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# Towards Rational Design of Selective Molecularly Imprinted Polymers (MIPs) for Proteins: Computational and Experimental Studies of Acrylamide-Based Polymers for Myoglobin

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#### 1. Statistical Thermodynamics

In the SI we present a detailed derivation of the expressions used to compute the monomer binding probabilities and the average number of bound monomers. The related formulas are usually expressed in terms of binding constants. In contrast, here, we employ estimates of the binding free energies  $\Delta G_{PL}$ .

#### 1.1 Protein with M binding sites.

We consider a protein P with *M* independent, non-equivalent sites, indexed by the integers  $i = \{1, ..., M\}$ . We assume that each protein site can be empty or bind one monomer L molecule. Binding of monomer L at site *i* is associated with a change in the standard-state free energy change  $\Delta G_{PL}(i)$ . The resulting grand-canonical partition function system is (1)

$$\Xi_{PL} = \prod_{j=1}^{M} \left( 1 + \mathbf{x} e^{-\beta \Delta G_{PL}(j)} \right)$$
(SI-1)

where  $x \equiv [L]/C_0$ , with [L] the *free* monomer concentration in solution and  $C_0 = 1 \frac{\text{mol}}{\text{L}} = \frac{1}{1660 \text{ A}^3}$  the standard-state concentration. The quantity  $\Delta G_{PL}$  is

defined in Eq. 1 of the main text.

The average number of monomer L molecules bound to a protein molecule is

$$\overline{M}_{L}(x) = \sum_{i=1}^{M} \frac{x e^{-\beta \Delta G_{PL}(i)}}{1 + x e^{-\beta \Delta G_{PL}(i)}}$$
(SI-2)

To employ Eq. (SI-2), we need to know the concentration x of *free* monomer. Since each protein molecule binds on average  $\overline{M}_{L}(x)$  ligand molecules, the concentration of *bound* monomer is  $[L]_{bound} = [P]_{tot} \overline{M}_L(x)$ , where  $[P]_{tot}$  is the total protein concentration. The concentration of free monomer is then:

$$[L] = [L]_{tot} - [L]_{bound} = [L]_{tot} - [P]_{tot} \overline{M}_{PL}(x)$$

with  $[L]_{tot}$  the total monomer concentration in solution. Using Eq. (SI-2), we obtain:

$$\frac{[L]}{C_0} = x = \frac{[L]_{tot}}{C_0} - \frac{[P]_{tot}}{C_0} \sum_{i=1}^{M} \frac{x e^{-\beta \Delta G_{PL}(i)}}{1 + x e^{-\beta \Delta G_{PL}(i)}}$$
(SI-3)

A similar equation has been derived for equivalent sites in reference (2). Eq (SI-3) can be solved self-consistently to yield the concentration  $x = x^*$  of free monomer in solution. The solution  $x = x^*$  can then be used in eq. (SI-2), to compute the average number of bound monomers at a protein molecule.

#### 1.2 Solutions with two types of monomers

Suppose that the solution contains two different monomers  $L_1$  and  $L_2$ . These monomers can bind at the same protein sites with free energy changes  $\{\Delta G_{PL_1}(i)\}$  and  $\{\Delta G_{PL_2}(i)\}$ , respectively. The grand-canonical partition function is now

$$\Xi_{PL_1L_2} = \prod_{i=1}^{M} \left( 1 + x_1 e^{-\beta \Delta G_{PL_1}(i)} + x_2 e^{-\beta \Delta G_{PL_2}(i)} \right)$$
(SI-4)

[compare with expression (SI-1)].

The average numbers of monomers of types  $L_1$  or  $L_2$ , bound to a protein molecule, are

$$\overline{M}_{L_1}(x_1) = \sum_{i=1}^{M} \frac{x_1 e^{-\beta \Delta G_{PL_1}(i)}}{1 + x_1 e^{-\beta \Delta G_{PL_1}(i)} + x_2 e^{-\beta \Delta G_{PL_2}(i)}}$$
(SI-5a)

and

$$\overline{M}_{L_2}(x_2) = \sum_{i=1}^{M} \frac{x_2 e^{-\beta \Delta G_{PL_2}(i)}}{1 + x_1 e^{-\beta \Delta G_{PL_1}(i)} + x_2 e^{-\beta \Delta G_{PL_2}(i)}}$$
(SI-5b)

By analogy with Eq. (SI-3), the free-ligand concentrations can be evaluated by solving self-consistently the system of equations:

$$x_{1} = \frac{[L_{1}]_{tot}}{C_{0}} - \frac{[P]_{tot}}{C_{0}} \sum_{i=1}^{M} \frac{x_{1}e^{-\beta \Delta G_{PL_{1}}(i)}}{1 + x_{1}e^{-\beta \Delta G_{PL_{1}}(i)} + x_{2}e^{-\beta \Delta G_{PL_{2}}(i)}}$$

$$x_{2} = \frac{[L_{2}]_{tot}}{C_{0}} - \frac{[P]_{tot}}{C_{0}} \sum_{i=1}^{M} \frac{x_{2}e^{-\beta \Delta G_{PL_{2}}(i)}}{1 + x_{1}e^{-\beta \Delta G_{PL_{1}}(i)} + x_{2}e^{-\beta \Delta G_{PL_{2}}(i)}}$$
(SI-6)

The solutions  $x_1 = x_1^*$  and  $x_2 = x_2^*$  can be substituted in Eqs. (SI-5), to yield the average monomers of types  $L_1$  and  $L_2$  bound to a protein molecule. Note that in general the free monomer concentrations differ  $(x_1^* \neq x_2^*)$ , even if the solutions are prepared with equal total monomer concentrations  $[L_1]_{tot} = [L_2]_{tot}$ .

The probability for site i to be occupied by monomer  $L_1$ , regardless of the occupancy state of other sites, is given by:

$$p(\text{ site i is occupied by } L_1) = \frac{X_1 e^{-\beta \Delta G_{PL_1}(i)}}{1 + X_1 e^{-\beta \Delta G_{PL_1}(i)} + X_2 e^{-\beta \Delta G_{PL_2}(i)}}$$
(SI-7a)

Similarly, the probability for site i to be occupied by monomer  $L_2$ , regardless of the occupancy state of other sites, is

$$p(\text{site i occupied by } L_2) = \frac{x_2 e^{-\beta \Delta G_{PL_2}(i)}}{1 + x_1 e^{-\beta \Delta G_{PL_1}(i)} + x_2 e^{-\beta \Delta G_{PL_2}(i)}}$$
(SI-8b)

The ratio of the above probabilities is

$$\frac{p(\text{site i is occupied by } L_1)}{p(\text{site i is occupied by } L_2)} = \frac{x_1 e^{-\beta \Delta G_{PL_1}(i)}}{x_2 e^{-\beta \Delta G_{PL_2}(i)}} \cong \frac{e^{-\beta \Delta G_{PL_1}(i)}}{e^{-\beta \Delta G_{PL_2}(i)}}$$
(SI-9)

The last equality holds if the *free* concentrations of the two monomers are approximately equal. Eq. (SI-9) was employed in the calculation of site-specific relative binding probabilities of the two co-monomer solutions reported in Table 2.

### 2. Tables S1 to S4

	,	0					
	X	Y	Z				
Site 1	17.620	-23.778	12.972				
Site 2	20.229	-23.142	32.053				
Site 3	7.083	-30.952	23.353				
Site 4	8.547	-21.480	-2.489				
Site 5	2.500	-13.810	7.800				
Site 6	2.968	-7.392	27.891				
Site 7	24.100	-36.302	20.533				
Site 8	26.717	-35.043	11.150				
Site 9	5.538	-18.823	34.450				
Site 10	-2.100	-29.760	1.650				
Site 11	-4.436	-29.653	13.807				
Site 12	-5.550	-15.110	18.950				
Site 13	4.850	-17.796	18.379				
Site 14	14.183	-22.010	27.283				

**Table S1:** Glide protein grid coordinate centers in Å, defined by inner box 10 Å  $\times$  10 Å  $\times$  10 Å, for each binding site.

Table S2: The top-ranked GlideScore value for each of the five monomers (AAm, NHMAm, NHEAm, DMAm,
TrisNHMAm), docked using Glide-SP into each of the 14 binding sites of myoglobin as predicted by SiteMap. <sup>a</sup>

	Binding Site													
	1	2	3	4	5	6	7	8	9	10	11	12	13	14
AAm	-1.79	-2.70	-2.17	-2.23	-2.72	-2.51	-2.04	-1.95	-2.14	-2.28	-2.17	-2.00	-1.85	-1.42
NHMAm	-1.01	-1.33	-1.46	-1.10	-0.93	-1.60	-1.41	-0.61	-1.36	-1.46	-2.06	-1.69	0.37	n/a
NHEAm	n/a	-1.54	-1.55	-1.07	-0.93	-1.21	-1.31	-0.78	-1.12	-1.70	-0.49	-1.60	n/a	n/a
DMAm	n/a	-2.74	-2.90	-2.53	-2.17	-2.65	-2.02	-2.03	-1.89	-2.68	-2.08	-1.88	n/a	n/a
TrisNHAm	-0.31	0.51	0.65	0.46	0.13	0.02	0.81	0.86	-0.98	-0.26	1.32	-0.77	n/a	n/a

<sup>a</sup> GlideScores that are positive are highlighted in bold. *n/a* indicates that no binding poses were obtained.

Monomer	Binding Site													
	1	2	3	4	5	6	7	8	9	10	11	12	13	14
AAm														
$\Delta E_{PL}^{MM}$	-21.76	-32.66	-20.4	-25.89	-33.01	-25.96	-24.70	-18.62	-21.67	-28.98	-15.78	-16.86	-17.46	-20.49
$\Delta G_{PL}^{solv}$	10.65	12.96	5.71	13.01	14.80	12.05	13.63	5.22	9.66	15.17	5.61	5.00	10.65	14.83
$-T\Delta S_{PL}^{RRHO}$	13.59	10.72	10.94	10.65	10.92	11.07	12.65	10.13	10.82	10.21	10.25	10.19	14.12	14.85
$\Delta G_{PL}$	2.5	-9.0	-3.8	-2.2	-7.3	-2.8	1.6	-3.3	-1.2	-3.6	0.1	-1.7	7.3	9.2
NHMAm														
$\Delta E_{PL}^{MM}$	-21.94	-34.33	-29.5	-13.49	-43.54	-35.21	-31.93	-27.81	-27.04	-33.86	-19.65	-24.13	10.50	n/a
$\Delta G_{PL}^{solv}$	6.83	10.84	10.84	16.74	21.45	18.32	18.57	10.71	9.73	11.80	6.69	11.91	9.77	n/a
$-T\Delta S_{PL}^{RRHO}$	11.74	12.66	13.56	12.98	12.26	13.98	14.01	11.77	12.47	12.13	12.27	12.74	16.50	n/a
$\Delta G_{PL}$	-3.4	-10.8	-5.1	16.2	-9.8	-2.9	0.7	-5.3	-4.8	-9.9	-0.7	0.5	36.8	n/a
NHEAm														
$\Delta E_{PL}^{MM}$	n/a	-34.85	-34.54	-31.24	-41.55	-36.82	-33.29	-27.53	-27.88	-36.83	-15.97	-29.8	n/a	n/a
$\Delta G_{PL}^{solv}$	n/a	11.33	13.65	12.96	20.04	18.75	18.76	8.94	11.35	11.52	10.54	14.30	n/a	n/a
$-T\Delta S_{PL}^{RRHO}$	n/a	12.23	13.89	12.38	12.91	14.91	14.01	11.97	12.17	12.42	12.29	12.35	n/a	n/a
$\Delta G_{PL}$	n/a	-11.3	-7.0	-5.9	-8.6	-3.2	-0.5	-6.6	-4.4	-12.9	6.9	-3.2	n/a	n/a
DMAm														
$\Delta E_{PL}^{MM}$	n/a	-34.54	-25.67	-30.38	-29.2	-27.69	-24.79	-23.49	-32.66	-35.18	-29.67	-25.79	n/a	n/a
$\Delta G_{PL}^{solv}$	n/a	14.32	5.18	11.84	10.31	11.84	10.46	6.97	17.32	14.12	15.30	12.78	n/a	n/a
$-T\Delta S_{PL}^{RRHO}$	n/a	11.46	11.35	12.04	11.59	12.01	15.02	11.08	11.08	10.41	11.08	10.70	n/a	n/a
$\Delta G_{PL}$	n/a	-8.8	-9.1	-6.5	-7.3	-3.8	0.7	-5.4	-4.3	-10.7	-3.3	-2.3	n/a	n/a
TrisNHAm														
$\Delta E_{PL}^{MM}$	-38.13	-32.84	-38.49	-42.96	-43.49	-44.76	-25.38	-42.58	-22.26	-45.93	-32.97	-28.51	n/a	n/a
$\Delta G_{PL}^{solv}$	14.44	12.48	14.44	18.02	17.35	21.63	10.15	18.39	7.20	8.14	14.49	10.33	n/a	n/a
$-T\Delta S_{PL}^{RRHO}$	13.40	15.17	13.36	14.22	14.03	15.76	14.74	12.90	13.75	12.93	13.32	12.99	n/a	n/a
$\Delta G_{PL}$	-10.3	-5.2	-10.7	-10.7	-12.1	-7.4	-0.5	-11.3	-1.3	-24.9	-5.2	-5.2	n/a	n/a

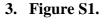
**Table S3**: The breakdown of  $\Delta G_{PL}$  into its contributions for each of the five monomers (AAm, NHMAm, NHEAm, DMAm, TrisNHMAm) calculated using MM-GBSA at each the 14 predicted binding sites of myoglobin.<sup>a</sup>

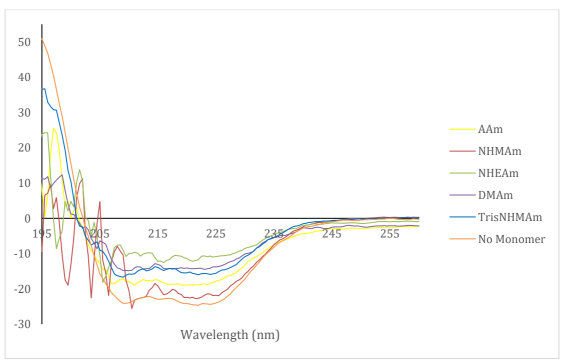
<sup>a</sup> Values that are positive (unfavourable binding) are highlighted in bold. *n/a* indicates that there were no predicted binding poses.

**Table S4:** The  $\Delta E_{Hbond}$  contributions for each of the five monomers (AAm, NHMAm, NHEAm, DMAm, TrisNHMAm) calculated using MM-GBSA at each the 14 predicted binding sites of myoglobin.<sup>a</sup>

	Monomer										
<b>Binding Site</b>	AAm	NHMAm	NHEAm	DMAm	TrisNHMAm						
1	-1.02	-0.51	n/a	n/a	-1.70						
2	-1.51	-2.13	-1.97	-0.89	-1.00						
3	-1.18	-1.01	-1.02	-0.49	-1.70						
4	-1.18	-1.30	-1.34	-0.74	-1.06						
5	-1.96	-1.42	-1.34	-0.74	-1.06						
6	-1.28	-1.77	-1.66	-1.03	-1.26						
7	-1.5	-2.04	-2.25	-1.21	-0.48						
8	-0.76	-1.08	-1.09	-0.54	-0.96						
9	-2.39	-1.53	-1.66	-2.45	-0.30						
10	-1.80	-1.74	-1.71	-1.08	-2.35						
11	-1.63	-1.32	-0.79	-0.58	-0.74						
12	-1.84	-1.25	-0.99	-1.04	-0.82						
13	-2.04	n/a	n/a	n/a	n/a						
14	-1.83	n/a	n/a	n/a	n/a						

<sup>a</sup> n/a indicates that there were no predicted binding poses.





**Figure S1:** CD spectroscopic analysis of myoglobin after being mixed with the five monomers: AAm (yellow), NHMAm (red), NHEAm (green), DMAm (purple), TrisNHMAm (blue) and no monomer (orange), at a protein:monomer ratio of 1:1081, the same as the polymerisation solution used in hydrogel MIP formation.

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