

# Retention of Histidine-Containing Peptides on a Nickel Affinity Column

Igor A. Kozlov, Bahram G. Kermani, Peter C. Melnyk, David L. Barker, Chanfeng Zhao, John P. Hachmann, and Michal Lebl\*

Illumina, Inc., 9885 Towne Centre Drive, San Diego, CA 92121

## Abstract

The retention of histidine-containing peptides in immobilized metal-affinity chromatography is studied using several hundred modeled peptides. Retention is driven primarily by the number of histidine residues; however, the amino acid composition in the immediate vicinity plays a significant role. Specifically, the arginine and tryptophan content has to be taken into consideration. During the course of this study, an alternative tag that can be used similarly to a polyhistidine tag is discovered.

## Introduction

Immobilized metal-affinity chromatography (IMAC) is an important tool for the purification of proteins containing residues that form metal complexes (histidine, cysteine, and tryptophan) (1–4). The polyhistidine tag is extremely useful in molecular biology where it serves to facilitate the isolation of genetically engineered proteins from complex mixtures and can be used for the targeted immobilization of these proteins (4,5).

A high-throughput method for the purification of peptide-oligonucleotide conjugates is developed. One of our strategies is to place three histidines at the amino terminus of the peptide and three histidines at the 5' end of the oligonucleotide. When joined together, the six histidines should form a tag that can be bound to a Ni-Sepharose affinity column. After washing away the unreacted components, the purified peptide-oligo conjugate can be eluted with a gradient of increasing imidazole concentration. For this strategy to be successful, the concentration of imidazole that elutes three histidines (His3) must be significantly less than the concentration that elutes six histidines (His6). In addition, the elution concentration for His6 should be relatively insensitive to the identity of amino acids surrounding the His6 tag.

Surprisingly, there is little information available in the literature on the relative affinity for Ni-Sepharose, in the presence of imidazole gradients, of polyhistidine-containing peptides and the influence of surrounding amino acids. Consequently, an

array of model compounds containing different numbers of histidines in various sequential arrangements and in combination with various other amino acids is synthesized. This manuscript presents comprehensive information that should facilitate the design and purification of engineered peptides and proteins.

## Experimental

Fluorenylmethyloxycarbonyl (Fmoc) amino acids, (benzotriazol-1-yloxy)tris(dimethylamino)phosphonium hexafluorophosphate (BOP) reagent, and Rink resin (0.42 mmol/g) were purchased from Novabiochem (EMD Biosciences, San Diego, CA). Solvents were from VWR International, Inc. (West Chester, PA). 4-Methylpiperidine was from Sigma-Aldrich (Milwaukee, WI).

Rink resin (300 mg) was added into a mixture of dimethylformamide (DMF) and dichloromethane (DCM) (10 mL total) to form a non-sedimenting suspension, which was distributed into the wells of flat bottom polypropylene microtiterplates (Evergreen Scientific, Los Angeles, CA). The plates were placed into a centrifugal synthesizer (6,7). An additional 100  $\mu$ L of DMF was added into the plate wells (beads sedimented), and the plate was centrifuged with a tilt of 6 degrees. A standard protocol was used for the synthesis to remove the Fmoc protecting group. 4-Methylpiperidine was used instead of piperidine (8). Individual Fmoc protected amino acids (0.3M solution in 0.3M HOBt in DMF) were pipetted to the wells, and a solution of BOP (0.6M in DMF) and 1.2M diisopropylethylamine (DIEA) in DMF was delivered to each well. Plates were oscillated five times and allowed to rest for 50 s. During oscillation, the plates were rotated at a speed at which the liquid does not overflow over the wall of the well and solid support moves towards the outer side of the well. When the rotation was stopped, liquid returned to the horizontal position and beads distributed at the well bottom, thus mixing the well content. This procedure was repeated 30 times. The plate was centrifuged, and the addition of amino acids and reagents was repeated. After another 30 cycles of oscillation and pausing, the reagents were removed by centrifugation and washing, and deprotection was repeated to prepare the plate for the next cycle of synthesis.

\* Author to whom correspondence should be addressed: email mlebl@illumina.com.

Table I. Structure of Model Peptides and Their Retention (Eluting Imidazole Concentration) on Nickel Column

#	Peptide	Im concentration (M)	#	Peptide	Im concentration (M)	#	Peptide	Im concentration (M)
1	GAGAHGAGAGY	0.083	46	GAHHGAGHHAY	0.150	96	RREHHHHHEEY	0.180
2	GAGAHHGAGAY	0.098	47	GHHAGAGHHAY	0.155	97	AAKHHHHAEAY	0.180
3	GAGAHHHGAGY	0.138	48	RRHHGGHHRRY	0.215	98	DDHHHHDDKY	0.180
4	GAGAHHHHGAY	0.190	49	RRHHGGHHHEEY	0.128	99	DDKHHHHKDDY	0.180
5	GAGHHHHHGAY	0.240	50	RSHHRSFHRSY	0.220	100	AKAHHHHAAAY	0.180
6	GAHHHHHHGAY	0.275	51	RSHHESHHSY	0.168	101	SSSHHHHSSSY	0.183
7	GHHHHHHHGAY	0.328	52	ESHHRSHHESY	0.135	102	EKHHHHKEKY	0.183
8	GHHHHHHHHAY	0.370	53	GAHHEEHHGAY	0.115	103	AAKHHHHAAEY	0.183
9	HHHHHHHHHAY	0.408	54	GAHHEGHHGAY	0.133	104	RDDHHHHDDRY	0.185
10	HHHHHHHHHHY	0.460	55	GAHHHHHGAY	0.138	105	RSRHHHHSEY	0.188
11	HGAGAGAGAGY	0.090	56	GAHLLHHHGAY	0.143	106	EERHHHHREEY	0.188
12	HHAGAGAGAGY	0.085	57	GAHHERHHGAY	0.145	107	AKAHHHHAAEY	0.188
13	HHHGAGAGAGY	0.120	58	GAHHGGHHGAY	0.150	108	KKHHHHDDKY	0.188
14	HHHHAGAGAGY	0.165	59	GAHHSGHHGAY	0.150	109	KKHHHHKDDY	0.188
15	HHHHHGAGAGY	0.210	60	GAHHGHHGAY	0.150	4	GAGAHHHHGAY	0.190
16	HHHHHHGAGAY	0.243	61	GAHHLGHHGAY	0.150	110	ILVHHHHVLIY	0.190
17	HHHHHHHGAGY	0.288	62	GAHHFGHHGAY	0.163	111	KEHHHHKEKY	0.193
18	HHHHHHHHAGY	0.325	63	GAHHFFHHGAY	0.173	112	RAAHHHHAAAY	0.193
			64	GAHRRHHHGAY	0.190	113	AARHHHHADAY	0.193
2	GAGAHHGAGAY	0.098				114	AARHHHHDAAY	0.193
19	GAGAHGAGAY	0.108	65	HHHAGAGHHHY	0.263	115	KKHHHHDKDY	0.193
20	GAHGAGAGAY	0.098	66	HHHAGAHHHGY	0.255	116	AAHHHHAAKY	0.193
21	GAHGAGHAGAY	0.095	67	HHHAGHHHAGY	0.258	117	AARHHHHEAAY	0.195
22	GAHGAGAHGAY	0.098	68	HHHAHHHGAGY	0.263	118	KKHHHHKEKY	0.195
			69	HHHGHHHGAGY	0.273	119	RRDHHHHDDDY	0.195
3	GAGAHHHGAGY	0.138	16	HHHHHHHAGAY	0.243	120	AARHHHHAAAY	0.195
23	GAHHGHAGAGY	0.128				121	GGGHHHHGGGY	0.198
24	GAHHGAGAGY	0.123	4	GAGAHHHHGAY	0.190	122	AARHHHHAEAY	0.198
25	GAHHGAGHAGY	0.120	70	EEEEHHHEEY	0.127	123	KKHHHHKEEY	0.198
26	GAHHGAGAHGY	0.121	71	ESEHHHHHEEY	0.128	124	KKHHHHKEEY	0.198
			72	EEHHHHHEEY	0.128	125	ARAHHHHAAAY	0.198
4	GAGAHHHHGAY	0.190	73	EEHHHHHEEKY	0.140	126	AARHHHHAAEY	0.200
27	GAHHGHHAGAY	0.155	74	REEHHHHHEEY	0.145	127	RRRHHHHDDDY	0.200
28	GAGHHGHHGAY	0.165	75	KEHHHHHEEY	0.150	128	RAAHHHHAAEY	0.203
29	GAGHHAHHGAY	0.163	76	EKKHHHHKEEY	0.163	129	KAHHHHAAKY	0.203
30	GAHHAHHGAGY	0.165	77	KDHHHHDDDY	0.163	130	AAKHHHHAAKY	0.205
31	GAGHHPHHGAY	0.158	78	DDHHHHDDDY	0.165	131	AAKHHHHKAAAY	0.205
32	GAGHHEHHGAY	0.088	79	FFLHHHHHEEY	0.168	132	DKHHHHKDKY	0.205
33	GAGHDHGHGAY	0.088	80	KKKHHHHHEEY	0.168	133	AAKHHHHAKAY	0.206
34	GAGHWHHHGAY	0.203	81	KSKHHHHHEEY	0.168	134	ARAHHHHAAEY	0.208
35	GAGHHKHHGAY	0.173	82	REEHHHHHEERY	0.168	135	DDRHHHHRDDY	0.208
36	GAGHHMHHGAY	0.163	83	DDDHHHHDDDY	0.168	136	KDHHHHDKKY	0.208
37	GAGHHNHHGAY	0.168	84	AAHHHHAAAY	0.168	137	RRRHHHHHEEY	0.210
38	GAGHHQHHGAY	0.160	85	KKHHHHDDDY	0.170	138	AKAHHHHAAKY	0.212
39	GANNHAHHNAY	0.155	86	KKHHHHHEEY	0.175	139	KKHHHHDKKY	0.218
40	GAQHHAHHQAY	0.143	87	KAHHHHHAAEY	0.175	140	DKKHHHHKDKY	0.218
41	GAGHHHHHGAY	0.158	88	AAKHHHHAAAY	0.175	141	KSKHHHHKSKY	0.225
42	GAGHHLHHGAY	0.155	89	AAKHHHHADAY	0.175	142	KKHHHHKEKY	0.225
43	GAGHHPHHGAY	0.170	90	AAKHHHHDAAY	0.175	143	EKKHHHHKKEY	0.228
44	GAGHRRHHGAY	0.180	91	AAAHHHHAAAY	0.178	144	AARHHHHHAARY	0.242
			92	AAKHHHHHEAAY	0.178	145	RAAHHHHHAARY	0.244
4	GAGAHHHHGAY	0.190	93	RDDHHHHDDDY	0.178	146	AARHHHHHARAY	0.244
27	GAHHGHHAGAY	0.155	94	PPPHHHHPPPY	0.180	147	AARHHHHRAAY	0.244
45	GAHHGAAHHGAY	0.150	95	FFLHHHHKSKY	0.180	148	ARAHHHHHAARY	0.249
						149	RERHHHHRERY	0.253
						150	RRRHHHHDDRY	0.253

**Table I. (continued) Structure of Model Peptides and Their Retention (Eluting Imidazole Concentration) on Nickel Column**

#	Peptide	Im concentration (M)	#	Peptide	Im concentration (M)	#	Peptide	Im concentration (M)
151	RRRHHHHRDDY	0.253	183	GAGARRRRRY	0.070	213	RRGASGASGASGHHHHY	0.200
152	RRRHHHHRDRY	0.255	184	GAGARRRRRY	0.085	214	EEGASGASGASGHHHHY	0.175
153	RREHHHHERRY	0.260	185	GAGAWWGAGAY	0.120	215	RREESGASGASGHHHHY	0.183
154	FFLHHHHSRY	0.263	186	GAGAWWGAGAY	0.000			
155	RRRHHHHEERY	0.265	187	GAGAHWGAGAY	0.095	216	PRREEGGRWGY	0.080
156	KKKHHHKKKY	0.267	188	GAGAHWWGAGY	0.138	217	PRREEGRWGY	0.080
157	FFLHHHHLFFY	0.270	189	GAGHHWGAGAY	0.135	218	PGDYDDDRRQY	0.000
158	RRRHHHHEREY	0.270	190	GAGHHWWGAGY	0.213	219	PGDYDDKRRQY	0.000
159	RRRHHHHEEY	0.273	191	GAGARWGAGAY	0.085	220	QPRKIRPEGRY	0.000
160	RDRHHHHRDRY	0.280	192	GAGARWWGAGY	0.135	221	QADKGEPEGRY	0.000
161	RRDHHHHRDRY	0.283	193	GAGRRWGAGAY	0.098	222	QPRMIRPEGRY	0.000
162	DRRHHHHRDRY	0.288	194	GARRWRRGAY	0.223	223	FNAEFNEIRRY	0.000
163	RSRHHHHSRY	0.295				224	GNAEPNEIRRY	0.000
164	ERRHHHHRREY	0.313	195	HHASGASGASGHHHY	0.140	225	NNFGKLFVQY	0.000
165	RRRHHHHRDRY	0.315	196	HASGASGASGHHHY	0.172	226	NNFGKDKEVKY	0.000
166	RRRHHHERRY	0.340	197	ASGASGASGHHHHY	0.197	227	EQKLISEEDLY	0.000
167	RRRHHHKKKY	0.368	198	ASGASGASGHHGHY	0.143	228	YPYDVPDYAY	0.000
168	RRRHHHRRRY	0.426	199	ASGASGASGHHGHY	0.133	229	LEHDCY	0.000
			200	ASGASGASGHHGHY	0.135	230	LEHDGGY	0.000
169	HRAGAGAGAY	0.108	201	ASGASGASHGASGHHY	0.133	231	ALEHDGGY	0.000
170	HRHGAGAGAY	0.148	202	ASGASGASHGASGHHY	0.128	232	LLEHDGGY	0.000
171	HRHRAGAGAY	0.168	203	ASGASGHHASGASGHHY	0.130	233	KLEHDGGY	0.000
172	HRHRHGAGAY	0.220	204	ASGASHHGASGASGHHY	0.130	234	ELEHDGGY	0.000
173	HRHRHRAGAY	0.243	205	ASGASHHGASGASGHHY	0.130	235	SLEHDGGY	0.000
174	HGAGAGAGAY	0.100	206	ASGHHASGASGASGHHY	0.133	236	PLEHDGGY	0.000
175	HHGAGAGARY	0.133	207	ASHHGASGASGASGHHY	0.130	237	HRIFLAGDKDY	0.098
176	HHAGAGARRY	0.140	208	AHHSASGASGASGHHY	0.130	238	HRIFLAGDEDY	0.093
177	HHAGAGARRY	0.190	209	HAHSHGHASGASGASGY	0.173	239	KRKGDEVDGVY	0.000
178	HHHAGARRRY	0.210	210	HASHGASHGASGASGY	0.150	240	RKGEVDGVQDY	0.000
179	AGAGAGAGARY	0.000	211	HASGHASGHASGHASGY	0.148	241	KGDEVDGVDEY	0.000
180	AGAGAGARRRY	0.000	212	HASGAHSGASHGASGHY	0.148	242	GDEVDGVDEY	0.000
181	GAGAGARRRY	0.000	197	ASGASGASGHHHHY	0.197	243	DEVDGVDEY	0.000
182	GAGAGARRRY	0.000						

At the end of the synthesis, the plate was dried in vacuo and 150  $\mu$ L of mixture K (9) (trifluoroacetic acid–thioanisole–water–phenol–ethanedithiol, 82.5:5:5:5:2.5, v/v) was added. The plate was capped and shaken on the plate shaker for 3 h. The suspension was transferred by multi channel pipettor to a filter plate (Orochem Technologies, Lombard, IL). The filtrate was collected in a deep well plate (VWR) and precipitated with ether (600  $\mu$ L). After standing in a refrigerator for 2 h, a pellet was formed by centrifugation. The supernatant was removed by a surface suction device and the pellet was resuspended in ether (600  $\mu$ L) and centrifuged again. The process of supernatant removal and resuspension was repeated three times. The product was dried in a Speedvac (ThermoSavant, Waltham, MA), dissolved in 200  $\mu$ L of H<sub>2</sub>O or 50% dimethylsulfoxide (DMSO)–50% H<sub>2</sub>O, and samples of 20  $\mu$ L were taken into 180  $\mu$ L of water. Twenty microliters were injected onto high-performance liquid chromatography (HPLC) column (Waters, Milford MA,  $\mu$ Bondapak, C18, 10  $\mu$  particle, 125 Å pore, 3.9  $\times$  150 mm, gradient 0.05% TFA in H<sub>2</sub>O to 70% acetonitrile, 0.05% TFA in 15 min, flow rate, 1.5 mL/min, detection by UV at 217 nm). Retention on the reversed-phase column was measured for all studied peptides. No significant

deviation from the predicted retention based on the amino acid composition of the peptide have been observed, suggesting that there is no steric or conformational effect on the retention. MS was performed at HT-Labs (San Diego, CA).

The retention of peptides using IMAC was evaluated using an HPLC equipped with a 1-mL volume HisTrap column (Amersham Biosciences) with the detection at 260 nm. The peptides were injected in 0.02M sodium phosphate buffer pH 7.4 containing 0.5M NaCl. The concentration of imidazole was increased linearly from 0 to 0.5M during 20 min.

## Results

Synthesized model peptides and the concentration of imidazole needed for their elution (extrapolated from the retention time in gradient elution) are given in Table I. All sequences were synthesized on tyrosine-modified resin to simplify UV detection (280 nm) of peptides eluted with increasing imidazole concentrations.

The first issue to be addressed was the dependency of the

retention on the content of histidines in the sequence. Figure 1 illustrates that the concentration of imidazole needed for elution of polyhistidine peptides depends linearly on the number of consecutive histidines (peptides 1 to 18). There is a notable difference in the elution of peptides containing amino terminal histidine versus peptides with all histidines in nonterminal positions. A free amino group on the *N*-terminal histidine residue reduces the affinity of these peptides for the nickel column. In longer sequences, this effect may be equivalent to having one fewer histidine in a nonterminal position.

Next, the arrangement of histidines within a series of undecapeptides were studied. As can be seen (Table I), separation of two histidines by one amino acid residue improves the retention slightly (peptides 2 and 19); separation by more than one amino acid residue does not have a significant effect (peptides 20–22).

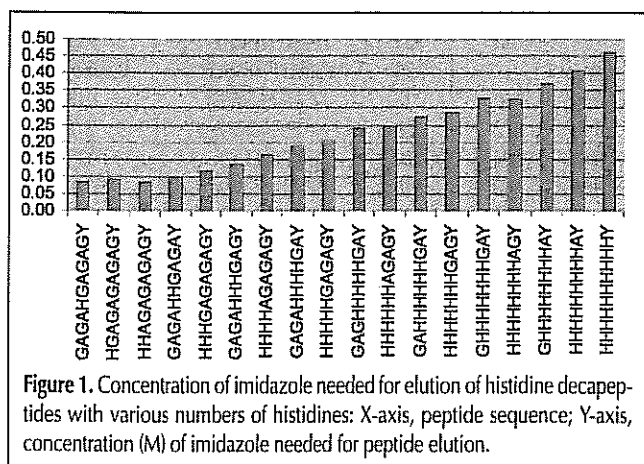


Figure 1. Concentration of imidazole needed for elution of histidine decapeptides with various numbers of histidines: X-axis, peptide sequence; Y-axis, concentration (M) of imidazole needed for peptide elution.

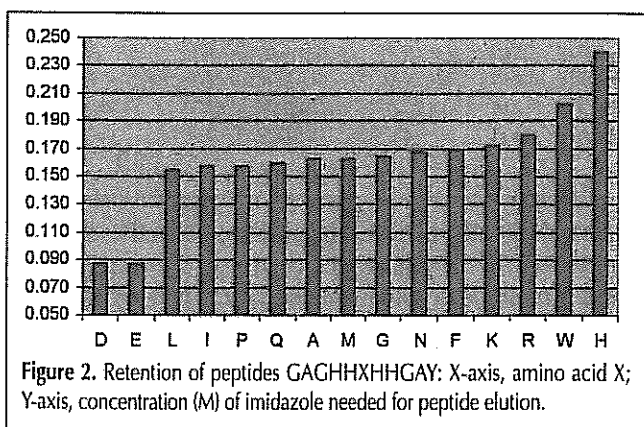


Figure 2. Retention of peptides GAGHHXHHGAY: X-axis, amino acid X; Y-axis, concentration (M) of imidazole needed for peptide elution.

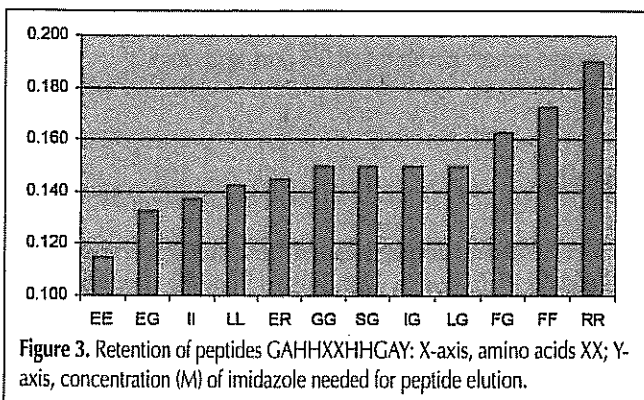


Figure 3. Retention of peptides GAHHXXHHGAY: X-axis, amino acids XX; Y-axis, concentration (M) of imidazole needed for peptide elution.

Separation of two and one histidine residue by one amino acid residue decreases the retention; however, separation by two to four amino acid residues does not decrease the retention further (peptides 3, 23–26). Separation of four histidine residues into two doublets decreases the retention; the number of intercalating residues does not make a significant difference (peptides 4, 27, and 45–47). The separation of two histidine doublets in the frame of a 17-mer was also studied, and it was found that the separation by two amino acids has the same effect as separation by 12 residues (peptides 197–208). Four histidine residues distributed regularly throughout the 17-mer (separated by two, three and four amino acid residues) have approximately the same retention, though a peptide with four histidines separated by one amino acid residue is slightly more retained (peptides 209–212). This result is easily explainable by the better availability of imidazole side chains for interaction with metal ions when histidine residues are separated by one amino acid residue. The preferred *trans* conformation of the amide bond brings amino acid side chains in 1–3 positions closer than side chains in the 1 and 2 positions.

A series of peptides with six histidine residues separated into two triplets by one to four amino acid residues do not have significant differences in their retention, although any separation slightly increases retention compared with the peptide with six adjacent histidines (peptides 16 and 65–69). This result was the most promising for our idea of combining two fragments, each containing a histidine triplet. This result permitted the use a variety of linking chemistries without concern for the spacing between the two triplets. As long as the initial components are released from the nickel column at a concentration of imidazole < 0.14M (peptide 3 or 13), and the conjugate is eluted by an imidazole concentration > 0.25M, the purification scheme might be very simple: (i) introduce the conjugation mixture to the column in a solution containing 0.14M imidazole; (ii) wash out the unreacted starting materials; (iii) and elute the purified His6 product with a solution containing 0.3M imidazole. To be able to apply this concept successfully, elution of the conjugation product should be relatively insensitive to the composition of the components surrounding the His6 tag. In the case of peptide conjugation, this requires relative insensitivity to the surrounding amino acids.

Consequently, over a hundred peptide sequences were designed and synthesized to determine the effect of amino acids surrounding the polyhistidine tag on its affinity to the nickel column. The effect of one or two amino acids joining two histidine doublets was evaluated in detail. Figure 2 shows the effect of an amino acid in the sequence GAGHHXHHGAY. The negative effect (decreased retention) of aspartic and glutamic acids (D and E) is very significant, and there is a slight positive effect (increased retention) of basic residues lysine (K) and especially arginine (R). A tryptophan (W) residue increased retention by approximately 50% of the increase achieved by addition of another histidine. This is in agreement with the effect of tryptophan observed in IMAC studies of various proteins (3,4).

Figure 3 shows the effect of a two amino acid linkage in the sequence GAHHXXHHGAY (peptides 53 to 64). The influence of glutamic acid and arginine is confirmed, and phenylalanine (F) appears to contribute to increased retention. It is more the effect of the aromaticity of the phenylalanine moiety than just the

hydrophobicity of the dipeptide, that is two leucines (L) or two isoleucines (I) do not increase retention significantly. Two isoleucines actually decrease the retention, probably due to steric factors (sequence HHHHH is significantly less flexible than HHGGHH and/or HHIGHH). The combination of arginine and glutamic acid residues was studied with sequences 48 to 52. Compensation of the effects of basic and acidic residues was independent of the relative proximity of these residues.

The effect of surrounding a polyhistidine sequence with a negative, positive, hydrophilic, and hydrophobic residue was studied by constructing an additional 98 peptides. Sequences of the form (X)XXHHHHXXXY (peptides 70 to 168) are sorted in Table I by increasing retention. Obviously, charged amino acids have the largest impact on the retention of the model peptides. Arginine and lysine residues increase retention, and glutamic acid and aspartic acid residues decrease retention. The effect of arginine compares with that of two lysine residues, and the effect of glutamic acid is slightly stronger than the effect of aspartic acid. The opposing effect of basic and acidic residues is illustrated by the finding that retention of RSRHHHHESE is the same as GAGAH-HHHGA. Hydrophobic residues increase retention significantly: FFLHHHHLFF has almost the same retention as GAHHHH-HHGA, although it has two fewer histidine residues. The 17-mer peptides 197 and 213–215 reveal an interesting effect of placing arginine and/or glutamic acid residues at a long distance from the polyhistidine cluster. Surprisingly, the effect of arginines is diminished significantly, becoming almost negligible, though the effect of glutamic acid still persists.

The combination of histidine and arginine residues was studied with peptides 169–178. When the retention of these peptides was compared with the retention of peptides with histidine alone (peptides 11–16), it was found that HR is retained better than HH, and that HRH is more effective than HHH. In addition, HHHH is very similar to HRHR, and HHHHHH and HRHRHR are retained identically. The pentamer sequence HRHRH has slightly more affinity than HHHHH. Peptides 174–178 show that histidines and arginines work cooperatively in the frame of a decapeptide as well. With peptides 179–184, the effect of pol-yarginine sequences and their affinity for the nickel column was studied. As can be seen, retention is observed only at the level of pentaarginine, showing that retention of the HR-containing peptides is caused by a cooperative effect of histidine and arginine.

Peptides 185–194 were designed to study the effect of tryptophan (W) on retention to the nickel column. A single W in the sequence did not facilitate retention; however a doublet of tryptophans is equivalent in its effect to a triplet of histidines. The HHH sequence is retained almost identically to sequences HWW and HHW. Furthermore, RW is retained as well as RRW, and RWW is equivalent in retention to the sequence HHH. Finally, RRWRR has nearly the affinity of five histidines in a row. This result may be useful in the genetic engineering of proteins, where tags alternative to pentahistidine may be desirable.

Using our high-throughput peptide synthesizer, several thousand peptides were recently synthesized for use in our protease and protein kinase assays. We tested some of these (peptides 216–243), each containing a terminal tyrosine, for their retention on the nickel column (Table I). Only peptides containing RW

or HR showed any affinity. Retention of peptides with one histidine residue in proximity of acidic residue (D or E) was completely eliminated.

## Conclusion

An exhaustive study of the affinity of histidine-rich peptide sequences for a nickel Sepharose column was performed, using elution by a gradient of increasing imidazole concentration as a measure of affinity. Retention of histidine-containing peptides depends on the arrangement of histidines within the sequence, with consecutive histidines not necessarily the best arrangement, and on the type of amino acid connecting histidine clusters. Affinity also strongly depends on the amino acid composition of neighboring sequences. Tryptophan and arginine, and to a lesser extent lysine and phenylalanine, increase affinity, while affinity is decreased by glutamic and aspartic acids. The findings indicate that it is possible to construct an affinity tag (e.g., RRWRR) that performs with the same efficiency as pentahistidine, but does not contain any histidines.

## Acknowledgments

The project was partially supported by NIH SBIR grant R44 AI056869-02.

## References

1. J. Porath, J. Carlsson, I. Olsson, and G. Belfrage. Metal chelate affinity chromatography, a new approach to protein fractionation. *Nature* **258**: 598–99 (1975).
2. E. Hochuli, H. Dobeli, and A. Schacher. New metal chelate adsorbent selective for proteins and peptid containing neighboring histidine residues. *J. Chromatogr.* **411**: 177–84 (1987).
3. E. Sulkowski. Purification of proteins *Trends Biotechnol.* **3**: 1–7 (1985).
4. M.C. Smith, T.C. Furman, T.D. Ingolia, and C. Pidgeon. Chelating peptide-immobilized metal-ion affinity-chromatography for recombinant proteins. *J. Biol. Chem.* **263**: 7211–15 (1988).
5. E. Hochuli, W. Bannwarth, H. Dobeli, R. Gentz, and D. Stuber. Genetic approach to facilitate purification of recombinant proteins with a novel metal chelate adsorbent. *Biotechnol.* **6**: 1321–25 (1988).
6. M. Lebl. Centrifugation based automated synthesis technologies. *J. Assoc. Lab. Automation* **8** 30–36 (2003).
7. M. Lebl. New technique for high-throughput synthesis *Bioorgan. Med. Chem. Lett.* **9**: 1305–10 (1999).
8. J.P. Hachmann and M. Lebl. Alternative to piperidine in Fmoc solid phase synthesis. *J. Combinatorial Chem.* **8**: 147 (2006).
9. D.S. King, C.G. Fields, and G.B. Fields. A cleavage method which minimizes side reactions following Fmoc solid phase peptide synthesis. *Internat. J. Peptide Protein Res.* **36**: 255–66 (1990).

Manuscript received March 10, 2006;  
revision received July 20, 2006.