The equilibrium concentration was used to calculate the mass balance to determine the amount of adsorbed protein in each trial.

2.4.3.2. Bead loading and volume determination for microfluidic chip. A suspension of 0.5% v/v chromatographic resin in storage buffer was prepared, by pipetting appropriate amounts of settled resin and storage buffer. The solution was vortexed and approximately 150 μ L of this solution was promptly pipetted directly into the microchip inlet using a micropipette. By suspending the resin and rapidly pipetting it into the microchip, it is avoided that the resin settles in the bottom of the pipette tip. This would block the microchip and hinder the entrance of the resin in the channel. Downstream of the channel there are frits. These structures, spaced 50 μ m from each other vertically and horizontally, will trap the beads inside the microchannel. After a sufficient amount of resin is loaded into the channel, the tip is removed and the microchip is analyzed in the microscope to estimate the total bead volume.

An accurate bead volume determination is paramount for the determination of the adsorption isotherms. Pictures of the loaded channel allow to estimate the number of beads and their radius. These images are then loaded into a MatLab script for image analvsis. Using the built-in function *imfindcircles* and appropriate parameters, it is possible to estimate the number of beads detected and the radius, in pixels, of each bead. Using an appropriate calibration (for example the channel's width, of 1000 µm), it is possible to correlate the bead radius in pixels with the distance per pixel (µm/px). With the information of the bead radius in micrometers, the bead volume is calculated using the formula for the volume of a sphere. An example of this bead volume determination can be found in Fig. 2. The beads are then washed with 200 µL of equilibration buffer. The interstitial porosity of the channel was not calculated since the experiments focus on studying the adsorption equilibrium of different proteins to a chromatographic resin. The microfluidic chip is used to mimic the batch adsorption mode of operation. Therefore, this parameter was considered to not be important since it provides information for the performance of packed columns but no relevant information for the equilibrium data between ligand and solute of interest.

2.4.3.3. Mass balance in microfluidic chip. The mass balance for the microfluidic chip is similar to the one presented in Eq. (1). However, for the microchip system, the initial protein concentration is determined for the tubing system, which doesn't account for the buffer volume present in the microchip. Therefore, an adjustment



Fig. 1. Microfluidic experimental setup for the determination of protein adsorption isotherms. A) Schematics of the setup. Yellow lines represent the PharMed® BPT tube that is connected to the peristaltic pump; black lines represent the PEEK tubing. Arrows show the direction of fluid flow through the system. B) Zoomed in image of the microfluidic chip with schematics for resin loading into the microchip. The resin is loaded using a micropipette tip and the chromatographic beads are trapped by the frits further down the channel.





Fig. 2. Image Analysis software. Top Left: Original image used as input for the program. Top right: visualization of the bead radius determination by the MatLab function imfindcircles. Bottom: Histogram distribution of the bead radius.

to the previous mass balance equation is needed:

$$q_{eq} = \frac{c_{l,initial} \times V_{tubing} - c_{l,eq} \times V_{system}}{V_r}$$
(2)

$$V_{system} = V_{tubing} + V_{\mu-chip} + V_{liquid\ plugs}$$
(3)

Where V_{tubing} is the volume of the tubes and the detector of the spectrophotometer, and $V_{\mu-chip}$ is the volume of the microchip. With these equations, all the buffer volume present in the final system (after the tubes are connected to the microchip) is considered. The volume of the "liquid plugs" ($V_{liquid plugs}$) in the mass balance was determined experimentally by measuring the height of the plug and considering a diameter of each inlet of 1.25 mm. This was also confirmed experimentally by running a test without resin inside the microchip and calculating the final liquid volume with the concentration after recirculation (data not shown). The total liquid volume for each microfluidics experiment was 56 µl and the resin volume per trial was around 200 nl (slight variations from experiment to experiment). The microchip system has a significantly larger volume than the microchip itself, mostly due to the tubing needed to operate the system. To reduce this vol-

Table 1

Overview of liquid and adsorbent volumes used in each experiment with each of the different methods.

Method	Adsorbent Volume (µl)	Liquid Volume (µl)	
LHS	20.8	800	
Eppendorf	20.8	800	
µ-chip	0.2	1.12	
μ-chip system (chip+tubing)	0.2	56	

ume further, the tube length reduction of the peristaltic pump can be considered. The total adsorbent and liquid volume used in each methodology is summarized in Table 1.

3. Results and discussion

3.1. Microchip design and operation

A microfluidic chip was designed to determine protein adsorption isotherms in batch uptake mode. The liquid phase (buffer solution containing protein) is permanently contacted with the solid



Fig. 3. Microfluidic chip design. A – Schematics of the whole microchip. B – Microscope image of the frits and outlet. The black circle is the hole punched with the punching tool and where the tube will be fit. C – Zoomed in image of the whole frit structure used. D – Zoomed in image of the top frits and indication of distance between frits. B, C and D are microscope pictures of the same structure, under different magnifications.

phase (chromatographic resin). Usually this is achieved by means of stirrers (e.g. magnetic stirrer in a glass flask), or by shaking the solution in shake flasks or orbital shakers (e.g., for the case of LHS). In the presented microfluidic chip the batch uptake is achieved by continuously pumping (by means of a peristaltic pump, Fig. 1) the protein solution through a microchannel where the chromatographic resin is trapped. The channel has a height of 100 µm, which can accommodate a plethora of resins used in the biopharmaceutical field (e.g. SP Sepharose FF (SP Seph FF) has an average diameter of 90 µm). To trap the beads, frits (pillar-like structures) were included in the design of the microchip. There are different methods one can apply to achieve this [22,23]. The chosen method was to have 5 frit columns next to each other. Compared to the studied alternatives (shafts, which is achieved by differences in height channels, narrowing the channel height to trap the resin), the frit design showed less flow constrains when performing simple flow simulations (data not shown). Each frit has a diameter of 50 µm, and the vertical and horizontal distance between frits was also set at 50 µm (Fig. 3). The liquid inlet (Fig. 3-A, right) was designed to have a circular shape that would allow for the bead loading, whereas the liquid outlet (Fig. 3-A, left) was designed with triangles to easily guide the liquid to the outlet and serve as an extra barrier in case of defected frits that would fail to trap the beads, thus avoiding major fouling in the tubes.

The liquid flow was ensured by a peristaltic pump. The liquid flowed from the inlet (Fig. 3A, right circular shape) to the outlet (Fig. 3A, left circular shape with triangle shapes). The beads were successfully trapped by the frit system, which is important to avoid fouling of the PEEK tubes used in the system. Besides SP Seph FF, beads of average particle diameter of 75 µm were also successfully trapped in the frit system. PDMS-based microfluidic chips can fail at relatively low pressures, so it is important that the pressure inside the flowing channel is as low as possible [24]. Therefore, a flow rate of 50 µl/min was employed, which enabled a good balance between a relatively low pressure in the channel, low bead compression against the frits, and fast enough recirculation through the system.

3.2. Image analysis

To determine protein adsorption isotherms and ensure results are reliable, it is important to have a good characterization of the concentration of protein in the liquid phase as well as the total resin volume that the protein solution contacts. For the first, inline monitoring of the absorbance values is used whereas for the second an image analysis program was developed. The crude images were uploaded to the program and the only processing required was the conversion from red green blue (RGB) to black and white and snipping the area to evaluate.

By using the function described in Section 2.4.3.2, it was possible to determine the radius of the chromatographic beads, given in pixels. The calibration used the channel's width (as shown in Fig. 3A) to convert the radius to µm. Fig. 2 shows the output of the program. The implementation of an appropriate calibration is important for the microfluidic chip results. Furthermore, ImageJ software was also used as an alternative to the described program to determine the resin volume. It had a worse performance for the intended purpose, as it was labor intensive and less accurate (data not shown). Therefore, it was decided that the most suitable method for chromatographic resin volume determination in the microchip was with the aforementioned program.

3.3. Determination of time to equilibrium

The monitoring of the adsorption of protein to the chromatographic resin was achieved by inline monitoring of the absorbance values at a wavelength of 280 nm (A_{280}). This allows to monitor in real time the adsorption of protein. In Fig. 4 it is possible to see the absorbance signal of two different systems throughout the time the experiments were performed. From this figure it is obvious that each system has its own characteristic time to equilibrium, which is the time it takes for the adsorption and desorption of protein molecules to the chromatographic resin to reach an equilibrium, meaning that the concentrations in the liquid and solid phase no



Fig. 4. Time to equilibrium of Lysozyme (A) and mAb (B) on Sp Seph FF. A – Lysozyme in 10 mM Phosphate buffer, pH 6.5, $[Na^+] = 50$ mM; B – mAb in 25 mM Acetate Buffer, pH 4.5, [NaCI] = 5 mM. Dashed line in A shows a closer look at the first 10 min of recirculation, where the signal oscillation can be explained by the presence of two phenomena: i) already some adsorption from the first pass of protein through the chromatographic beads and ii) local dilution of the solution due to the presence of protein-free solution present inside the microchip that is reaching the detector. Lysozyme systems reached the plateau around the 100-minute mark whereas mAb system reached the plateau around the 1000-minute mark.

longer change with time. The time to equilibrium varies dependent on the protein, buffer, and resin used in each experiment

After recirculation is started (time = 0 min), there is a lag time before a signal drop is observed. This is because the system was primed with a protein solution, which is still present in the tubes between the microchip and the detector. The first valley that can be seen in Fig. 4, and is highlighted in Fig. 4-A, is representative of two phenomena: i) the adsorption of the first "protein front" upon passing through the chromatographic resin for the first time, and ii) a local dilution of the solution due to the presence of proteinfree buffer inside the microchip. The signal gradually smoothens as a result of a better mixing and adsorption throughout the duration of the experiment. Eventually, the signal flattens and reaches a plateau, which indicates the time to equilibrium. It is noteworthy that for the tested resin and systems, the mAb showed the longest time to equilibrium (around the 1000 min mark), whereas for lysozyme the systems reached equilibrium before the 120 min mark. This difference can be explained by the smaller effective diffusivities that mAbs have compared to Lysozyme on SP Seph FF [25]. The larger mAb molecules take longer to migrate through the pores into the center of the particles, contributing to a longer time needed to access all available binding sites.

3.4. Protein adsorption isotherms

3.4.1. Lysozyme on SP Seph FF

The adsorption of Lysozyme to SP Seph FF was studied. Lysozyme has a very high pI, meaning that at the tested conditions its net charge will always be positive. Since SP Seph FF is a Cation-Exchange (CEX) resin, it was expected that a favorable adsorption behavior would be observed [26].

The isotherm data presented in Fig. 5 shows the adsorption isotherms of Lysozyme determined with the 3 different methodologies, as well as the fitted results to a Langmuir isotherm model (Eq. (4)). The adsorption of Lysozyme to SP Seph FF is highly favorable, visible by the rectangular shape of the three isotherm curves.

$$q = \frac{q_{max}KC}{1+KC} \tag{4}$$

The regressed parameter values for the q_{max} and K estimated from the fitting of the experimental data to Eq. (4) are summarized in Table 2. From the aforementioned figure and table, it is noticeable that the microfluidic chip estimated higher adsorption



Fig. 5. Adsorption isotherms of Lysozyme on SP Seph FF, in 10 mM Phosphate buffer pH 6.5, $[Na^+] = 50$ mM, using 3 different methodologies: (\blacktriangle) microfluidic chip, (\blacksquare) LHS and (\blacklozenge) Eppendorf tubes. Dashed lines represent the fitting of the different data sets to the Langmuir model. The parameters of the fitting can be found in Table 2.

Table 2	
Langmuir parameters of Lysozyme and mAb adsorption isotherms on SP Seph	ı
FF.	

Lysozyme			
Method	q _{max} (mg/ml)	K (ml/mg)	
µ-chip	136.4	17.8	
LHS	98.8	33.1	
Eppendorf	86.8	71.7	
mAb			
Method	q _{max} (mg/ml)	K (ml/mg)	
u-chip	148.0	5.6	
LHS	165.6	9.3	
Eppendorf	137.7	15.4	

values of Lysozyme to the SP Seph FF for the studied system. Although the estimated values for the microchip are higher than for the other two methodologies, these are still within the same order of magnitude. The fitted isotherms also show a similar shape, meaning that the microchip also shows the highly favorable adsorption behavior of Lysozyme to SP Seph FF.

Proteins can interact with PDMS and non-specific adsorption of proteins to the PDMS surface has been previously reported [27]. To understand if this phenomena was occurring in our system, some experiments with no resin inside the microchannel were performed. The results showed a flat signal over the duration of the trial (data not shown). Furthermore, there is a very low area of PDMS that contacts the solution at any given point in time and the liquid is being continuously pumped through the channel. Not only it is unlikely that there is non-specific adsorption, but this is greater when the solution is contacted statically with the PDMS surface [27]. The fluid flow through the microfluidic chip would contribute to prevent the non-specific adsorption of Lysozyme to the PDMS surface.

3.4.2. mAb on SP Seph FF

The adsorption of mAbs to SP Seph FF was also studied using the same setup. Initial experiments showed some instability in the UV signal of the solution containing mAb throughout the recirculation time. A recent study showed that different kinds of peristaltic pump tubes can affect the solutions [28]. The authors saw that the amount of nano and microparticles in solution increased over the pumping trials with water and buffer solutions, meaning that some particle shedding from the tubes to the solution is occurring. The study showed that there is particle shedding from the tube's material to the solution, and from the tested tubes, the one that shed the most particles was PharMed® BPT. Although the amount of particles present in protein-free solutions increased with the peristaltic pumping, this was much more pronounced when protein solutions were pumped. This could help explain that the pumping and particle shedding can influence the stability of the studied protein solution, potentially leading to more aggregation. Another study by Deiringer and Friess hypothesizes that protein particle formation is caused by the formation of a protein film in the surface of the tube and consequent tearing of the film caused by the pump rollers, thus releasing parts of this film in solution [29]. However, both studies show that the addition of surfactants to the proteins solutions would significantly reduce this phenomenon.

Keeping these studies' observations and our own in mind, the adsorption studies with the mAb were carried out with a solution containing 0.01% Tween® 20 to increase the solution's colloidal stability. LHS studies showed comparable adsorption between the solutions with and without the added surfactant (data not shown). Similarly to what was observed for Lysozyme, it was expected that mAb would have a favorable adsorption to SP Seph FF at the tested pH, as it is positively charged in these conditions. The results of the inline monitoring of the signal with the microfluidic chip, shown in Fig. 4, were used to define the time to equilibrium needed for this system.

Fig. 6 shows the adsorption isotherms of mAb with the three different methodologies, as well as the fitted results to a Langmuir model. From these results, it is possible to see that the adsorption behavior of mAb to SP Seph FF is favorable, characterized by the slope of the linear part of the isotherm. For this protein-resin pair it is noticeable the microchip results are more in line with what was determined by the other two methodologies. The regressed parameter values for the q_{max} and K estimated from the fitting of the experimental data to Eq. (4) are summarized in Table 2.

This system showed the most agreement between the three methodologies, with the Langmuir fitting for the microfluidic chip placed between the fitted Langmuir isotherms for the LHS and Eppendorfs. The microfluidic chip can, therefore, capture the adsorption behavior of the Lysozyme and mAb.



Fig. 6. Adsorption isotherms of mAb on SP Seph FF, in 25 mM Acetate buffer pH 4.5 and 5 mM NaCl, using 3 different methodologies: (\blacktriangle) microfluidic chip, (\blacksquare) LHS and (\bullet) Eppendorf tubes. Dashed lines represent the fitting of the different data sets to the Langmuir model. The parameters of the fitting can be found in Table 2.

3.5. Cost considerations

The adsorption isotherms results show that the microfluidic chip, the LHS, and the Eppendorf tubes are different methodologies that achieve very similar results for the intended purpose. It would be good to have a comparison between the three methodologies in terms of costs and compare the key features of each. For the cost considerations, four different methodologies were considered: LHS (both renting and purchasing one), microfluidic chip, and Eppendorf tubes. Three cost categories were considered: equipment costs, material costs and labor costs. These cost considerations take into account the current technology state of each methodology used for the studies adsorption studies.

The equipment costs comprise all the necessary equipment for the operation of each of the methodologies. The renting price of the LHS includes all the necessary equipment integrated in the LHS that is used for the determination of adsorption isotherms. For the microfluidics systems a shorter depreciation period was considered for the tubings and wafers due to the greater wear that these are subjected to and, therefore, shorter lifetimes. The equipment necessary for the fabrication of the mastermold, which is often very expensive, was not considered. The costs of the device used for the preparation of the resin plugs for the LHS and Eppendorf studies were not considered. The materials cost estimation is based on the materials and consumables needed to perform the experiments. The labor cost estimation is based on the hours needed of active labor, which is the time that an operator needs to actively work for the determination of the adsorption isotherms. For all the systems the reagent preparation (buffer and protein solutions) was considered. For the LHS studies, the operation was considered as the necessary time to prepare the LHS, resin plug preparation and supervision of initial stages of the operation, to ensure everything runs without errors. For the microfluidics studies, the chip production was considered, as well as microchip operation. The latter includes the time that the operator is required to operate the system and does not include the incubation time. For the Eppendorf studies, besides the reagent preparation, the pipetting of the solutions used for the study was also considered. The operation included the preparation of the resin plugs, the loading of the plugs to different Eppendorf tubes, and operating the rotor. The removal of the supernatant and subsequent dilution (when needed) of the solution in the 96 well-plate was also considered. A rate of 25€/h of labor costs was considered for the present study. The considered equip-

Table 3

Overview of the needed equipment, materials and labor hours needed for the determination of protein adsorption isotherms for the different methodologies studied.

Method	Equipment	Cost (ϵ)	Depreciation period	Part used for isotherm determination	yearly equipment $cost (e)$		
LHS	LHS	400,000	10	0.5	20,000		
LHS (rent)	LHS	10,000	-	1	10,000		
Microfluidic chip	Peristaltic Pump	3508	10	1	350.8		
	UV-Detector	15,000	10	1	1500		
	Other (tubes, connectors, etc.)	1000	1	1	1000		
	Wafer	410	0.5	1	820		
Eppendorf	Multi-well Plate Reader	10,000	10	1	1000		
	Rotor	586	10	1	58.6		
Materials							
Method	Materials needed			Amount ne	Amount needed per isotherm		
All	Resin			Dependent	Dependent on methodology		
	Buffer				Dependent on methodology		
	Protein			Dependent	Dependent on methodology		
LHS (Purch. & Rent.)	Fi	Filter plate			0.5 (pieces)		
	U	0.5 (pieces	0.5 (pieces)				
	Deep-Well Plate			0.5 (pieces)			
	Di	isposable Tips	0.5 (pieces)				
Microfluidic chip	PI	PDMS + Curing Agent			7.5 g		
Eppendorfs	Er	48 (pieces	48 (pieces)				
		V plate		0.5 (pieces	0.5 (pieces)		
Labor Hours							
Method	Ta	sk		Hours spen	nt per Task per isother		
All		eagent prep		1			
LHS (Purch. & Rent.)		Operation		1			
Microfluidic chip	M	icrochip produ	iction	0.5	0.5		
		peration		9	9		
Eppendorfs	Pi	petting to Epp	endorf tubes	1	1		
	Re	emoving super	natant	1.5	1.5		
	di	lution to UV-P	late	0.75			
	0	peration		1			

ment, materials and labor hours per isotherm for each methodology are summarized in Table 3.

The cost determination of the protein isotherms was based on the three components described above. This was done for two different scenarios: one where a "cheap" protein was considered (e.g. Lysozyme, which costs $0.03 \ \epsilon/mg$) and another where an "expensive" protein was considered (e.g. mAb, which was estimated to cost 2.29 ϵ/mg , based on the average price of the infliximab biosimilar in 2016 [30]). This was considering the determination of 100 protein adsorption isotherms per year. Besides this, an estimation of the variation of the cost per isotherm with the number of isotherms determined per year was also performed.

It is possible to see that, out of all the 4 options, the LHS have the highest material cost, for both scenarios (Fig. 7). This is because of the significantly larger liquid and resin volumes used by the technique (800 μ l and 20.8 μ l versus 56 μ l and 0.2 μ l, respectively) per data point. The Eppendorfs also use the same amount of liquid and resin volumes, but require less disposables than the LHS, hence why the materials costs are lower. This difference becomes less evident for the "expensive" protein scenario since the costs related to the sample represent the higher portion of materials costs for these two methods (LHS and Eppendorfs). However, the lesser consumables use comes at the expense of added labor and, consequently, higher labor costs.

The advantage of automation is reflected in the lower labor costs of the LHS in both scenarios and proves to be an advantage when low-value proteins are studied. On the other hand, the miniaturization of the adsorption studies proves to be very advantageous when high-value proteins are studied. Increasing protein costs will change the cost driver of LHS and Eppendorf to the materials whereas the cost driver of the microfluidic chip will always be the labor costs. The labor needed for the isotherms studies considered the microfluidic chip methodology as it was previously described. However, this could be reduced by increasing the automation of the system, bringing the Labor costs down. Therefore, the cost performance of the LHS is dependent on the materials prices. Consequently, a high degree of miniaturization is preferred for very expensive proteins and a high degree of automation is preferred for cheap materials.

An increased number of isotherms to be determined per year also contributes to a "dilution" of the costs per isotherm for the LHS (see Fig. 8). This is mainly due to a decreasing contribution of the equipment costs for an increasing number of isotherms determined per year. As expected, the microfluidics costs per isotherm plateau at a relatively low number of isotherms to be determined per year; this is because the labor costs are the main cost driver and the labor per isotherm is independent of the number of isotherms to be determined per year. A similar trend can be observed for the Eppendorfs, as the main cost driver for the cheap protein is the labor, but for the expensive protein is the materials. For the renting of the LHS, it was considered that only 120 isotherms could be determined in a 3-month rental period (2 isotherm per day for a 60 working day period). Renting the equipment two times a year would already bring the yearly costs of the equipment at the same level as purchasing the LHS, which would then be considered to be more cost-effective. It is important



Fig. 7. Breakdown of the costs for the determination of protein adsorption isotherms, for a LHS (purchasing and renting), microfluidic chip and Eppendorf tubes, for a base case of 100 isotherms determined per year. The costs are normalized relative to the methodology of the scenario that has the highest overall cost (LHS for the Expensive Protein scenario). Left – Scenario for the "expensive" protein; the most expensive method for this scenario was the LHS (purchasing), with an isotherm to be estimated to cost 665*c*. Right – Scenario for the "cheap" protein; the most expensive method for this scenario was the microfluidic chip. The percentages show what is the percent contribution of the Labor, Materials and Equipment cost for each methodology in each scenario (e.g. for LHS in the "cheap" protein scenario, 8% of the isotherm costs were Labor costs, 15% Material costs and 77% Equipment costs).



Fig. 8. Variation of the price per isotherm with increasing number of isotherms determined per year, for the 4 different methodologies considered. A minimum number of 20 isotherms to be determined per year was considered and for the LHS rent, a maximum number of 120 isotherms per year was considered. A – Scenario for the "cheap" protein; B – Scenario for the "expensive" protein.

to mention that maintenance costs are not being considered for this study, which could impact more the yearly equipment costs of owning the equipment and for renting this would not represent added costs.

3.6. Key features of the different methodologies

LHS remain the *status-quo* for chromatographic process development for industry and academia. The LHS rely on their high degree of automation and precision and have a higher Technology Readiness Level than the other two methodologies presented in this study. Furthermore, and since it has been widely used by industry and academia for more than a decade, there is a wide variety of consumables dedicated to chromatographic process development. However, the equipment has a high price tag, not affordable for every lab.

Microfluidics has been taking steps in the field of bioengineering and there are some microfluidics devices that are commercialized (mainly for analytics [14]). At a lower TRL, the microfluidic chips present, generally, a low level of automation, and on-chip automation can hinder the versatility of the devices. If the operation of the microchips can have a higher level of automation, the labor costs can be reduced further. Nevertheless, microfluidics still offers the highest degree of miniaturization possible, which in the case of this study, that used an external source for fluid pumping, still reduced the liquid volume to 56 µl. The tested devices were used only once, but further testing for the removal of the beads from inside the channel, by reverse flow, could prove that the devices are reusable. Since PDMS has a high compatibility with acid and base, the microfluidic chip can be cleaned with most solutions used for the removal of proteins and regeneration of chromatographic resins. The use of solvents may pose a problem for PDMS, as these can react with PDMS (e.g. acetone). However, using lower concentrations of these solves (e.g. 70% v/v EtOH) for short periods of time should be sufficient for the intended purpose and still safe to the PDMS chip. Using different materials for the microfluidic chip (e.g. glass or quartz) would also improve the microchip's mechanical and chemical resistance.

If the devices can be reused, it would help to reduce the labor costs. Furthermore, a common approach for microfluidics is parallelization, which for this case could also be an interesting option, provided that a system could be designed to allow this. Microfluidics is still trying to pave the way into HTS for chromatographic processes and the high skill required to design and produce the designs makes it a less attractive alternative. This could be overcome with outsourcing and mass production of the devices or by finding alternative production techniques (like 3D printers).

Of all the methodologies in this work, Eppendorfs proved to be the one with the lowest level of automation. The need to prepare all the solutions by hand, removal of supernatant and subsequent dilution in the UV-plate (when needed) required tremendous amounts of pipetting, leading to lengthy runs in the lab and tedious and cumbersome work. However, it made up by being the methodology with the most easily accessible materials (which are common in every lab) and the lowest equipment costs of the three methodologies.

4. Conclusions

Lysozyme and mAb adsorption isotherms were successfully determined using three different methodologies: microfluidic chip, Liquid-Handling Station, and Eppendorf tubes. An in-house designed and produced microfluidic chip showed comparable results to the other two methodologies. Inline monitoring of the absorbance of the protein solution allowed the microfluidics setup to estimate the time to equilibrium required for the different systems. This can be especially relevant when systems that are not reported in the literature are studied (e.g., the mAb used in this study), thus allowing for saving time and material for such study in a LHS. Industry perceives microfluidics as a viable option to contribute to HTPD [31], and our study showed that microfluidics can compete with the *status-quo* of HTPD for chromatography.

For the correct estimation of adsorbed protein, accurate protein concentration in the liquid phase is needed (usually achieved by using spectrophotometry) as well as accurate resin volume determination. For the latter, devices for the generation of resin plugs are used for the LHS (and the same device was used for the Eppendorf experiments in this study), but microfluidics resin volume determination was often imprecise [18] or cumbersome [17]. By implementing the Image Analysis in our studies, we were able to accurately estimate the resin volume in each microfluidics experiment.

The microfluidic chip allowed to perform protein adsorption studies to chromatographic resin using only 200 nl of resin and with a total system volume of 56 μ l. This represents a 100-fold reduction in resin and a 15-fold reduction in solution. Such large reductions in material and sample proved to be advantageous when considering the cost of isotherm studies for expensive proteins. However, the large labor costs meant that it was not cost-competitive when cheap proteins were studied. Of the three methods used, microfluidics presents the lowest TRL. Further studies could help increase automation of the proposed system, thus reducing labor costs of microfluidics. The large level of automation offered by LHS is still very attractive, and depending on the de-

sired total use, investing in one can be the best option. However, if no intensive studies are needed year-round, renting a LHS can be a viable option and the most cost-competitive (Fig. 8). Although the Eppendorf studies showed a good cost performance, the experimental work involved in the isotherm studies does not allow for a good throughput and more workers would be needed.

Of the three methodologies, Eppendorfs are the least attractive but could still be used for lower throughputs if companies or academia are not willing to invest in a LHS or the microfluidic chip. For intensive studies on protein adsorption isotherms, the LHS is still the best option, since an increase in throughput (if needed) does not directly translate in an increase in labor. On the other hand, if the studied protein is very expensive and/or the amount of material is very low (e.g. due to low expression levels or in early stages of process development), the proposed microfluidic chip, in its current TRL stage, is an attractive alternative to the LHS. Further studies and improvements in the microfluidic chip production and possible parallelization could increase the competitiveness of the microfluidic chip, even for studies involving cheaper samples. This study showed that the microfluidic chip was able to generate protein adsorption isotherms using a fraction of the materials required by other methodologies.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

CRediT authorship contribution statement

Tiago Castanheira Silva: Conceptualization, Methodology, Software, Validation, Formal analysis, Investigation, Writing – original draft, Visualization. **Michel Eppink:** Conceptualization, Resources, Writing – review & editing. **Marcel Ottens:** Conceptualization, Resources, Supervision, Writing – review & editing, Funding acquisition.

Acknowledgements

This work has received funding from the European Union's Horizon 2020 research and innovation program under the Marie Skłodowska-Curie grant agreement No 812909 CODOBIO, within the Marie Skłodowska-Curie International Training Networks framework.

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