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1	Spectroscopic and Quartz Crystal Microbalance (QCM) Characterization of Protein-
2	based MIPs
3	
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13	
14	Abstract
15	We have studied acrylamide-based polymers of varying hydrophobicity (acrylamide, AA; N-
16	hydroxymethylacrylamide, NHMA; N-isopropylacrylamide, NiPAm) for their capability of
17	imprinting protein. Rebinding capacities (Q) from spectroscopic studies were highest for
18	bovine haemoglobin (BHb) MIPs based on AA, Q = 4.8 $\pm$ 0.21 < NHMA, Q = 4.3 $\pm$ 0.32 <
19	NiPAm, $Q = 3.6 \pm 0.45$ , while also demonstrating low selectivities for non-template proteins
20	( $<30 \pm 5\%$ ), with the exception of bovine serum albumin (BSA, $>76 \pm 0.5\%$ ). When applied
21	to the QCM sensor as thin-film MIPs, NHMA MIPs were found to exhibit best discrimination

22 between MIP and non-imprinted control polymer (NIP) in the order of NiPAm < AA <

23	NHMA. The extent of template removal and rebinding, using both crystal impedance and
24	frequency measurements, demonstrated that 10% (w/v):10% (v/v) sodium dodecyl
25	sulphate:acetic acid (pH 2.8) was efficient at eluting template BHb (with $80 \pm 10\%$ removal).
26	Selectivity studies of NHMA BHb-MIPs revealed higher adsorption and selective recognition
27	properties to BHb (64.5 kDa) when compared to non-cognate BSA (66 kDa), myoglobin
28	(Mb, 17.5 kDa), lysozyme (Lyz, 14.7 kDa) thaumatin (Thau, 22 kDa) and trypsin (Tryp, 22.3
29	kDa). The QCM gave frequency shifts of $\sim 1500 \pm 50$ Hz for template BHb rebinding in both
30	AA and NHMA MIPs, whereas AA-based MIPs exhibited an interference signal of $\sim$ 2200 ±
31	50 Hz for non-cognate BSA in comparison to a $\sim$ 500 ± 50 Hz shift with NHMA MIPs. Our
32	results show that NHMA-based hydrogel MIP are superior to AA and NIPAM

34 Keywords: Molecular imprinted polymer (MIP); Hydrogel; Protein; Biosensor; QCM

#### 35 **1. Introduction**

### 36 1. Introduction

In recent years, molecularly imprinted polymers (MIPs) have allowed selective extractions that rival immunoaffinity-based separations, and have shown clear advantages over real antibodies for sensor applications: they are easy to fabricate, intrinsically stable, robust, and are able to operate in extreme environments [1], [2] and [3]. MIPs could provide an alternative, inexpensive, fast, and efficient diagnostic method for highly sensitive analytical procedures within the pharmaceutical area [3].

43

44 When imprinting complex bio-macromolecules some of the most significant drawbacks in MIP technology are the unprecedented degree of influence that the variation in pH [4], ionic 45 strength and local matrix effects all have on the gel properties [5], [6], [7] and [8]. This can 46 affect the three dimensional shape and chemical characteristics of the template molecule 47 during polymerisation. This is particularly true when imprinting large bio-macromolecules 48 such as proteins. Proteins are relatively labile, and have variable conformations which are 49 sensitive to solvent environments, pH and temperature, all of which present a variety of 50 challenges [5]. It has been thought that low imprinting capacities associated with bio-51 macromolecules could be caused by the use of charged functional monomers causing non-52 specific electrostatic interactions [5]. Moreover, as with antibodies, MIPs have also shown a 53 degree of cross-selectivity, in that they can bind molecules similar to the native template and 54 55 cause non-specific binding. It is thought that this is due in part to an excess of functional monomer molecules being randomly distributed and frozen within the imprinted cavity 56 during polymerisation that have an affinity for non-template molecules [3] and [9]. Thus, 57 more sophisticated monomers capable of forming better, stronger and more stable 58

interactions that offer better positioning and complementary functionality are widely being
sought. Once these parameters are optimised, application to biosensors and analysis of actual
biological samples would be more realistic [6], [10], [11] and [12].

62

63	Biosensors for proteins are currently expensive to develop because they require the use of
64	expensive antibodies [3] and [13]. However, as MIPs are becoming more and more promising
65	as viable alternatives to natural receptors new MIP-based sensor strategies are being
66	developed [3]. The main advantage of biosensors is the ability to sample outside the
67	laboratory environment with minimal user input. One important part of bio-sensing is
68	transducers, which monitor the reaction between bio-selector and analyte. Among various
69	physical transducers (electrochemical, peizoelectric, etc.), mass sensitive devices such as
70	surface acoustic wave (SAW), surface plasmon resonance (SPR) and quartz crystal
71	microbalance (QCM) have become popular for sensing applications [14], [15], [16] and [17].

72

Following the thorough analysis of QCM systems for use in fluids over the past 2 decades, 73 74 this has allowed for more esoteric applications including bio-sensing [16]. In most cases quartz resonators are integrated to oscillator circuits to form a QCM for micro weighing 75 applications. Normally, an equivalent circuit model is fitted to the impedance curve, and the 76 obtained parameters can be used for calculating the resonant frequency and dissipation (D) of 77 the quartz crystal i.e. mass and viscoelastic properties of the deposited layers [15] and [16]. 78 79 Determining the impedance curve has many advantages; first and foremost it has expanded the range of measurable parameters from rigid thin films, to biologically relevant films of 80 81 "soft" viscoelastic material. These QCM couplings have been widely used for biomaterials 82 and biosensor studies [10], [12], [16], [18] and [19], where surface confined bio-molecular

interactions have provided an insight into dissolution of polymer coatings, DNA 83 hybridisation, cell response to pharmacological substances, molecular interactions of drugs 84 85 and their delivery. The QCM has also been utilised as an immunosensor, where analytes are recognised by antibodies, which are immobilised on a thin layer deposed on a crystal surface. 86 Resulting mass changes are transformed into an electronically measurable quantity. The 87 objective behind the majority of QCM research is to use sensor technology to develop a rapid 88 89 method for the measurement of bio-molecular affinity reactions, and an in-depth analysis of electrochemical deposition, adsorption and reaction mechanisms of polymers coated on 90 91 electrodes as 'thin films' [10], [12], [18], [19] and [20].

92

93 The focus of this paper is the tailoring of QCM electrode surface chemistry (i.e. specialised polymer coatings), with a view that these devices can discriminate proteins for bio-sensing 94 and basic surface-molecular interaction studies. In this work, we demonstrate the application 95 96 of the QCM technique to distinguish between the behaviour of MIPs and NIPs in the 97 presence of cognate and non-cognate proteins. Bovine haemoglobin (BHb, 64.5 kDa) was chosen as a model protein for its well-known function in the vascular system as a carrier of 98 99 oxygen, also in aiding the transport of carbon dioxide and regulating blood pH [3] and [13]. Bovine serum albumin (BSA, 66 kDa), a non metalloprotein of similar molecular weight to 100 BHb, served to test the selectivity of the BHb-MIP to BSA compared to template BHb, and 101 102 was compared across a family of acrylamide-based polymer hydrogels.

103

104 2. Experimental

105 2.1. Materials

107	Acrylamide (AA), N-hydroxymethylacrylamide (NHMA), N-iso-propylacrylamide (NiPAm),
108	N,N'-methylenebisacrylamide (bis-AA), ammonium persulphate (APS), N,N,N',N'-
109	tetramethylethylenediamine (TEMED), sodium dodecyl-sulphate (SDS), glacial acetic acid
110	(AcOH), bovine haemoglobin (BHb), bovine serum albumin (BSA), hen egg-white lysozyme
111	(Lyz), thaumatin from Thaumatococcus danielii (Thau), bovine pancreatic trypsin (Tryp) and
112	equine heart myoglobin (Mb) were all purchased from Sigma-Aldrich, Poole, Dorset, UK.
113	Sieves (75 $\mu m$ ) were purchased from Endecotts Ltd. and Inoxia Ltd., UK. AT-cut quartz
114	crystal pieces (9 MHz fundamental resonance) with gold-on-chrome electrodes were supplied
115	by Nihon Dempa Kogyo Company Ltd. (Tokyo, Japan).
116	
117	2.2. HydroMIP preparations
118	
119	Hydrogel MIPs were synthesised by separately dissolving AA (54 mg), NHMA (77 mg),
120	NiPAm (85.6 mg) and bis-AA as cross-linker (6 mg), (8.5 mg) and (9.5 mg), respectively
121	along with template protein (12 mg) in 960 $\mu$ L of MilliQ water. The solutions were purged
122	with nitrogen for 5 min, followed by an addition of 20 $\mu L$ of a 10% (w/v) APS solution and
123	20 $\mu L$ of a 5% (v/v) TEMED solution. Polymerisation occurred at room temperature (RT, 22
124	$\pm$ 2 °C) giving total gel densities (%T) of 6%T, AA/bis-AA (w/v); 8.5%T, NHMA/bis-AA
125	(w/v); 9.5%T, NiPAm/bis-AA (w/v), and final crosslinking densities (%C) of 10%C (9:1,
126	w/w) for all hydrogels.

For every MIP created a non-imprinted control polymer (NIP) was prepared in an identical 128 manner but in the absence of protein. After polymerisation, the gels were granulated 129 separately using a 75 µm sieve. Of the resulting gels, 500 mg were transferred into 1.5 mL 130 131 centrifuge Eppendorf tubes and conditioned by washing with five 1 mL volumes of MilliQ water followed by five 1 mL volumes of 10% (w/v):10% (v/v) SDS:AcOH (pH 2.8) and 132 another five 1 mL volume washes of MilliQ water to remove any residual 10% (w/v):10% 133 134 (v/v) SDS:AcOH eluent and equilibrated the gels. Each wash step was followed by a centrifugation, whereby the gels were centrifuged using an Eppendorf mini-spin plus 135 136 centrifuge for 3 min at 6000 rpm (RCF:  $2419 \times g$ ). All supernatants were collected for analysis by spectrophotometry to verify the extent of template removal. It should be noted 137 that the last water wash and eluent fractions were not observed to contain any protein. 138 139 Therefore we are confident that any remaining template protein within the MIPs did not 140 continue to leach out during the rebinding studies.

141

142 2.3. Rebinding studies

143

Once the gels (500 mg) were equilibrated, a 1 mL template protein solution prepared in 144 MilliQ water containing 3 mg of protein was added to the target MIPs and NIP controls and 145 was allowed to associate at RT ( $22 \pm 2$  °C) for 20 min. Selectivity studies were also 146 conducted to assess the relative imprinting factor of the original protein template. This was 147 148 achieved by loading non-cognate proteins on a BHb imprinted gel. Gels were then washed with four 1 mL volumes of MilliQ water. Each reload and wash step for all MIPs and NIP 149 150 controls was followed by centrifugation at 6000 rpm (RCF:  $2419 \times g$ ) for 3 min. All supernatants were collected for analysis by spectrophotometry. 151

# 153 2.4. Spectrophotometric analysis

155	Calibration curves in MilliQ water and 10% (w/v):10% (v/v) SDS:AcOH were prepared for
156	BHb, BSA, Lyz, Tryp and Mb. Spectral scans revealed peak wavelengths for BHb in MilliQ
157	water and 10% (w/v):10% (v/v) SDS:AcOH to be 405 nm and 395 nm, respectively. Peak
158	wavelengths for BSA in MilliQ water and 10% (w/v):10% (v/v) SDS:AcOH were found to be
159	288 nm and 290 nm respectively. Peak wavelengths for Lyz in MilliQ water and 10%
160	AcOH:SDS were found to be 291 nm and 296 nm respectively. Peak wavelengths for Tryp in
161	MilliQ water and 10% (w/v):10% (v/v) SDS:AcOH were found to be 293 nm. Peak
162	wavelengths for Mb in MilliQ water, 10% (w/v):10% (v/v) SDS:AcOH were found to be 410
163	nm, and 396 nm respectively. Analysis and subsequent determination of protein
164	concentration in appropriate media was performed at specific peak wavelengths using a UV
165	mini-1240 CE spectrophotometer (Shimadzu Europa, Milton Keynes, UK).
166	
167	2.5. Quartz crystal microbalance (QCM) analysis of thin film MIPs
168	
169	QCM crystals were sealed and air capped (single-sided) with PVC glue in-order to prevent
170	short circuiting when the QCM was submerged in solution [10]. Poly AA, NHMA and
171	NiPAm gels for BHb were synthesised using the hydrogel production procedures outlined in
172	Section 2.2. Before polymerisation, MIPs and NIPs were deposited as thin films onto the
173	capped QCM crystals. Thin-films were achieved by beading and compressing 10 $\mu$ L of the
174	polymerising solutions directly onto the crystals. QCM frequency and impedance

175	measurements were taken using an Agilent 4194A Impedance Analyser. An in-house written
176	QBasic programme was used to drive the analyser and collect series resonance frequency and
177	impedance data in real time.

179 2.5.1. Elution and rebinding studies

180

MIP and NIP polyAA thin-film capped crystals were firstly immersed in MilliQ water,
followed by 10% (w/v):10% (v/v) SDS:AcOH in order to remove imprinted protein primarily
from the surface of the polymer. This was followed by another submersion in MilliQ water to
remove any residual surfactant and to re-condition the hydrogel. After subsequent
stabilisation of the QCM response, template protein was reloaded by immersing the QCM in
a 3 mg/mL BHb solution and the response trace was recorded at RT (22 $\pm$ 2 °C).

188 2.5.2. Selectivity studies

189

190 Continuous real-time scans were conducted in-order to assess characteristic impedance 191 changes of the gels during surface exposure to wash, elute and protein rebinding conditions. 192 During a typical run, the MIP thin-film capped crystals were submerged sequentially in 193 various solutions such as 10% (w/v):10% (v/v) SDS:AcOH, MilliQ water or 3 mg/mL protein 194 solutions (cognate BHb and non-cognate BSA, Thau, Lyz and Tryp) for a set time of 5 min 195 each, and crystal impedance and frequency responses were recorded at RT ( $22 \pm 2$  °C). The 196 latter procedure was followed for AA, NiPAm and NHMA based MIP hydrogels for BHb.

### 198 3. Results and discussion

199 3.1. MIP selectivity

The molecular imprinting effect is characterised by the rebinding capacity (Q) of protein to the gel polymer (mg/g) exhibited by the protein-specific MIP and the control NIP, and is calculated using Eq. (1), where Ci and Cr are the initial protein and the recovered protein concentrations (mg/mL) respectively (which specifies the specific protein bound within the gel), V is the volume of the initial solution (mL), and g is the mass of the gel polymers (g).

205

$$Q = \frac{[C_i - C_r]V}{g}$$
(1)

207

Fig. 1A shows the rebinding capacities and imprinting factors of polyacrylamide (polyAA) MIP and NIPs for several different proteins. The internal measure of the imprinting factor between MIP and NIP serves to demonstrate that the MIP possesses selective cavities for the rebinding of template molecule compared to NIP controls. It can clearly be seen that there is a distinctive rebinding capacity variation for each imprinted protein template within a polyAA MIP. This has previously been attributed to protein size, cross-linking density, and the initial degree of association within the polymer matrix [4].

215 Gels based on N-hydroxymethylacrylamide (NHMA) exhibited similar rebinding trends,

216 whereas poly-N-isopropylacrylamide gels (polyNiPAm) demonstrated lower rebinding

217 capacities. Thus, bulk gel characterisation revealed the highest rebinding capacities for BHb

MIPs based on polyAA ( $Q = 4.8 \pm 0.21$ ), followed by polyNHMA ( $Q = 4.3 \pm 0.32$ ),

polyNiPAm (Q =  $3.6 \pm 0.45$ ). These gel imprinting trends are in agreement with those previously published [4], [9] and [10].

221

Selectivity studies were also conducted to confirm a BHb specific imprinting effect and to assess the relative imprinting factor of cross-selective binding profiles. The cross-reactivity of the BHb-imprinted MIPs for non-cognate proteins was quantified using relative imprinting factors (k), Eq. (2), where IFBHb is the imprinting factor for BHb, and is calculated by IF = [Ci - Cr]MIP/[Ci - Cr]NIP, and IFx is the imprinting factor of the cross-reacting noncognate proteins on a BHb MIP. For the template BHb k = 1, and for non-cognate proteins that are less-specific for the BHb MIP, k < 1.

229

230 
$$k = \frac{\mathrm{IF}_{BHb}}{\mathrm{IF}_{x}}$$
(2)

The data (Fig. 1B) suggests that both non-cognate trypsin (Tryp) and lysozyme (Lyz) proteins 231 have relatively low affinities for the BHb-specific MIP, k  $\approx 0.2 \pm 0.05$ . However, bovine 232 233 serum albumin (BSA), which is a similar size to BHb, exhibited a high degree of interference binding (cross selectivity) resulting in high k values of  $0.65 \pm 0.05$ . Myoglobin (Mb) also 234 exhibited some degree of cross-selectivity; this can be attributed to the size of Mb, which is a 235 quarter that of BHb (17.5 kDa), and its similarity to a single BHb sub-unit [4]. Interestingly 236 though, when reversed, a polyAA BSA-MIP exposed to non-target BHb protein had 237 relatively low affinity. It would appear that BSA has a high ability to bind non-specifically to 238 a BHb MIP, whereas BHb does not exhibit the same ability within a BSA MIP. 239

Competitive binding studies using a 50:50 mixture of BHb:BSA (3 mg/mL total) on a MIP-241 BHb were also conducted (Fig. 1B). The addition of BSA caused an obvious capacity 242 decrease of BHb binding on the BHb-MIP, suggesting that the rebinding of BHb was 243 displaced by the competing BSA or by protein-protein interactions [21]. As the size, 244 structure, and specificity of the imprinted cavities should be in favour of the BHb template, it 245 is rational that the addition of BSA as a competing protein would not bind to the BHb-246 247 specific imprinted cavities. Gai et al. previously demonstrated that BSA does not bind specifically to a BHb MIP, but rather displaces the non-specific recognition sites of cavities 248 249 and the nonspecific binding of BHb to BHb-MIP [21]. Moreover, although BSA and BHb share similar sizes (66 kDa and 64.5 kDa, respectively), it should be noted that BSA has a pI 250 of 4.6 and BHb a pI of (6.8–7.0). Since competitive binding was performed under MilliQ 251 252 water (pH 5.4), conditions are in favour of BSA [21] and [22]. Above its pI, BSA becomes negatively charged and the groups exist as single bondNH2 and single bondCOO-, this 253 overall negative net charge induces more favourable and complementary hydrogen bonding 254 interactions, resulting in increased specific as well as non-specific binding [4]. 255

256

257 3.2. QCM sensor application of MIPs

Thin film BHb MIPs were prepared on the surface of a QCM chip and the sensor was exposed sequentially to MilliQ water, 10% (w/v):10% (v/v) SDS:AcOH and 3 mg/mL protein solutions at RT ( $22 \pm 2 \,^{\circ}$ C). We have previously published on the thickness of the thin films on sensor chips with an average thickness of  $138 \pm 9 \,$ nm [6]. Given that for a 9 MHz crystal the shear wave decay length is 250 nm at RT [23], we are within the sensing region of the QCM to measure both bulk and surface effects within the MIP film.

Fig. 2A and B shows the QCM impedance and frequency responses following immersion in a 265 solution of 10% (w/v):10% (v/v) SDS:AcOH in order to remove imprinted protein from the 266 surface of the polymer. Previous investigations have shown that optimum conditions for 267 protein removal of up to 80% have been achieved using 10% (w/v):10% (v/v) SDS:AcOH 268 [9]. Using this acid/surfactant combination the positively charged protein attaches to the 269 negatively charged surface of SDS micelles and disrupts the hydrophobic bonds. Since there 270 271 is a significant shift in both resonance frequency and impedance it can be assumed that some of the BHb imprinted template has been successfully removed from the MIP. 272

It is worth noting the two distinct differences in the impedance response when compared with 273 the frequency response. Firstly, the impedance response has much reduced noise in the signal 274 in contrast to the frequency response. Secondly, there are significant additional transitions 275 (e.g. at 350 and 650 s) in the signal which are being observed in the Z response, but not in the 276 277 frequency response. It has been suggested that whereas the frequency response predominately 278 demonstrates the QCM mass response only within an adlayer, the electrical impedance gives a combination response of the mass effect as well as subsequent changes in the viscoelasticity 279 280 of the adlayer possibly due to molecular relaxations within the adsorbed layer over a longer timescale following initial immersion [23], [24] and [25]. 281

282

After subsequent stabilisation of the QCM response, the template BHb was then reloaded by immersing the QCM in a 3 mg/mL BHb solution and the response trace recorded. Fig. 2C and D compares the final QCM impedance and frequency responses to template BHb exposure of each MIP and its corresponding NIP. It can be seen that upon addition of a 3 mg/mL BHb solution to the BHb MIP caused significant QCM responses compared with NIP thin-film hydrogels. This suggests that MIP thin-film gels are affected by specific binding of target

BHb. This distinct difference between responses exhibited by MIP and the NIP control strongly supports that binding and elution of target protein gave rise to distinct impedance transitions. The  $200 \pm 50$  Hz frequency shift observable by both MIP and NIP during the initial loading step (Fig. 2D) is suggestive of a solution viscosity effect.

293

294 Real time impedance response following sequential immersion in solutions of BHb, 10% (w/v):10% (v/v) SDS:AcOH and BSA were also measured. Three distinct types of responses 295 were observed depending on the acrylamide-based monomer used. The key difference 296 between the polymers is their hydrophilicity dictated by the hydrophilic hydroxyl group in 297 NHMA and the hydrophobic isopropyl group in NiPAm. AA sits between the two in degree 298 299 of hydrophilicity (polyNHMA > polyAA > polyNiPAm), which agrees with the order of best performance of the polymers as BHb MIPs in previous QCM studies [10]. Fig. 3 compares 300 the final QCM response to cognate and non-cognate protein exposure of each MIP with its 301 302 corresponding NIP. Interestingly, the NiPAm MIP and NIP both show a near zero frequency 303 response to template BHb and non-cognate BSA, indicating that NiPAm is equally unselective for both proteins as is the control non-imprinted polymer. The non-response of 304 NiPAm to either BHb or BSA suggests that there is a resistance to either protein to bind to 305 the polymer. The striking difference in selectivities between cognate and non-cognate 306 proteins for NHMA and AA suggests that the hydroxyl group in NHMA plays a significant 307 308 role in the selective binding of BHb and the lack of binding of BSA.

Moreover, variations of the series resonance frequency demonstrated to be highly dependent on the test solution used (Fig. 4A). The impedance data is presented here because in comparison to the frequency response, there is much reduced noise in the signal following each solution phase immersion. It can be seen that MIP thin-films exposed to a 10%:10%

313 (v/v) SDS:AcOH solution exhibited an immediate significant decrease in impedance (500  $\pm$ 314 100  $\Omega$ ); this is possibly due to the increase in the viscosity of the solution caused by the 315 presence of SDS micelles in the solution. Moreover, it can clearly be seen that the 316 introduction of non-template BSA also exhibits an impedance response within the poly AA 317 MIP, suggesting some non-specific binding within the BHb-HydroMIP. Thus, there is a high 318 degree of cross-selectivity present for our AA-based MIPs (>70%), and this interference is 319 absent when NHMA-based MIPs are used (<20%) as seen in Fig. 3.

To further test the BHb-MIP selectivity, we investigated the rebinding of template BHb after 320 exposing the MIP with non-target BSA. The resulting quantified imprinting effect of BHb for 321 322 polyAA MIP thin-film gels can be seen in the impedance responses (Fig. 4B). The comparison demonstrates that both HydroMIP and HydroNIP films act differently under 323 water wash, elution and load (solution immersion) treatments following BSA loading. It can 324 325 be seen that when non-target BSA is loaded first, the QCM impedance response is now negligible. Interestingly, impedance responses are also almost negligible when BHb is 326 introduced after prior exposure to BSA. Although the loaded BSA did not associate 327 specifically with the BHb-MIP thin-film surface, an interesting and lasting effect inhibits 328 329 BHb from easily binding to recognition sites. Indeed, BSA is similar to the template BHb in 330 size, but the spatial arrangement of the effective groups on its exterior are different from BHb, and the recognition sites in the BHb-MIP cavities are not complementary in shape to 331 BSA [22]. Therefore, little to no selectivity of BSA on BHb-MIPs should be expected. 332 333 Therefore, in the case of AA-based MIPs, the inhibition effect is most likely due to the ability of BSA to exhibit protein binding on the MIP surface but not within cavities. Formation of a 334 335 BSA biolayer above (but not within) the cavities would block subsequent cavity-selective MIP binding for its cognate protein [22]. This is further indication that BHb-MIPs distinguish 336 proteins not only based on molecular size, but also on the synergistic effect of shape memory, 337

and multiple weak hydrogen bonding interactions specific to template protein inmacromolecular recognition [21], [22] and [26].

340

Moreover, further studies to interrogate the recognition capabilities of MIPs were carried out 341 using a range of non-metalloproteins chosen for their different sizes and functionalities 342 compared to BHb, BSA and Mb. Of these proteins: lysozyme (Lyz), a glycoside hydrolase 343 enzyme (14.7 kDa) that is part of the innate immune system, and exists as a natural form of 344 protection from pathogens like Salmonella, E. coli, and Pseudomonas [9], [10] and [27]. 345 Thaumatin (Thau), a sweetener or flavour modifier (22 kDa) often used in crystallisation 346 studies due to its ease of use and crystal formation [27]. Trypsin (Tryp), a serine protease 347 enzyme or proteinase 'digestive enzyme' (23.8 kDa) commonly imprinted within MIPs [9], 348 [10] and [27]. Fig. 5 shows that the BHb-MIPs based on all three polymers (AA, NHMA, and 349 NiPAm) are essentially non-responsive to the addition of the three smaller proteins Thau, Lyz 350 351 and Tryp respectively. An average NIP response was calculated based on all three polymers 352 and used as an illustration to demonstrate the negligible responses exhibited by the MIP properties. The negligible responses exhibited by the QCM sensor concur with the qualitative 353 data and confirm that these small proteins exhibit no selective specific/non-specific binding 354 characteristics to a BHb-imprinted MIP. 355

356

357 4. Conclusions

A family of acrylamide-based MIPs have been characterised for their imprint efficiency using spectrophotometric and QCM sensor techniques for biosensor development. Varied rebinding capacities and relative imprinting factors have been achieved using bulk characterisation. We

361 have demonstrated that MIP selectivity is a function of the hydrophilicity of the acrylamide monomer used to form the MIP. Three distinct types of QCM responses were observed 362 depending on the acrylamide used (polyNHMA > polyAA > polyNiPAm), which agrees with 363 364 the order of best performance of the polymers in previously published QCM studies. The selectivity of BHb-MIP for BHb and BSA was also compared via QCM, along with several 365 other proteins. Results demonstrated BHb-MIP to have better selective adsorption and 366 367 recognition properties to BHb than BSA when using the hydrophilic NHMA as a MIP polymer matrix. Therefore, the QCM sensor was able to indicate MIP surface activity and 368 369 provide physical interpretation in terms of hydrophilicity of the polymer matrix that forms the MIP and protein selectivity. Our QCM sensor also has the ability to assess the extent of 370 specific protein binding by sensing surface-specific bound cognate protein to MIPs compared 371 372 to non-imprint NIP controls. We expect, once fully developed, that the benefits of sensitivity, specificity and stability of MIPs coupled with discriminatory techniques, such as QCM, will 373 be crucial to the future impact of portable diagnostics for personal healthcare and use by 374 375 health professionals. The technology also presents major potential benefits to environmental and food monitoring. 376

377

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381

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465 Fig. 1.

466 (A) Rebinding capacities (Q) and imprinting factors of MIP<sub>polyAA</sub> and NIP<sub>polyAA</sub> hydrogels for several
 467 proteins in MilliQ water media: bovine haemoglobin (BHb), bovine serum albumin (BSA), myoglobin

- 468 (Mb), lysozyme (Lyz), trypsin (Tryp); (B) relative imprinting factors (k) for a range of BHb-
- 469 MIP<sub>polyAA</sub> cross-reactants in MilliQ water media. Results illustrate higher MIP selectivities for cognate
- 470 BHb and the degree of cross-selectivity for other non-template analytes. Data represents
- 471 mean  $\pm$  S.E.M., n = 3.
- 472



473

474 Fig. 2.

475 QCM response to the immersion of polyAA-BHb hydrogel thin-film MIP in 10% (w/v):10% (v/v)

476 SDS:AcOH in order to follow protein elution (arrow indicates time of MIP immersion): (A) impedance

477 (ΔZ), (B) frequency (Δf); QCM responses to BHb (3 mg/mL) loading onto a BHb imprinted polyAA

478 hydrogel thin-film (arrow indicates time of immersion in protein solution): (C) impedance ( $\Delta Z$ ) and (D)

- 479 frequency ( $\Delta f$ ).
- 480







- 483 Fig. 3.

484 QCM frequency shift responses of NiPAm, AA and NHMA polymer MIPs and NIPs to cognate BHb

485 and non-cognate BSA loading (3 mg/mL) after 5 min of exposure. Data represents

486 mean  $\pm$  S.E.M., n = 3.

487 488



489

490 Fig. 4.

491 Real time QCM impedance responses: (A) direct BHb rebinding and BSA cross-selectivity on a BHb-

492 MIP<sub>polyAA</sub>; (B) BSA cross-selectivity on a BHb-MIP<sub>polyAA</sub> followed by BHb rebinding.





496 Fig. 5.

497 QCM response of functionalised acrylamide BHb MIPs to non-cognate proteins thaumatin (Thau),

498 Iysozyme (Lyz), and trypsin (Tryp) after H<sub>2</sub>O washes and an SDS:AcOH elute.

