



Article Deciphering Interactions Involved in Immobilized Metal Ion Affinity Chromatography and Surface Plasmon Resonance for Validating the Analogy between Both Technologies

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Abstract: Various peptides can be obtained through protein enzymatic hydrolysis. Immobilized metal ion affinity chromatography (IMAC) is one of the methods which can be used to separate metal chelating peptides (MCPs) in a hydrolysate mixture. In this context, this work aims to understand deeply the interactions in IMAC and surface plasmon resonance (SPR) in order to validate experimentally the analogy between both technologies and to be further able to perform IMAC modeling in the next work using peptide sorption isotherm parameters obtained from SPR. Indeed, chromatography modeling can be used to predict separation of MCPs in IMAC and the knowledge of peptide sorption isotherm obtained from SPR is a crucial step. For this purpose, 22 peptides were selected and investigated in IMAC using HisTrap X-Ni²⁺ and HiFliQ NTA-Ni²⁺ columns and were also studied in SPR as well. Results showed that peptides with histidine residues had good affinity to Ni²⁺, while the high positive charge of peptides was responsible of ionic interactions. Further, most of the peptides with good retention time in IMAC showed a good affinity in SPR as well, which validated experimentally the SPR-IMAC analogy.

Keywords: IMAC; SPR; metal chelating peptides; metal complexation; chemical interactions; separation; sorption isotherm

1. Introduction

Bioactive peptides naturally encrypted within protein sequences can be produced by enzymatic hydrolysis of animal or plant proteins. Proteolysis is widely used to produce hydrolysates. These complex mixtures of peptides present various biofunctional and biological activities, such as antimicrobial, antioxidant, anticarcinogenic, antitumoral, antidiabetic and mineral absorption activities [1]. Among the biofunctional peptides, the so-called metal chelating peptides (MCPs) are endowed with the ability to complex metal ions involving some metal-coordination bonds, which confers them various potential applications. Indeed, according to the metal ion complexed, MCPs can be used in various domains, notably in health to transport metals like iron, calcium and zinc in the body [1], in cosmetics and in food systems to avoid lipid oxidation [2]. The MCPs are produced from various protein resources such as soybean [3,4], sesame [5], whey [6] and tilapia and its co-products [7].

Despite their huge potentials, the recovery of MCPs in hydrolysates is challenging due to their low concentration in these complex mixtures of peptides, constituted from 10 to 100 various peptide sequences. Separation techniques such as IMAC are often used to



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Copyright: © 2024 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https:// creativecommons.org/licenses/by/ 4.0/). recover biofunctional peptides in hydrolysates [1]. Over the past decades, IMAC has been used to purify and separate proteins (e.g., recombinant proteins) and polypeptides based on the interaction between proteins and immobilized metal ions like Ni²⁺, Cu²⁺, Co²⁺ [8,9]. For recombinant protein separation, histidine tag is commonly added to their C- or N-terminus part in order to increase their affinity to metal ion immobilized onto the chromatographic phase, thus improving their yield of purification. More recently, IMAC has been developed to separate MCPs from protein hydrolysates based on the differences in peptide affinity to a given immobilized metal ion, such as $Fe^{3+}[3,4,10]$, $Ca^{2+}[4]$, $Zn^{2+}[5,6]$ or $Cu^{2+}[6]$. Peptide-metal ion affinity depends on the peptide size, number, position and nature of the amino-acid (AA) residues present in a peptide sequence and can be predicted based on the theory of hard and soft acid and base (HSAB). According to this HSAB theory [9], metals behave like Lewis acids, while peptides behave like Lewis bases, both classified in three categories: hard, borderline and soft. This theory helps to predict which metal ion can form the most stable complex with a peptide, based on the type of residues present in the sequence. Therefore, AA residues having carboxylate groups (glutamic acid and aspartic acid) are more likely to chelate metals such as Fe³⁺ since both are classified in the hard acid-base groups; sulfur (thiolate or thioether) groups (cysteine, methionine) are more prone to chelate metals like Cu⁺, both classified in soft acid-base groups, while imidazole and indole groups (histidine, tryptophane) are more likely to chelate Cu²⁺, Ni²⁺, Co²⁺, or Zn^{2+} , which belong to the borderline acid-base group. In this study, we are interested in peptides that can chelate Ni²⁺ (Ni²⁺ being a borderline Lewis acid) and thus contain His and Trp residues according to the HSAB theory. Nevertheless, it should be noted that the complexation of metals by the indole ring of Trp is rarely observed in proteins [11].

In the meantime, other methods based on a peptide/protein-metal ion interactions as in IMAC have been developed to screen MCPs in hydrolysates, among them the phage surface display [12], the electrically switchable nanolever technology [13] and surface plasmon resonance [14–16]. SPR is an optical biosensor which enables to quantify the interaction between a peptide/protein and an immobilized metal ion [17]. As main advantages, this technology uses a low quantity of sample and buffer and enables a quick investigation of the presence of MCPs in hydrolysates, which is a crucial step before launching time-consuming separation.

Due to the advantages and efficacy of SPR stated above, our team recently initiated an approach to predict peptides concentration profile at the outlet of the column during their separation in IMAC [15,18]. In this approach, IMAC chromatography was modeled using a transport-dispersive model and peptide binding affinity constants (K_A). These peptide binding affinity constants (K_A) were obtained from SPR sorption isotherms and used as model input data considering the theoretical SPR-IMAC analogy. Indeed, SPR and IMAC are notably based on peptide-metal ion interactions and present various similarities, especially when they use the same metal ion (e.g., Ni²⁺, Fe³⁺, Cu²⁺) immobilized on the same complexing agent (e.g., nitrilotriacetic acid (NTA)) fixed onto a support and the same loading buffer and elution buffer as well [15]. However, IMAC is used for MCP separation, while SPR can be used for MCP screening only.

Therefore, this work aims to decipher the phenomenon of peptide-metal ion complexation occurring inside an IMAC chromatographic column with a deeper investigation using SPR (a technology also based on peptide-metal ion interaction) in order to be further able to predict peptide separation in IMAC. For this purpose, synthetic peptides—potentially Ni²⁺ chelating peptides present in pea proteins—were investigated for their affinity for Ni²⁺ in IMAC and SPR, respectively, with the better understanding of interactions helping to experimentally validate the analogy between SPR and IMAC.

2. Materials and Methods

2.1. Synthetic Peptides Investigated and Other Chemicals

The peptide sequences investigated in IMAC and SPR were obtained from in silico proteolysis of pea (*Pisum Sativum*) proteins. Pea proteins were chosen for their absence of

allergens compared to other plant proteins like soy, according to the Official Journal of the European Union [19]. Major pea proteins, like legumin, vicilin, albumin and convicilin, were identified in the literature [20–22] and their sequences were obtained from UniProtKB (protein sequences presented in Supplementary Data S1).

From these pea protein sequences, peptides were obtained by in silico enzymatic proteolysis, and potential Ni²⁺ chelating peptides were identified based on HSAB theory; they presented at least one histidine or tryptophan in their sequences, which might lead to Ni²⁺ complexation [17]. In addition, they were selected as small-sized Ni²⁺-chelating peptides (MCPs, 2–10 amino-acid residues), less subject to further in vivo hydrolysis that could reduce their activities [23]. In the end, 16 peptides theoretically endowed with various Ni²⁺ metal-chelation capacities and 4 peptides with potentially no affinity for Ni²⁺ (without histidine/tryptophan, negative control) were selected and synthetized by Genecust (Dudelange, Luxemburg). In addition, 2 peptides, GHHHHHHHHHY (GH8AY) and HHHHHH, were used as positive controls since HHHHHH is a good Ni²⁺ chelating peptide commonly used as a tag for recombinant protein purification [24] and GH8AY presents a high number of histidine and shows a good affinity for Ni²⁺ [25]. The synthetized peptides, the protein from which they are derived and their physico-chemical properties (molecular weight, MW; isoelectric point, IP; and percentage of global charge at pH 7.4) are summarized in Table 1. In addition, their chemical structures are illustrated in Figure 1. Pea protein names and sequences were obtained from UniprotKB, and the molecular weight and isoelectrical point were calculated from PepCalc free software (https://pepcalc.com).

The percentage of global charge at pH 7.4 was determined by ACD/Percepta Software version 14.0 (ACD/Labs) (https://www.acdlabs.com/), which can calculate the acid dissociation constants (pKa) of all ionizable groups in the sequence under standard conditions. The predictions of pKa values are based on a database of more than 20,000 compounds. The plot of percentage of net charge as a function of pH can be generated based on the pKa values. From these data, at pH 7.4, all percentages of positive, negative and zwitter-ionic charges (i.e., considered as globally neutral) were summed up (%) to obtain the percentage of global positive charges and global negative charges of the peptides and are presented in Table 1.

Concerning other chemicals, the following were purchased from Sigma Aldrich (St Louis, MO, USA): potassium phosphate monobasic (KH₂PO₄), nickel (II) sulfate (NiSO₄.X H₂O), imidazole bio–Ultra GC > 99.5%, hydrochloric acid 37% (HCl), ethylenediamine tetra acetic acid disodium salt dihydrate (EDTA-Na₂·2H₂O). Others, namely sodium hydroxide (NaOH), sodium chloride (NaCl) and di-sodium hydrogen phosphate dodecahydrate (Na₂HPO₄·12H₂O), were purchased from VWR Chemicals (Leuven, Belgium). Ultrapure water (18.2 MΩ/cm) was obtained using a Milli-Q System (Millipore, Bay City, MI, USA). Specifically, for SPR experiments, 500 µM NiCl₂ Biacore kit solution, 50 µM EDTA Biacore kit solution and BIA-maintenance kit solution were provided by GE Healthcare (Uppsala, Sweden). Sodium dodecyl sulfate (SDS) and Tween[®] 20 were provided by Sigma Aldrich (St Louis, MO, USA).

2.2. *Immobilized Metal Ion Affinity Chromatography* 2.2.1. HPLC Equipment and IMAC Columns Used

IMAC experiments were carried out using an HPLC system (LC-20AD, Shimadzu, Japan) at a flow rate of 1 mL/min, an oven temperature set at 25 °C and a Diode Array Detector (SPD-M20A, Shimadzu, Japan). Three wavelengths were selected: 214 nm and 232 nm (i.e., peptide bonds) and 280 nm (aromatic ring) [26]. For each observed peak, fractions were collected using a fraction collector (FRC-10A, Shimadzu, Japan) and analyzed by LC-MS (Section 2.3) in order to confirm the peptide presence and the absence of impurities.



Figure 1. Chemical structures of some synthetic peptides investigated by IMAC and SPR, with histidine and tryptophan residues, functional groups being highlighted with red circle. Peptides in panels (**A**–**E**) have more than two His or Trp residues that confer them the ability to chelate Ni²⁺; peptides in panels (**F**,**G**) have one His that confers low ability to chelate Ni²⁺, while peptides in panels (**H**,**I**) do not have His or Trp.

Table 1. Synthetic peptides investigated and selected from pea protein sequences with their theoretical physico-chemical properties. Pea protein names and sequences were obtained from UniprotKB; the molecular weight and isoelectrical point were calculated from PepCalc software, while the percentage of global charges at pH 7.4 was calculated using Percepta software. n.c.: not concerned.

Unip	rotKB			Pe	epCalc	Percepta
Protein Code	Protein Name	Peptide Sequence	n Residues	MW (g/mol)	Iso-Electric Point (PI)	% of Global Charge at pH 7.4
n.c	n.c	GHHHHHHHHAY	11	1406.43	8.21	80% +, 3% -
n.c	n.c	НННННН	6	840.87	7.97	70% +, 6% -
P05692	Leguminin J	GRHRQKHS	8	1005.12	12.13	100% +
O24294_PEA	Legumin (Minor small)	QRHRK	5	723.9	12.13	100% +
P13915	Convicilin	KRHGEWRPS	9	1152.27	11.22	100% +
P02857	Leguminin A	HGSLHKNA	8	862.95	9.88	65% +, 0% -
P02857	Leguminin A	KGKSR	5	574.67	11.57	100% +
P02857	Leguminin A	HGSLH	5	549.59	7.72	26% +, 25% -
P13915	Convicilin	RHGEWRPS	8	1024.09	10.39	44% +, 0% -
P02857	Leguminin A	AEHGSLH	7	749.77	6.05	11% +, 48% -
P05692	Leguminin J	AP HW NIN	7	850.92	7.88	20% +, 21% -
P05692	Leguminin J	GLH	3	325.36	7.81	34% +, 13% -
P05692	Leguminin J	GLHLPS	6	622.71	7.80	19% +, 21% -
P05692	legumin J	YPVGR	5	590.67	9.57	52% +, 0% -
O24294_PEA	Legumin (Minor small)	KERESH	6	784.82	7.60	24% +, 27% -
P13918	vicilin	FPGSA	5	477.51	3.38	0%-, 56% -

Unip	rotKB			PepCalc		Percepta	
Protein Code	Protein Name	Peptide Sequence	n Residues	MW (g/mol)	Iso-Electric Point (PI)	% of Global Charge at pH 7.4	
P62926	Albumin-1 A	LTKNV	5	573.68	10.12	70% +, 0% -	
O24294_PEA	Legumin (Minor small)	VSHVN	5	554.60	7.78	18% +, 25% -	
Q7M1N3_PEA	Legumin L1 beta chain	LLH	3	381.47	7.81	34% +, 13% -	
O24294_PEA	Legumin (Minor small)	LAHS	4	426.47	7.81	22% +, 19% -	
P13918	Vicilin	QSHF	4	517.53	7.52	15%+, 34% -	
P13918	vicilin	KETQH	5	641.67	7.60	24% +, 26% -	

Table 1. Cont.

The 22 peptides were studied in IMAC and experiments were carried out with two different columns in order to understand various interactions well: HiFliQ NTA-Ni²⁺ (Generon, Slough, UK; NTA as complexing agent) and HisTrap[™]FF crude (GE Healthcare Bio-Sciences, Sweden; unknown complexing agent). HisTrap X-Ni²⁺ and HiFliQ NTA-Ni²⁺ nomenclature was used for columns loaded with Ni²⁺, which were used to investigate the peptide–metal ion interactions. In addition, all of the 22 peptides were also analyzed on two columns without Ni²⁺ as control columns and HisTrap X and HiFliQ NTA nomenclature was used in this latter case. These control experiments were performed in order to determine whether there were interactions between the peptide and other parts of the column like the matrix or the complexing agent. For the majority of peptides, these interactions were not observed or were very weak since peptides were directly eluted from the columns. Different experiments were required to investigate these former interactions in IMAC, while in SPR these former interactions and direct interactions between the peptide and Ni²⁺ could be obtained within one experiment.

Regarding other columns specifications, both columns had a volume of 1 mL, their binding capacity was 40 mg or 50–75 mg of 6xHis-tagged protein for HiFliQ NTA-Ni²⁺ and HisTrap X-Ni²⁺, respectively (Supplementary Data S2 for other specifications).

The HiFliQ NTA-Ni²⁺ column was initially selected since the metal ion complexing agent was known (NTA) and identical to the one used on the SPR sensor chip, enabling a real analogy between IMAC and SPR. Yet, due to the absence of retention on HiFliQ NTA-Ni²⁺ of some peptides, a second IMAC column, the HisTrap X-Ni²⁺, was investigated as well. Yet, in this latter case, the metal ion complexing agent (X) and its denticity were not communicated by the manufacturer.

2.2.2. IMAC Cycle and Elution Program

An IMAC cycle is composed of 4 main steps: equilibration, injection + unbound component removal, elution and washing. For each peptide investigated, the IMAC column was first equilibrated using the PBS 1X pH 7.4 buffer (Na₂HPO₄.12H₂O 6.7 mM; KH₂PO₄ 1.25 mM; NaCl 15 mM) (equilibration step). Fifty μ L of peptide solutions prepared at 20 mM in PBS 1X buffer was loaded (injection), and then the PBS 1X buffer was injected for 5 min to remove unbound components. For eluting peptides that have affinity for Ni²⁺, a gradient of imidazole was used, from 0 to 600 mM imidazole (prepared in PBS 1X pH 7.4) for 60 min. Finally, extra binding peptides were removed by 600 mM imidazole for 5 min (washing step) and the column was equilibrated using the PBS 1X pH 7.4 buffer for the next run (equilibration step).

Note that during the unbound component removal phase and gradient elution step, the fractions were collected and identified by mass spectrometry in order to confirm or not the presence of the peptide or an impurity in the collected fraction.

After 10 peptides assays using HisTrap X-Ni²⁺, the column was stripped with 50 mM EDTA in PBS 1X pH 7.4 buffer to remove the immobilized Ni²⁺ and regenerated with

100 mM NiSO₄ in PBS 1X pH 7.4 buffer according to the provider specifications. Meanwhile, after 5 peptide assays, HiFliQ NTA-Ni²⁺ was stripped with 100 mM EDTA pH 8 and regenerated with 10 mM NiSO₄ solution according to the provider specifications.

2.2.3. Treatment of IMAC Data

For each peptide series investigated, a blank was first studied, i.e., PBS 1X buffer injected only without sample in order to correct the deviation of the baseline in elution phase due to the slight imidazole absorbance at the wavelengths investigated (214 nm and 232 nm). Thus, a baseline correction was systematically applied to each peptide chromatogram by subtracting the blank at the wavelength of interest. Then, for each investigated peptide and for each column, a retention time (t_R , min) was determined corresponding to the maximum of the peak.

2.3. LC-MS

To confirm the presence of a peptide in fractions collected in IMAC experiments during the gradient elution step and to ensure that the peak was not related to any impurities, fractions were analyzed on a Thermo Scientific UHPLC-HRMS system composed of a VanquishTM liquid chromatography unit coupled to a photodiode array detector (PDA) and an Orbitrap ID-XTM TribridTM high-resolution mass spectrometer operating in electrospray ionization mode (ESI).

For each collected fraction, 10 µL was injected on a Thermo ScientificTM AcclaimTM 120 C18 column (100 mm \times 2.1 mm–2.2 μ m) maintained at 40 °C. The flow rate was set at 200 μ L.min⁻¹ and mobile phases consisted of water modified with trifluoroacetic acid (0.1%) for A and acetonitrile modified with trifluoroacetic acid (0.1%) for B. Peptides were eluted using a 10 min gradient from 5% to 98% of B before a 5 min column wash at 98% of B, and finally followed by a 6 min equilibration step at 5% of B. Mass analysis was carried out in ESI positive ion mode (ESI+) and mass spectrometry conditions were as follows: spray voltage was set at 3.5 kV; source gases were set at 35, 7 and 10 (in arbitrary units.min⁻¹) for sheath gas, auxiliary gas and sweep gas, respectively; vaporizer temperature and ion transfer tube temperature were both set at 300 °C. MS scans were performed from 150 to 2000 m/z at 7.5 K resolution (at 200 m/z, FWHM) with following parameters: RF-lens, 35%; maximum injection time, 50 ms; data type, profile; normalized AGC target, 25%. The mass spectrometer calibration was performed using the Thermo Scientific Pierce TM FlexMix TM calibration solution. MS data acquisition was carried out with the Xcalibur v. 3.0 software (Thermo Scientific, Waltham, MA, USA). For all 22 peptides investigated, the peak associated with the dead time of the HPLC system for IMAC was observed around 1.2 min. The non-retained peptides were detected in this peak at 1.2 min, and other peptides were detected in another peak and their t_R value was used for their classification.

2.4. Surface Plasmon Resonance

SPR experiments were carried out on a Biacore X100 instrument (GE Healthcare, Uppsala, Sweden) using an NTA sensor chip (GE Healthcare, Uppsala, Sweden) at 25 °C as described in the literature [14]. Briefly, Ni²⁺ was first loaded on the NTA sensor chip using a 500 μ M NiCl₂ Biacore kit solution. The running buffer was a PBS 1X (pH 7.4, 0.005% Tween[®] 20) prepared from PBS 10X buffer (67 mM Na₂HPO₄·2H₂O, 12.5 mM KH₂PO₄, 150 mM NaCl, pH 6.6). The flow rate was set at 20 μ L/min, the running buffer was filtered onto a 0.22 μ m membrane (low protein binding, non-pyrogenic, PALL) and two channels were available on the chip, one previously loaded with Ni²⁺ for investigating the interactions between the peptides and Ni²⁺ and another one without Ni²⁺ for investigating interactions between peptides and the matrix or complexing agent.

In order to study the binding affinity of a peptide, seven concentrations varying from 10^{-6} M to 10^{-2} M depending on the peptide were prepared in the running buffer and each concentration was loaded onto the Ni²⁺-NTA chip to evaluate the affinity of the peptide for Ni²⁺ already immobilized on NTA. The chip surface was regenerated with a 500 mM

imidazole solution prepared using ultrapure water, followed by a second regeneration with EDTA solution prepared by adding 87 μ L of EDTA solution at 50 μ M (Biacore kit solution) in 10 mL running buffer (PBS 1X). After each cycle, the SPR chip surface was washed with SDS 0.5% v/v at a flow rate of 40 μ L/min followed by an extra wash with 50 μ M EDTA and equilibrated with the running buffer before studying the next peptide concentration.

For each peptide concentration, the response (Resonance Unit, R.U.) was recorded as a function of time (s) to form a sensorgram (Figure 2A), and then the response at equilibrium was taken to build a sorption isotherm (response as a function of peptide concentration, Figure 2A). The isotherm was fitted with a 1:1 binding model using BIAevaluation software version 4.1. in order to determine the dissociation constant (K_D , M). The equation of 1:1 binding model is as follows:

$$R = \frac{R_{\max}C}{K_D + C} \tag{1}$$

where R is a response during steady state phase (R.U.), C is the peptide concentration (M) and R_{max} is the saturation response (R.U.).



Figure 2. Surface plasmon resonance: (**A**) example of sensorgram with a color line used for each investigated concentration, x meaning values at equilibrium; (**B**) sorption isotherm for a peptide (with good affinity) studied, plotted from response collected at equilibrium; (**C**) example of an isotherm for a peptide without affinity or very low affinity where saturation is not attained (peptide KETQH); (**D**) example of an isotherm where K_D could not be calculated due to the presence of high aspecific interactions compared to specific interactions (peptide AEHGSLH). For panel (**B**–**D**), the red points represent the experimental data, and the black line represents the fitted data.

2.5. Statistical Analysis

Statistical analysis was performed using Statgraphics $19^{\text{(B)}}$ Centurion (Statgraphics Technologies, Inc., The Plains, VA, USA). Mean values and standard deviations were introduced to Statgraphics; multiple sample comparisons using Bonferroni were performed to determine statistical differences between peptides retention times. The significance was evaluated statistically at p < 0.05 significance level.

3. Results and Discussion

3.1. Determination of Peptide Retention Time Using IMAC

3.1.1. Peptide Retention Time Using HisTrap X-Ni²⁺ Column

From IMAC experiments performed on 22 peptides using the HisTrap X-Ni²⁺ column, peptides can be classified in 4 groups according to their retention times (t_R): group 1 varying between 15 min and 31 min (GH8AY, HHHHHH and GRHRQKHS), group 2 varying within 5 min and 10 min (QRHRK, KRHGEWRPS, HGSLHKNA, KGKSR, HGSLH, RHGEWRPS), group 3 with retention times less than 4 min (AEHGSLH, APHWNIN, GLH, GLHLPS) and finally, group 4 with peptides not retained on the HisTrap X-Ni²⁺ column (YPVGR, KERESH, LLH, VSHVN, QSHF, LAHS, KETQH, LTKNV, FPGSA). Results are summarized in Table 2.

Table 2. Retention times of peptides investigated with HisTrap X-Ni²⁺ and HiFiQ NTA-Ni²⁺ columns and their corresponding groups. Letters (^{a-h}) indicate statistically significant differences (p < 0.05) between samples.

Peptides	HisTrap X-Ni ²⁺ Mean t_R (min)	Group	HiFliQ NTA-Ni ²⁺ Mean t_R (min)	Group
GH8AY	30.25 ± 0 g	1	22.93 ± 0 ^h	Ι
НННННН	$22.20\pm0.58~^{\rm f}$	1	17.62 ± 0.23 g	Ι
GRHRQKHS	15.79 ± 0.34 $^{ m e}$	1	8.72 ± 0.18 $^{ m e}$	Ι
QRHRK	10.01 ± 0.06 d	2	5.01 ± 0.62 ^{c,d}	Π
KRHGEWRPS	8.58 ± 0.29 c	2	5.24 ± 0.36 ^{c,d}	II
HGSLHKNA	7.71 ± 0.29 c	2	6.02 ± 0.05 $^{ m d}$	II
KGKSR	5.57 ± 0.02 $^{\mathrm{b}}$	2	not retained	III
HGSLH	$5.48\pm0.20^{\text{ b}}$	2	4.82 ± 0.24 b,c	II
RHGEWRPS	$5.41\pm0.19^{ ext{ b}}$	2	3.98 ± 0.22 ^{a,b}	II
AEHGSLH	3.93 ± 0.09 ^a	3	not retained	III
AP HW NIN	3.81 ± 0.22 a	3	3.50 ± 0.04 a	II
GLH	3.68 ± 0.46 a	3	not retained	III
GLHLPS	3.44 ± 0.67 ^a	3	not retained	III
YPVGR		4		III
KERESH		4		III
FPGSA		4		III
LTKNV		4		III
VSHVN	not retained	4	not retained	III
LLH		4		III
LAHS		4		III
QSHF		4		III
KETQH		4		III

In IMAC, the higher the retention time (t_R) of the peptide in the column is, the stronger the interaction between the peptide and the immobilized metal ion is and thus the higher the imidazole concentration required for peptide elution is. Therefore, HHHHHH used as positive control in this study is well retained in the column, with a high retention time (t_R) of 22.20 min. The good affinity of HHHHHH for Ni²⁺ is in agreement with the literature since this peptide is used in the purification of recombinant proteins as a His-tag [27] due to the presence of multi-histidine residues (Figure 1A) [28]. Concerning other peptides belonging to the same group 1 as HHHHHH, i.e., GH8AY and GRHRQKHS, these sequences also showed high retention times and thus have a good affinity for Ni^{2+} . Generally, the affinity for Ni²⁺ depends on the composition of the peptide sequences, i.e., the number of histidine or tryptophan and their position in the sequence [27,29]. Therefore, the good affinity for Ni²⁺ observed for HHHHHH, GH8AY and GRHRQKHS is due to the presence of more than one histidine residue; in addition, these histidine residues are separated with 3 other AA residues in the case of peptide GRHRQKHS (Figure 1B), which can facilitate the formation of the peptide-Ni²⁺ complex by macrochelate arrangement. Concerning HHHHHH and GH8AY, both peptides contain many histidine residues with a possibility to form a complex from two non-vicinal H, thus leading to high affinity for Ni²⁺. In addition to having at least 2 H in their sequence, group 1 peptides present a high percentage of positive charges (greater than 70%, Table 1) at pH 7.4.

In addition, group 2 is divided into two subgroups: the first one is composed by peptides QRHRK, KRHGEWRPS (Figure 1C) and HGSLHKNA (Figure 1D), while the second one is composed by KGKSR, HGSLH and RHGEWRPS. These latter peptides have retention times smaller than those of group 1, which are referred to as having a medium affinity for Ni²⁺. To explain the trend in the first subgroup, notably the t_R value of QRHRK (10.01 min, 1 H residue) being higher than that of HGSLHKNA (7.71 min, 2 H residues), the overall positive charge of the peptide QRHRK must be considered; this latter peptide is closer in amino acid composition to peptide GRHRQKHS of group 1. Indeed, the retention time of a peptide depends not only on the number of H present in the sequence but also on the global positive charge of the peptide and the global charge of the X-Ni²⁺ complex on the surface of the solid phase, which might be negative, leading to ionic interactions. Thus, QRHRK and KRHGEWRPS peptides having one histidine but also 100% positive charges at pH 7.4 (Table 1) were retained at the same level as HGSLHKNA having 2 H but a lower percentage of positive charges (65%) at the same pH. This lower percentage of overall positive charges can explain the lower retention time of HGSLHKNA ($t_R = 7.71$ min and 65% of positive charges at pH 7.4) compared to the GRHRQKHS peptide of group 1 ($t_R = 15.79$ min and 100% of positive charges at pH 7.4). To explain the trend in the second subgroup, with the exception of KGKSR (Figure 1H), peptides HGSLH and RHGEWRPS (Figure 1E) present two potential complexing AA residues: either 2H or 1H with a weak complexing residue W, and their common point is an overall positive charge less than 45%. Again, to explain the unusual behavior of KGKSR, which does not contain H, one must consider the positive charges of this peptide (100% at pH 7.4) to explain its higher retention time ($t_R = 5.57$ min) compared to other peptides without histidine (not retained, group 4, Table 2).

In group 3, peptides AEHGSLH, APHWNIN, GLH (Figure 1F) and GLHLPS (Figure 1G) showed lower retention times, allowing us to consider their lower affinity for Ni²⁺ than group 2. With the exception of AEHGSLH, these peptides present 1H or 1H with a weak complexing residue W and an overall positive charge less than 35%. Peptide AEHGSLH has the lower positive charge (11%), which explains its lower retention time despite having 2H.

Meanwhile, the higher positive charge at pH 7.4 observed for peptides of group 1, 2 or 3 is mainly due to the presence of arginine (Arg, R) and lysine (Lys, K) residues, which remain protonated on a large pH range (pKa of side chain of Arg is 12.10, and the one of Lys is 10.67). Indeed, the data demonstrate that the 100% positively charged KRHGEWRPS peptide with potentially two metal complexing AA residues (H or W) shows good retention time (8.58 min, group 2); however, this retention time is reduced by decreasing the positive charges when the lysine (K) residue is removed from this latter sequence to give the peptide RHGEWRPS (5.41 min, 44% of positive charges, group 2). The same behavior is observed for peptide HGSLHKNA (7.71 min, 65% of positive charges, group 2) and peptide HGSLH (5.48 min, 26% of positive charges, group 2, Tables 1 and 2) without KNA residues at the C-term. According to Karavelas and colleagues [30], Lys residue does not participate in forming a coordination complex between Lys-containing peptides and Ni²⁺ in solution. The lack of coordination complex formation involving Lys residues is explained by the fact that the ε -amino function of Lys is too far to form stable chelate with another donor atom of the sequence [31]. Thus, the presence of Lys residues in the peptides GRHRQKHS, QRHRK, KRHGEWRPS, HGSLHKNA and KGKSR might increase retention time rather due to ionic interactions with a negatively charged X-Ni²⁺ complex of the solid phase surface and not due to direct coordination interaction with Ni²⁺. However, the presence of Glu (E) in the sequences did not improve the t_R values in IMAC since Glu is a hard base and Ni²⁺ is a borderline acid, and thus they do not form a strong coordination interaction. In addition, Glu decreased the overall positive charge of the peptide AEHGSLH (t_R = 3.93 min, 11% of

positive charges) versus peptide HGSLH ($t_R = 5.48 \text{ min}, 26\%$ of positive charges) due to the low pKa value (4.25) of COOH group on the side chain of Glu.

Peptides in group 4 are not retained at all in the HisTrap column despite the presence of His residues in the sequences for some of them (i.e., KERESH, VSHVN, LLH, LAHS, KETQH, QSHF). This can be explained by the low percentage of positive charges at pH 7.4 (<20–30%, Table 1) for peptides with only one His in their sequence and by the insufficient positive charges (<100%) for the peptides without H or W in their sequence (YPVGR, FPGSA, LTKNV, Tables 1 and 2), in agreement with the literature [28,32]. One H residue in the sequence is not enough when imidazole is used as elution buffer, meaning that imidazole, commonly used as competitive agent, is suitable to investigate only multi-Hiscontaining peptides with higher affinities for Ni²⁺ [28,33]. Therefore, the peptides of group 4 have very low affinity for Ni²⁺ and a very low imidazole concentration should be used to elute them.

In addition, as reported by Sun and co-workers [29], other parameters can affect peptide–metal interactions, such as pH, ionic strength and buffer composition, which may be involved in the separation. In the study presented here, the pH was set at 7.4 and ionic interactions were reduced using 15 mM NaCl in the loading and elution buffers. This NaCl concentration, initially selected to be in the same condition as SPR experiments, may not be enough to prevent ionic interactions and should be further optimized. Indeed, the concentration of NaCl between 300 mM and 1000 mM is generally used in IMAC in order to minimize ionic interactions [9,34]. It has also been reported that the increase in NaCl improves the yield of the elution of peptides in a complex mixture of peptides [33]. Ren and colleagues studied the separation of 14 peptides in mixture and only 10 peptides were eluted with a buffer containing 100 mM NaCl, while 13 of 14 peptides were eluted with a 500 mM NaCl-containing buffer.

Taken altogether, in the analytical conditions implemented for IMAC with the HisTrap X-Ni²⁺ column, the retention time is mainly influenced by the number of histidine and the overall positive charge of the peptide at pH 7.4. In the defined groups from 1 to 4 (vide supra), the $t_{\rm R}$ decreases when the number of histidine and/or the percentage of positive charges at pH 7.4 decrease. Additional comparisons of peptides can be added to support this statement, as shown for these peptides in different groups (i) at the same level (100%) of positive charges with a different number of H: t_R GRHRQKHS (group 1, 2 H, 100% of positive charges, 15.8 min) > t_R QRHRK (group 2, 1 H, 100% of positive charges, 10 min) > t_R KGKSR (group 2, 0 H, 100% of positive charges, 5.6 min); or (ii) at the same number of histidines in the sequence with different percentages of positive charges: t_R HGSLHKNA (group 2, 2 H, 65% of positive charges, 7.71 min) > t_R HGSLH (group 2, 2 H, 26% of positive charges, 5.48 min). More, W residues might not have a high influence on the peptide retention time considering that the KRHGEWRPS peptide was less retained (1 H and 1 W, 8.6 min) than the peptide QRHRK (1 H only, 10.01 min), knowing that both peptides are 100% positively charged at pH 7.4. If the indole part on the side chain of tryptophan had played a role in metal ion complexation, the retention time of KRHGEWRPS (1 H + 1 W, 100% of positive charges, 8.6 min) should have been closer to that of GRHRQKHS (group 1, 2 H, 100% of positive charges, 15.8 min).

3.1.2. Peptide Retention Time Using HiFliQ NTA-Ni²⁺ Column

Using the HiFliQ NTA-Ni²⁺ column, peptides can be classified in three groups named I, II and III (Table 2). The group I with peptides GH8AY, HHHHHH, and GRHRQKHS have t_R varying between 8 min and 23 min. In addition, group II is constituted of peptides HGSLHKNA, KRHGEWRPS, QRHRK, HGSLH, RHGEWRPS and APHWNIN, which showed retention times between 3 min and 7 min. Finally, other peptides in group III (KGKSR, AEHGSLH, GLH, GLHLPS, YPVGR, KERESH, FPGSA, LTKNV, VSHVN, LLH, LAHS, QSHF, KETQH) were not retained. The retention of peptides on Ni²⁺ in HiFliQ NTA-Ni²⁺ generally follows the same trends as for the HisTrap X-Ni²⁺ (Figure 3A), but peptides were more retained on this latter column. Indeed, peptide KGKSR, which was in

group 2 for HisTrap X-Ni²⁺, is not retained on HiFliQ NTA-Ni²⁺, and peptides in group 3 (AEHGSLH, APHWNIN, GLH, GLHLPS), which had low retention times in HisTrap X-Ni²⁺, are no longer retained on HiFliQ NTA-Ni²⁺, except peptide APHWNIN. Comparing peptide retention times between the two columns, it is observed that the differences in peptide retention times are important for peptides which have a high affinity for Ni²⁺ (GH8AY, HHHHHH, GRHRQKHS and QRHRK) with a mean delta t_R of 6.0 min, while the differences are moderate for other peptides with a mean delta t_R of 1.5 min. This difference in retention times—and thus indirectly in peptide affinities for Ni²⁺—is due to the fact that the columns may have different complexing agents: NTA in the HiFliQ NTA-Ni²⁺ column and an unknown complexing agent in the HisTrap X-Ni²⁺ one.

Whatever the column used, HiFliQ NTA-Ni²⁺ or HisTrap-X-Ni²⁺, it was generally observed that peptides with more histidine and a high percentage of global positive charges at pH 7.4—due to the presence of lysine and arginine residues in their sequences—are more retained. This indirectly indicates that they have a good affinity for Ni²⁺, which is in agreement with previous studies on immobilized Ni²⁺ chelation by peptides using IMAC and SPR [17,27], but arginine and lysine might be responsible of ionic interactions in the presence of X-Ni²⁺.



Figure 3. Comparison between retention times of investigated peptides in HisTrap X-Ni²⁺ and HiFliQ NTA-Ni²⁺ columns (**A**) and peptide-binding dissociation constants, with the red signs indicating peptides with K_D not determined and peptides not retained (**B**).

3.2. Determination of Peptide Affinity Parameters Using SPR

3.2.1. Non-Specific Interactions

The term specific interactions will be used for interactions between the peptide and Ni^{2+} , while non-specific stands for interactions between the peptide and other parts (matrix, complex agent). SPR results were treated by observing at the channel which gives both interactions (active channel, with Ni²⁺) and the channel which represents non-specific interactions (reference channel, without Ni²⁺) in order to understand the effect of nonspecific interactions on peptide retention time. Indeed, the retention time of a peptide in IMAC is obtained considering both specific and non-specific interactions in the column. The 22 peptides were investigated for their affinity for Ni²⁺ immobilized on an NTA chip. Examples of a peptide sensorgram on the active channel (Figure 4A), sensorgram on a reference channel (Figure 4B) and sensorgram for only specific interactions (Figure 4C) are presented below. In some cases, the response measured on the reference channel (without Ni^{2+}) is higher than the response on the active channel (with Ni^{2+}), which can lead to a negative response for a reference-subtracted sensorgram, which is supposed to illustrate only specific interactions. However, it is important to emphasize that the nonspecific bindings are not necessarily perfectly corrected by this former subtraction, since the non-specific bindings might not occur in the same way on the active channel and on the reference channel, thus complicating the discussion of results where higher responses are observed on the reference channel [35]. Considering the reference-subtracted sensorgram, the AEHGSLH and APHWNIN peptide isotherms obtained were inverse and all responses were negative, which can be associated with non-specific bindings. For other peptides like GRHRQKHS, HGSLH and HGSLHKNA, some corrected responses were negative and others were positive, but the overall isotherm still has the Langmuir isotherm shape, as expected. For others, like FPGSA or LTKNV, the responses are negative and there is no correlation between the points of the isotherm, and thus the peptides do not have affinity for Ni²⁺ in the concentration range investigated. Therefore, the negative response might not always refer to non-specific interactions. In addition, the fact that peptides with low molecular weight lead to low responses (overall, if they have little affinity for Ni²⁺) might prevent the results from being very accurate.

3.2.2. Specific Interactions

Each peptide was studied in the range of concentrations that helps to obtain a sorption isotherm as described in Section 2.4 (Figure 2B). From their sorption isotherm, K_D were obtained by fitting a 1:1 binding model and are summarized in Table 3, and the evolution of K_D is illustrated on Figure 3B. The K_D values are reliable when they are lower than half of the maximum concentration studied.

Group	Peptide	<i>K</i> _D (μM)	SE (K _D) (μM)	R _{max} (R.U.)	SE (R _{max}) (R.U.)	Concentration Range (µM)
А	НННННН	0.7	27.0	197.90	27.00	0.4–50
А	GH ₈ AY	1.1	0.5	164.8	11	0.01–100
А	GRHRQKHS	18.7	4.9	28.70	1.90	1–200
А	KGKSR	45.6	37.0	24.9	7.6	20-5000
В	YPVGR	156.6	420.0	15	9	5-5000
В	HGSLHKNA	195.5	64.0	72.38	5.70	5-1000
В	RHGEWRPS	303.9	210.0	81.3	14	20-1000
В	HGSLH	341.5	74.0	38.40	2.20	5-5000
В	QRHRK	910.0	230.0	64.30	5.30	20-5000
В	KRHGEWRPS	737.9	250.0	366.3	30	20-5000
С	GLH	1567.0	760.0	19.30	2.9	20-5000
С	KERESH	4835.0	5400.0	76.9	25	500-10,000
С	GLHLPS	5586.0	5700.0	54.5	22	100-10,000

Table 3. Peptide binding affinity data obtained using SPR and the range of concentrations in which peptides were investigated. n.d.: Not determined.

Group

Peptide

Concentration Range (µM)



600

Time(S)

800

1000

R_{max} (R.U.)

SE (R_{max}) (R.U.)

Table 3. Cont.

SE (*K_D*) (µM)

 K_D (μ M)

200 0

-200 -400 -600 -800 -1000

-1200

200

Resonance(R.U.)

Figure 4. Example of peptide sensorgram for channel with Ni^{2+} (specific + aspecific interactions) (**A**); peptide sensorgram on reference channel (without Ni^{2+} , aspecific interactions) (**B**); and corrected sensorgram (difference of response between 2 previous channels) (**C**).

The significance of the fitted K_D is expressed by standard error (SE) and this SE should be lower than 10% of the K_D value, otherwise K_D can have a wide range of values [36]. In these experiments, all peptides investigated show high values of SE (greater than 10% of the K_D), making it difficult to evaluate if one peptide has a higher affinity for Ni²⁺ compared to another one. Therefore, the affinity of peptides for Ni²⁺ is evaluated by groups of peptides, as described hereafter. First, group A, with K_D less than 85 µM taking into account the standard errors for fitting the data (SE), is composed of peptides HHHHHH, GH8AY, GRHRQKHS and KGKSR. Secondly, group B, with K_D varying between 94 and 1200 µM, is composed of peptides YPVGR, HGSLHKNA, <u>RHGEWRPS</u>, HGSLH, <u>QRHRK</u> and <u>KRHGEWRPS</u>. Then, group C, with K_D higher than 1200 µM, is composed of peptides GLH, KERESH, GLHLPS, QSHF and LAHS. Finally, group D is composed of peptides VSHVN, APHWNIN, AEHGSLH, KETQH, LLH, FPGSA and LTKNV, for which it was not possible to determine the K_D in the range of concentrations investigated. From K_D values, the corresponding association constant (K_A , M^{-1}) was also calculated as $K_A = 1/K_D$. The maximum resonance at equilibrium (R_{max} , R.U.) was obtained generally for the highest peptide concentration. Results are summarized in Table 3. The lower the dissociation constant (K_D) is or the higher the affinity constant (K_A) is, the better the affinity of peptides for Ni²⁺ is.

Concerning group A, peptides HHHHHH, GH8AY, GRHRQKHS and KGKSR showed lower K_D compared to other peptides and thus higher affinity for Ni²⁺. This affinity of peptides for Ni²⁺ can be explained by the presence of two or more amino-acid residues with aromatic rings (i.e., histidine, H) in their sequence. Indeed, according to the literature, HHHHHH was expected to have a good affinity for Ni²⁺ since it showed a lower K_D (0.014 ± 0.001) µM in previous studies [24,37] and GH8AY showed a good affinity for Ni²⁺ according to the literature [25]. The peptide GRHRQKHS shows affinity for Ni²⁺ due to the presence of H at the third position and the presence of a second H at the n + 4 position from the first H. According to former studies [17,24], two histidines separated by three residues (position n: n + 4) is a position that favors the affinity of a peptide for Ni²⁺. In addition, arginine (R) and lysine (K) residues can also participate to increase affinity via ionic interactions with the NTA-Ni²⁺ complex (negatively charged) due to their positive charges as explained in Section 3.1.2. Therefore, KGKSR shows a noticeable affinity due to the presence of two lysine (K) residues and one arginine (R) residues, which lead to the 100% positive charges of this sequence at pH 7.4 (Table 1).

In addition, the peptides of group B (YPVGR, HGSLHKNA, <u>R</u>HGEW<u>R</u>PS, HGSLH, <u>QRHRK</u> and <u>KR</u>HGEW<u>R</u>PS) also show moderate affinity for Ni²⁺. These peptides present at least two histidine (H) and/or tryptophan (W) residues that influenced their ability to chelate Ni²⁺ with, sometimes, additional arginine (R) or lysine (K) that can participate to form other interactions, except peptide YPVGR. Indeed, the affinity of **Y**PVG**R** might be due to the presence of arginine (R) and tyrosine (Y), which is an amino acid also reported to contribute to metal binding [29,38] due to the presence of an aromatic ring and a hydroxyl group on its side chain.

A third group of peptides, C, comprising GLH, KERESH, GLHLPS, QSHF and LAHS sequences shows very low affinity for Ni²⁺ since the saturation has not been reached in the range of concentrations investigated (maximum concentration: 30,000 μ M). The lower affinity for Ni²⁺ observed for these peptides is due to the presence of only one histidine residue in their sequences. In addition, the K_D values of peptides KERESH, GLHLPS and LAHS might not be reliable since they are higher or closer to half of the maximum concentration studied for each peptide [36].

Further, the last group, D, constitutes peptides VSHVN, APHWNIN, AEHGSLH, KETQH, LLH, FPGSA and LTKNV and does not show affinity for Ni²⁺ in the range of concentrations studied. For some peptides, the saturation was not attained and the isotherm was a straight line upward (Figure 2C), which means unknown interactions are increasing and higher concentrations might be needed to reach saturation. For other peptides, negative responses are observed and kept decreasing as the concentration increased (inversed isotherms (Figure 2D) or straight lines downward. Therefore, non-specific interactions, i.e., affinity for the matrix or NTA, are important compared to the affinity for Ni^{2+} . Indeed, AEHGSLH and APHWNIN show the same behaviors with inverse isotherms, characteristic of non-specific interactions increasing with peptide concentration (Figure 2D). These peptides with their respective AE and AP motif at their N-terminal part can probably bother the formation of a complex with Ni²⁺. Indeed, the study of Bal and colleagues [39] reported that alanine (A) at the N-terminus of a peptide can destabilize the complex formed between Cu²⁺ and a peptide; yet, Cu²⁺ and Ni²⁺ are transition metals having the same behaviors according to the HSAB theory. In addition, the proline residue (P) at positions 2, 3 or 4 is known to slow down the metal complexation by inducing the peptide chain to bend [39]. In addition, proline was reported to hinder the deprotonation of the N-terminus, sometimes involved in metal complexation [31]. As for KETQH and LLH, their histidine at

the C-terminal might complicate its accessibility to Ni²⁺ binding, while FPGSA and LTKNV do not contain any histidine to enhance Ni²⁺ chelation.

3.3. Understanding the Peptide-Ni²⁺ Interaction Based on the Analogy between SPR and IMAC

Considering that NTA is a complexing agent used on SPR chips, the HiFliQ NTA-Ni²⁺ column might be a good choice for IMAC experiments to study the SPR-IMAC analogy well. Unfortunately, peptides are better retained in the HisTrap X-Ni²⁺ column, and thus more data are available for this latter column (Figure 3A).

To understand peptide-Ni²⁺ complexation in both SPR and IMAC, the comparison was made between the value of retention times for each peptide in IMAC and the corresponding values of K_A determined in SPR. For this purpose, various trends characterizing the IMAC-SPR analogy were drawn between groups of peptides determined in the IMAC HisTrap X-Ni²⁺ (groups 1, 2, 3, and 4) and HiFliQ NTA-Ni²⁺ (groups I, II, III) and those determined in SPR (groups A, B, C, and D) (Table 4).

Table 4. Investigation of the validity of the analogy between IMAC and SPR. Comparison is based on couples formed based on groups of peptides in IMAC (group 1, 2, 3, 4 for HisTrap X-Ni²⁺ and group I, II, III for HiFliQ NTA-Ni²⁺) and SPR (group A, B, C, D).

Peptide	HisTrap X-Ni ²⁺	HiFliQ NTA-Ni ²⁺
GH8AY	1A	IA
НННННН	1A	IA
GRHRQKHS	1A	IA
QRHRK	2B	IIB
KRHGEWRPS	2B	IIB
HGSLHKNA	2B	IIB
KGKSR	2A	IIIA
HGSLH	2B	IIB
RHGEWRPS	2B	IIB
AEHGSLH	3D	IIID
APHWNIN	3D	IID
GLH	3C	IIIC
GLHLPS	3C	IIIC
YPVGR	4B	IIIB
KERESH	4C	IIIC
FPGSA	4D	IIID
LTKNV	4D	IIID
VSHVN	4D	IIID
LLH	4D	IIID
LAHS	4C	IIIC
QSHF	4C	IIIC
KETQH	4D	IIID

The first couple, 1A, is composed of peptides GH8AY, GRHRQKHS and HHHHHH, which have high constant affinity in SPR and are also highly retained in IMAC since they are eluted with a high quantity of imidazole compared to the other peptides. The second couple, 2B, constitutes peptides QRHRK, KRHGEWRPS, HGSLHKNA, HGSLH and RHGEWRPS, which have moderate affinity for Ni²⁺ in SPR and are also well retained in IMAC. The third couple, 3C, is composed of the peptides GLH and GLHPS, which have low affinity for Ni²⁺ in SPR and have low retention in IMAC as well. The fourth couple, 4D, concerns peptides FPGSA, LTKNV, VSHVN, LLH and KETQH, which have no affinity for Ni²⁺ in SPR and are not retained in IMAC.

However, beyond these direct and major trends correlating IMAC and SPR, sometimes trends are not so obvious with some exceptions observed for peptides having lower/no affinity for Ni²⁺ in IMAC (group 3, 4) or SPR (C, D). These peptides are KGKSR (2A), AEHGSLH (3D), APHWNIN (3D), YPVGR (4B), KERESH (4C), LAHS (4C) and QSHF (4C). It can be observed that some peptides can have lower retention in IMAC and do not have

Concerning HiFliQ NTA-Ni²⁺, a similar trend was observed and couples are illustrated in Table 4, where couple IA contains peptides with higher affinities that are the same as those for couple 1A, and couple IIB contains peptides with moderate affinities, like for couple 2B. Due to less data for this HiFliQ NTA-Ni²⁺ column, other peptides are in couple IIIC or IIID with very low/no affinities.

The main exceptions concern KGKSR and YPVGR (two peptides without H); despite their good affinity constant in SPR, KGKSR is eluted with a low quantity of imidazole for the HisTrap X-Ni²⁺ column and it is not retained in the HiFliQ NTA-Ni²⁺ column, while YPVGR is not retained in either column. Indeed, for these two peptides, the affinity determined in SPR mainly involves weak ionic interactions with Ni²⁺, which cannot be corrected even with the signal observed on the reference channel (uncharged in Ni²⁺). Yet, in IMAC, these weak ionic interactions between Ni²⁺ and peptides (KGKSR or YPVGR) are easily destroyed upon elution with imidazole (a competitive agent), which explains their low/absence of retention.

Taken all together, these results demonstrate experimentally for the first time the analogy between SPR and IMAC. Indeed, for around 70% of the peptides investigated in the HisTrap X-Ni²⁺ and for around 60% of peptides studied in the HiFliQ NTA-Ni²⁺, peptides which have good affinity for immobilized Ni²⁺ in SPR also have a good affinity for Ni²⁺ in IMAC, and peptides without affinity for Ni²⁺ in SPR do not have a good affinity for Ni²⁺ in IMAC, as is illustrated in Figure 3. In addition, the analogy is clear for peptides with higher affinity for Ni²⁺ in SPR or IMAC.

4. Conclusions

Among the 22 peptides investigated in IMAC and SPR, 70% (HisTrap X-Ni²⁺) and 60% (HiFliQ NTA-Ni²⁺) of the peptides validated the analogy between IMAC and SPR: peptides with high retention times in IMAC were also endowed with a high affinity for Ni²⁺ in SPR. Both in IMAC and SPR, the affinity for immobilized Ni²⁺ depends on the position and the number of histidine (H) residues present in the sequence, while tryptophan (W) residues have very low/no impact. In addition, the high percentages of global positive charges at pH 7.4, which is due to the presence of lysine (K) and arginine (R) residues in the sequences, were also responsible of high peptide retention times in IMAC and high affinities in SPR, which is related to ionic interactions. Therefore, an increase in the NaCl concentration will be further considered to decrease ionic interactions so that coordination interactions between the peptide and the immobilized metal ion can be observed solely. This will improve the SPR-IMAC correlations for further chromatographic simulation.

In addition, since the affinity of peptides is also related to the charge of the Ni(II)peptide complexes present at pH 7.4, it would be useful to also investigate the charge and composition of the supposed Ni(II)-peptide complexes formed at 7.4 by various analytical techniques to go deeper in the interpretation of SPR and IMAC results, notably via pH potentiometry measurement.

Supplementary Materials: The following supporting information can be downloaded at: https://www. mdpi.com/article/10.3390/inorganics12010031/s1, Data S1: Pea protein sequences from UniprotKB; Data S2: IMAC column specifications.

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