brought to you by Torrector

Electrophoresis

Global properties and propensity to dimerization of the amyloid-beta (12-28) peptide fragment through the modeling of its monomer and dimer diffusion coefficients and electrophoretic mobilities. Julio A. Deiber<sup>1</sup>, Marta B. Peirotti<sup>1</sup>, Maria V. Piaggio<sup>2</sup> <sup>1</sup>Instituto de Desarrollo Tecnológico para la Industria Química (INTEC), Universidad Nacional del Litoral (UNL), Consejo Nacional de Investigaciones Científicas y Técnicas (CONICET), Santa Fe, Argentina <sup>2</sup>Cátedra de Bioquímica Básica de Macromoléculas, Facultad de Bioquímica y Ciencias Biológicas, UNL, Santa Fe, Argentina

**Running title**: Properties and propensity to dimerization of amyloid-beta (12-28)

**Correspondence**: Dr. Julio A. Deiber, INTEC, Güemes 3450, 3000, Santa Fe, Argentina

E-mail: <u>treoflu@santafe-conicet.gov.ar</u>

Fax: +54-(0)342-4550944

Received: 12-Aug-2014; Revised: 20-Oct-2014; Accepted: 05-Nov-2014

This article has been accepted for publication and undergone full peer review but has not been through the copyediting, typesetting, pagination and proofreading process, which may lead to differences between this version and the Version of Record. Please cite this article as doi: 10.1002/elps.201400395.

Abbreviations: AAS, amino acid sequence; Aβ, amyloid-beta; AD,
Alzheimer´s Disease; IPCR, intraparticle charge regulation; PLLCEM,
Perturbed Linderstrøm-Lang Capillary Electrophoresis Model; PPCR, pair particle charge regulation.

**Key words:** amyloid-beta fragment / charge regulation phenomena / dimerization equilibrium / effective electrophoretic mobility / translational diffusion coefficient

Manuscript total number of words and figure legends is 5531

### Abstract

Neuronal activity loss may be due to toxicity caused mainly by amyloid-beta (1-40) and (1-42) peptides forming soluble oligomers. Here the amyloid-beta (12-28) peptide fragment (monomer) and its dimer are characterized at low pH through the modeling of their diffusion coefficients and effective electrophoretic mobilities. Translational diffusion coefficient experimental values of monomer and dimer analogues of this peptide fragment and monomer and dimer mixtures at thermodynamic equilibrium are used as reported in the literature for different monomer initial concentrations. The resulting electrokinetic and hydrodynamic global properties are employed to evaluate the amyloid-beta (12-28) peptide

fragment propensity to dimerization through a thermodynamic theoretical framework. Therefore equilibrium constants are considered at pH 2.9 to elucidate one of the amyloidogenic mechanisms involving the central hydrophobic region LVFFA of the peptide spanning residues 17-21 associated with phenylalanine at positions 19 and 20 in the amino acid sequence of amyloid-beta peptides. An analysis demonstrating that peptide aggregation is a concentration-dependent process is provided, where both pair and intraparticle charge regulation phenomena become relevant. It is shown that the modeling of the effective electrophoretic mobility of the amyloid-beta (12-28) peptide fragment is crucial to understand the effect of hydrophobic region LVFFA in the amyloidogenic process.

#### **1** Introduction

Global electrokinetic and hydrodynamic properties of peptides are needed to find out key biophysical mechanisms associated with bioprocesses involved in human pathologies (see for instance Refs. [1-4]). In this regard, the amyloidogenic process found in Alzheimer's Disease (AD) is a relevant case study [4]. At present the definite etiopathology of AD is not well known [5, 6], although several hypotheses have been proposed, and among them, the loss of neuronal activity due to toxicity caused by amyloid-beta (A $\beta$ ) peptides forming soluble aggregates is considered one of the most probable. In fact the histopathology is characterized by extracellular plaques formation caused by aggregation of mainly A $\beta$ (1-42) with an amino acid sequence (AAS) DAEFRHDSGYEVHHQKLVFFAEDVGSNKGAIIGLMVGGVVIA, and  $A\beta(1-40)$  where the terminal I(41) and A(42) are not present [4, 5]. In previous studies emphasis has been placed on the fact that an increase in A $\beta$  soluble oligomers, intermediates, and fibrils with a progressive formation of amyloid plaques was found, where addition of new aggregates occurred. Nevertheless the specific molecular mechanisms yielding extracellular plaques are not well understood yet. In a previous work we have shown [4] that electrokinetic and hydrodynamic properties of  $A\beta$  peptides evaluated from the modeling of their effective electrophoretic mobilities at well defined BGE properties were required to estimate the propensity to aggregation of these polyampholyte-

Electrophoresis

polypeptide chains having weak ionizing groups. It was shown that CE is one of the most appropriate experimental methods to achieve the evaluation of a well defined set of global properties needed in calculations involving peptide Brownian aggregation theories [4]. Also it was pointed out that A $\beta$  peptide concentration, aggregate sizes and near molecule pH due to the pair particle charge regulation (PPCR) phenomenon might be crucial parameters in the amyloidogenic process. The PPCR phenomenon is a relevant and still no well described effect in polypeptide interactions, which must be added to the classical intraparticle charge regulation (IPCR) phenomenon [7-12] giving the near molecule pH designated pH<sup>\*</sup> [4]. In the present work we continue this study concomitantly with another relevant experimental work [13] on this subject. Now the focus is placed on the propensity to dimerization of the amyloid-beta (12-28) peptide fragment. The rationale is that phenylalanine at positions 19 and 20 in the AAS of A $\beta$  peptides may be a zone of nucleation in the oligomerization process. Since some peptides contain specialized domains enhancing dimerization, we will refer to the homo-dimerization of the A $\beta$ (12-28) peptide fragment common to most A $\beta$  peptide AASs relevant in AD. Further an analysis concerning the effect of initial monomer concentration on this peptide aggregation will be provided.

It is worth to observe that basic transport properties like diffusion and friction coefficients, intrinsic viscosity and electrophoretic mobility are interconnected

through some common phenomenological aspects, among which chain friction is the relevant information sensitive to hydration, size and shape of peptides visualized as different types of hydrodynamic particles [14, 15]. In particular the electrophoretic mobility is proportional to the effective charge number of ionizing groups in the AAS presenting a challenging for screening out the electrokinetic and hydrodynamic phenomena involved in the polyampholytepolypeptide heterochains [3, 4, 7-12, 14-29]. The comprehension of the interplay among these transport properties through rigorous models is useful in modern biotechnology as described previously in many publications. Their measurements through different analytical techniques like for instance CE [30-34] at well defined temperature T, pH, ionic strength I, electrical permittivity  $\varepsilon$ and viscosity  $\eta_s$ , are valuable results for studying biophysical mechanisms and process.

This work presents the evaluation of global properties of monomer and dimer analogues of the A $\beta$ (12-28) peptide fragment. For this purpose, experimental data of translational diffusion coefficient for monomer and dimer analogues and monomer and dimer mixtures at thermodynamic equilibrium provided by Ref. [13] were modeled to estimate the friction coefficients of these hydrated particles. In fact, monomer and dimer analogues were obtained so that diffusion coefficient measurements could be carried out avoiding monomer dimerization and conversely impeding dimer reversion to monomer. Further, diffusion

coefficients of the monomer and dimer mixtures were obtained from the A $\beta$ (12-28) peptide fragment at different initial monomer concentrations. This interesting proposal provided valuable experimental information to visualize the propensity to dimerization of the A $\beta$ (12-28) peptide fragment in the present work. In fact, friction coefficients were calculated via our previous theoretical developments [12, 14], which then allowed us the estimation of the corresponding effective electrophoretic mobilities. Although it is preferable to use directly effective electrophoretic mobilities values obtained from CE [30-34], this method has not been exploited appropriately for this type of subject despite it is offering the relevant experimental information as already indicated in the literature [4, 35-42]. Here we find this alternative valid because the Perturbed Linderstrøm-Lang Capillary Electrophoresis Model (PLLCEM) [7-12, 14, 15, 22, 25-27] may be also applied when the input data is the diffusion coefficient value (Supporting Information). This model provides electrokinetic and hydrodynamic properties necessary to study the equilibrium between monomer and dimer and to evaluate the associated total free energy change. For this purpose, the thermodynamic theory established previously [9, 43] was used as modified and presented below for the case of monomer and dimer thermodynamic equilibrium.

Accepted Artic

In this work, Section 2 is split in two parts: (i) Section 2.1 describes the monomer, dimer and their analogues, and the experimental information used here. (ii) Section 2.2 presents theoretical thermodynamic considerations of peptide dimerization involving electrokinetic and hydrodynamic global properties obtained from the PLLCEM for different initial monomer concentration. Section 3 discusses results of the dimerization process placing emphasis on the effect of the IPCR and PPCR phenomena. Finally Section 4 provides concluding remarks and suggestions for future researches.

#### 2 Materials and methods

## 2.1 The A $\beta$ (12-28) peptide fragment and diffusion coefficient experimental data

For the purpose of using a non-aggregating monomer analogue at low pH in Ref. [13], the monomer AAS expressed VHHQKLVFFAEDVGSNK was changed to VHHQKLVGGAEDVGSNK designated here monomer analogue and coded as  $[G^{19,20}]A\beta(12-28)$ . Thus the FF at position 19 and 20 of the A $\beta(1-42)$  peptide were changed to GG. Also to use a non-reverting dimer analogue, a modified dimer was generated through a disulfide-linked bond of two monomer analogues. These mutated monomer analogues had each one the F at position 20 changed to C giving the dimer analogue (VHHQKLVFCAEDVGSNK)<sub>2</sub> coded as  $([C^{20}]A\beta(12-28))_2$  in [13]. The diffusion coefficient of the mixture was composed of monomer and dimer indicating that the interconversion between

these species at equilibrium was a rapid kinetic process thus providing one mixture diffusion coefficient value only (see also discussion in [13] and below). Thus peptide chains are sampling rapidly both states giving weighed average values of mixture diffusion coefficients. Therefore experimental diffusion coefficients of these peptides solutions were obtained at 25°C via pulsed-field gradient nuclear magnetic resonance spectroscopy. The peptides were dissolved in D<sub>2</sub>O adjusted to pD 2.9 with DCl giving an ionic strength  $I \approx 1$  mM. Peptide solutions were prepared in the range  $C_0 = 0.5-3.25$  mM by serial dilution. For  $C_{o} > 3.25$  mM the peptide fragment solubility was highly diminished. Experimental values of monomer and dimer mixture diffusion coefficient D of A $\beta$ (12-28) peptide fragment reported in Ref. [13] was fitted through the polynomial  $D \approx (2.4432 - 0.1955C_o + 0.0224C_o^2) \times 10^{-10} \text{ m}^2/\text{s}$  with a correlation coefficient  $r^2 \approx 0.97$ . Further, the monomer  $D_M = 2.48 \times 10^{-10} \text{ m}^2/\text{s}$  and dimer  $D_D = 1.78 \times 10^{-10} \text{ m}^2/\text{s}$  diffusion coefficient analogues were provided, which in principle were independent from C<sub>o</sub> within the concentration range studied [13].  $D_2O$  properties required in our calculations are available in the Supporting Information. The A $\beta$ (12-28) peptide fragment, like A $\beta$ (1-40) and A $\beta$ (1-42) peptides, forms amyloid fibrils in vitro for pH>4. One reason is that their chain negative and positive ionizing groups may form salt bridges stabilizing the  $\beta$ sheet formations. Nevertheless at pH 2.9, the A $\beta$ (12-28) has a positive effective

charge number while the residual and terminal carboxylic groups are partially protonated producing dimers driven by hydrophobic forces [13].

### 2.2 Theoretical concepts and considerations

## **2.2.1** The A $\beta$ (12-28) peptide fragment dimerization from the

## thermodynamic point of view

Accepted Articl

In general the oligomerization of peptide chains is a concentration-dependent process that may be visualized from different thermodynamics and kinetics theories [4, 13]. Here we will refer to the dimerization process of A $\beta$ (12-28) peptide fragments at pH 2.9 where equilibrium between monomer and dimer structures can be found [13]. Therefore, monomer and dimer (species  $\alpha = M, D$ ) follow the dimerization stoichiometry  $2M \leftrightarrow D$ . Defining  $X_{\alpha}$  as the molar fraction of species  $\alpha$ , the dimerization equilibrium constant is  $K = X_D / X_M^2$ . The chemical potentials  $G_{\alpha}$  of these polyampholyte-polypeptide chains with  $N_{\alpha}$ amino acid residues are required.  $G_{\alpha}$  are associated with the conformations of chains in the BGE and the electrostatic interactions among charged ionizing groups and BGE ions, which are responsible of the IPCR phenomenon described previously in the literature [7-12]. Thus the pH sensed by any electrically charged chain is not necessarily equal to the bulk pH of the BGE. Therefore from [9], one can express  $G_{\alpha} = G_{\alpha}^* + RT \ln X_{\alpha} + e N_A Z_{\alpha} \zeta_{\alpha}/2$ , where  $N_A$ is the Avogadro constant,  $R = N_A k_B$  is the gas constant,  $k_B$  is the Boltzmann

constant, e is the elementary charge and  $G_{\alpha}^{*}$  is the reference chemical potential that depends on T and  $pH_{\alpha}^{*}$ , and hence on  $C_{o}$  when the PPCR phenomenon becomes relevant (see below). Also  $Z_{\alpha}$  is the net charge number and  $\zeta_{\alpha}$  is the mean field zeta (electrokinetic) potential of species  $\alpha$ . Therefore for a given dimerization degree  $\Delta G^{eff} = \Delta G^{*} + eN_{A}(Z_{D}\zeta_{D} - 2Z_{M}\zeta_{M})/2$  is the effective free energy change, where  $\Delta G^{*} = G_{D}^{*} - 2G_{M}^{*}$  is the conformational free energy change yielding  $K = exp(-\Delta G^{eff} / RT)$ . Here the electrostatic free energy change  $\Delta G^{el}$  due to the dimerization equilibrium, defined  $\Delta G^{el} = eN_{A}(Z_{D}\zeta_{D} - 2Z_{M}\zeta_{M})/2$ , is sensitive to  $pH_{\alpha}^{*}$  due to IPCR phenomenon (Supporting Information) expressed,  $pH_{\alpha}^{*} \approx pH + e^{2}Z_{\alpha}/\{ln(10)k_{B}T4\pi\epsilon a_{H\alpha}(1+\kappa a_{H\alpha})\}$  (1)

where  $a_{H\alpha}$  is the hydrodynamic radius of specie  $\alpha$  and  $\kappa = (2Ie^2N_A 10^3/\epsilon k_B T)^{1/2}$  is the Debye-Hückel parameter. Equation (1) applies for analyte infinite dilution only. For increasing values of  $C_o$ , this equation must be modified when a pair of charged particles comes near one another, giving an additional shift in the  $pH_{\alpha}^*$  due to the PPCR phenomenon. Assuming that interactions of pair particles of equal size predominate [4], the  $pH_{\alpha}^*$  may be expressed as follows,

$$pH_{\alpha}^* \approx pH + e^2 Z_{\alpha} \{1 + e^{-\kappa d}\} / \{\ln(10)k_B T 4\pi \epsilon a_{H\alpha}(1 + \kappa a_{H\alpha})\}$$

(2)

In this expression the average distance d between two particle centers of

Electrophoresis

species  $\alpha$  depends on  $C_o$  and the dimerization degree  $\xi,$  giving monomer  $C_M = C_o - 2\xi$ , dimer  $C_D = \xi$  and total concentrations  $C = C_o - \xi$ . Therefore d may be estimated from  $d \approx m (3/4 \pi C N_A 10^3)^{1/3}$  involving peptide volume fraction in the solution at each  $\xi$  value. In this regard parameter m is obtained by assuming that charged particles are in a dynamic lattice having each cell a radius  $r^* \approx (3/4 \pi C N_A 10^3)^{1/3}$ . Therefore  $m \approx 1.207$  by considering that the average approach of pair interacting particles is between  $r^*$  and  $\sqrt{2}r^*$  as derived from simple geometric calculations. In Eq. (2) the electrical potential of the interacting charged particles is  $\psi_{\alpha}(d) \approx \zeta_{\alpha} e^{-\kappa d}$  as a first approximation, where the hypothesis of interacting effective point charges far from contact is introduced. For  $C \rightarrow 0$  and  $d \rightarrow \infty$ , Eq. (1) is physically recovered at infinite dilution. Therefore, it is clear that the PPCR phenomenon introduces an additional shift of the  $pH_{\alpha}^{*}$  toward the isoelectric point pI of species  $\alpha$  chain, which is also a phenomenon controlling the interaction forces between peptides and proteins [3, 4, 25-27] as illustrated and discussed below in Section 3 for the peptide fragment under study. These approximations are used to illustrate simply the PPCR phenomenon and for clearness. A more precise numerical calculation may lead one to a difficult task, which is out of the purpose of the present work. From Eqs (1) and (2) one finds that the effective free energy must

depends on  $C_o$ , which is the formal demonstration that the aggregation of peptides is a concentration-dependent process (see also numerical validation below) because  $G_{\alpha}$  depends on  $\zeta_{\alpha}$  as well as on  $pH_{\alpha}^*$  including both the IPCR and PPCR phenomena. Therefore following our previous work [9] rewritten for this particular problem here, the molar fractions of monomer

 $X_M = (D - D_D)/(D_M - D_D)$  and dimer  $X_D = (D_M - D)/(D_M - D_D)$  are obtained from the calculation of the mixture diffusion coefficient  $D = X_D D_D + X_M D_M$ . This last expression applies when the rate of interconversion from monomer to dimer of any chain in the solution is in the fast change regime, which indicates that the monomer and dimer mixture diffusion coefficient is an average value [9, 13, 43]. Then within the CE framework the effective electrophoretic mobility  $\mu_{\alpha}$ for  $\alpha = M$ , D may be combined into an average mobility  $\mu = \mu_M X_M + \mu_D X_D$ , where  $\mu$  is the effective electrophoretic mobility of monomer and dimer mixture of the AD plaque-competent A $\beta$ (12-28) peptide fragment, while  $\mu_M$  and  $\mu_D$  are the effective electrophoretic mobilities of the monomer and dimer analogues, respectively, which are AD plaque-incompetent peptides fragment analogues, as expressed in [13].

#### 2.2.2 Electrokinetic and hydrodynamic global properties

To evaluate the electrokinetic and hydrodynamic global properties of monomer and dimer analogues, the PLLCEM was used. Its applicability to polyampholyte-polypeptide chains has been discussed in detail in [3, 4, 7-12, 14, 15, 22, 25-27] by pointing out the approximations introduced. This model can be used with either diffusion coefficient or effective electrophoretic mobility values as input data (Supporting Information). Here diffusion coefficient experimental values  $D_{\alpha} = k_B T / f_{\alpha}$  of species  $\alpha$  reported in [13] are used to obtain friction coefficient  $f_{\boldsymbol{\alpha}}$  of monomer and dimer analogues (Supporting Information Table 1). Then the following electrokinetic and hydrodynamic global properties of monomer and dimer analogues with  $\alpha = M$ , D are obtained: effective  $Z_{\alpha} = Z_{+\alpha} - |Z_{-\alpha}|$ , positive  $Z_{+\alpha}$ , negative  $Z_{-\alpha}$  and total  $Z_{T\alpha}$  charge numbers, effective charge number fraction  $\Delta\sigma_{\alpha}{=}\left|Z_{\alpha}\right|/N_{\alpha},$  total charge number fraction  $\sigma_{\alpha}$  =  $Z_{T\alpha}/N_{\alpha},$  approximate hydration  $H_{\alpha}$  (number of water molecules per chain) or  $\delta_{\alpha}$  (gram of water/gram of peptide), size estimated via the equivalent hydrodynamic or Stokes radius  $\,a_{H\alpha}\,,$  compact radius  $\,a_{c\alpha}\,$ accounting the chain mass only, friction ratio  $\Omega_{\alpha}$  (also designated asphericity) where both the stick and slip between fluid and hydrated particle may be considered [27], pH near molecule designated  $pH_{\alpha}^{*}$  due to both IPCR and PPCR phenomena. Therefore the estimation of  $\mu_M,\,\mu_D$  and  $\mu$  are readily calculated as

Electrophoresis

indicated in the Supported Information and Section 2.1. In addition species  $\alpha$  hydration is defined  $H_{\alpha} = H_{o\alpha} + H_{d\alpha}$  [4], where  $H_{o\alpha}$  is the number of water molecules captured by amino acid residues of the AAS, while  $H_{d\alpha}$  is the number of water molecules due to the degree of water occlusion or release by peptides [4]. Concerning the physical interplay between particle shape and hydration described previously [4] one may consider the minimum  $\delta_{\alpha}$  sampled by the chain ( $H_{d\alpha} = 0$  and  $\Omega_{\alpha} < 1$ ) and the maximum  $\delta_{\alpha}$  physically admissible ( $H_{d\alpha} > 0$  and  $\Omega_{\alpha} = 1$ ). Further partial BGE slip on particle surface applies when  $\Omega_{\alpha} > 1$  with  $H_{d\alpha} = 0$  for both spherical and aspherical hydrated particles [4, 27]. In this case  $H_{d\alpha} < 0$  is required to obtain BGE stick on particle surface with  $\Omega_{\alpha} = 1$  as long as  $a_{H\alpha} \ge a_{c\alpha}$  is satisfied.

### **3** Results and discussion

# **3.1 Electrokinetic and hydrodynamic global properties of monomer and dimer analogues**

Supporting Information Table 2 shows main global properties of monomer and dimer analogues at pH 2.9 obtained within the framework of Section 2.2 for  $H_{d\alpha} = 0$ , when the IPCR phenomenon is included only (Eq. (1)). Although an extended set of properties are presented in this table, we will not describe fully this numerical information here because one may use [3, 4, 7-12, 14, 15, 22, 25-27] for direct interpretations. Instead this discussion section will be centered on

Electrophoresis

physical aspects considered relevant for the visualization of the mechanisms
associated with the amyloidogenic process of interest in AD. Thus when the
IPCR phenomenon is considered, the charge numbers of residual $Z_{AspM} \approx -0.49$ ,
$Z_{GluM} \approx -0.28$ , $Z_{ArgM} \approx 1.00$ , $Z_{HisM} \approx 0.99$ , $Z_{LysM} \approx 1.00$ , and terminal
$Z_{NH_2M} \approx 1.00$ , $Z_{COOHM} \approx -0.60$ ionizing groups in the monomer analogue show
that ionizing amino groups are fully protonated, while ionizing carboxylic
groups are partially protonated yielding relatively a high effective charge
number fraction $\Delta \sigma_M \approx 0.21$ [14, 15, 22]. In this regard, one expects an enough
electrostatic repulsion among monomer analogue chains avoiding for instance
the aggregation via salt bridges stabilized with beta-sheet conformations (this
physical interchain interaction requires a very close approximation between
particles of around 4 Å [44]). These quantitative results are consistent with the
discussion in [13] and citations therein. In fact the dimerization of the monomer
occurs mainly via hydrophobic forces associated with the central region LVFFA
(the monomer analogue did not present dimerization at any $C_o$ because this
zone was changed to LVGGA). From this table one finds that $pH_M^* \approx 3.98$ is
lower than $pH_D^* \approx 4.31$ while both monomer and dimer analogues present the
same pI 7.09. It should be observed that at pI, charge regulation phenomena
cancel out in polyampholyte chains [3, 4, 25, 26]. Therefore it is clear that the
dimerization vielded a $pH_{D}^{*}$ closer to the pI as a consequence of the IPCR

Electrophoresis

Accepted Article

phenomenon (below in Section 3.3, it is shown that this shift is enhanced even  
more due to the PPCR phenomenon). The dimer analogue has 
$$Z_{AspD} \approx -0.67$$
,  
 $Z_{GluD} \approx -0.45$ ,  $Z_{ArgD} \approx 1.00$ ,  $Z_{HisD} \approx 0.99$ ,  $Z_{LysD} \approx 1.00$ ,  $Z_{NH_2D} \approx 1.00$  and

 $Z_{COOHD} \approx -0.76$ , showing that ionizing amino groups remained fully protonated but ionizing carboxylic groups increased their negative charge numbers yielding relatively lower  $\Delta \sigma_{\rm D} \approx 0.18$ , what imply less electrostatic intraparticle repulsion. Therefore the dimer seems to be a more stable particle than the monomer from the electrostatic point of view. This phenomenon may occur in the interactions of proteins and peptides in general. Its deeper elucidation and additional applications will be worth to be considered in future researches. In a forthcoming work we will show that this phenomenon is enhanced even more when oligomers of higher order are formed. This last aspect is also consistent with the fact that for pH>4 (nearer the pI) other higher oligomers must be consider for the A $\beta$ (12-28) peptide fragment ending up in complex structures like protofibrils and fibrils typically found in AD [13, 45]. This result is not surprising taking into account that at higher protocol pH the monomer will approach closer the pI 7.09 due to the  $pH_M^*$  shift expected. On the other hand it is not simple to visualize how the monomer may have a good propensity to aggregation once the central hydrophobic nucleation occurs, unless it is a consequence of an additional pH near molecule shift toward the pI as explained and discussed above.

From Supporting Information Table 2, one also finds that $\Omega_{\alpha} > 1$ with $H_{d\alpha} = 0$
and the monomer and dimer analogues can be considered as sampling a
spherical particle with partial BGE slip at the surface [27]. This result is
probably associated with a poor packing of $D_2O$ on the particle surface. The
$D_2O$ molar mass is higher than that of $H_2O$ causing the "cage effect" already
described previously in [27]. Further, when $H_2O$ was considered as BGE in
calculations, both $\Omega_{\alpha}$ became near unity from below. Thus BGE stick condition
was found showing that the apparent slip is due to solvent type used in the BGE
formulation. This conclusion is reinforced with the fact that the monomer and
dimer analogues are relatively low hydrophilic chains as indicated in the
Supporting Information, and hence one does not expect a high slip as it was
found mainly with hydrophobic chains in [27]. The discussion of this section
may be readily extended to the cases when the monomer and dimer analogues
are considered particles sampling the minimum hydration with $H_{\alpha} < 0$ as
required to reduce $\Omega_{\alpha} > 1$ to $\Omega_{\alpha} = 1$ (Supporting Information Table 3).
3.2 Monomer and dimer mixture effective electrophoretic mobility and

## 3.2 Monomer and dimer mixture effective electrophoretic mobility and properties

Supporting Information Table 2 shows  $\mu_M < \mu_D$  despite they are expect to be rather equal. There is a subtle phenomenon to be explained with this result by taking into account that from the experimental information in Ref. [13]  $D_M > D_D$ , which is the opposite relationship to that concerning effective This article is protected by copyright. All rights reserved.

electrophoretic mobilities. Thus while  $D_{\alpha} \propto 1/f_{\alpha}$  and the later inequality applies, one observes that  $\mu_{\alpha} \propto Z_{\alpha}/f_{\alpha}$  (here hydration and IPCR phenomenon must be included [4]) permitting a different inequality. Therefore results in this table show that the dimerization of the monomer analogue yields a higher increase of effective charge number than an increase in friction coefficient. The explanati is that  $pH_M^* < pH_D^*$  (3.98 compared to 4.31), as indicated above in Section 3.1, giving  $Z_D \approx 6.19$  less than twice  $Z_M \approx 3.62$ . In addition,  $\delta_M > \delta_D$  (0.55 compare to 0.51) affecting the volume size of the dimer and resulting in  $f_D \approx 2.31 \times 10^{-11}$  kg/s less than twice  $f_M \approx 1.66 \times 10^{-11}$  kg/s value. Consequently it is clear that in general global properties of the dimer cannot be taken twice those of the monomer due to both IPCR phenomenon and particle hydrations. It must be also pointed out here that in general for high  $Z_{\alpha}$  the ratio  $e\zeta_{\alpha}/k_BT>2.5$ , which is a value around the onset of ion polarization-relaxation effects (see details in [14, 46, 47]) and  $\mu_{\alpha}$  may become saturated taking an asymptotic value at very high  $\zeta_{\alpha}$ . Nevertheless, the cases considered here presented  $\kappa a_{H\alpha} < 0.02$  extending even more the admissible values of electrokinetic notentials [14] effective charge number than an increase in friction coefficient. The explanation giving  $Z_D \approx 6.19$  less than twice  $Z_M \approx 3.62$ . In addition,  $\delta_M > \delta_D$  (0.55 compared  $^{11}$  kg/s less than twice  $f_{M} \approx 1.66 \times 10^{-11}$  kg/s value. Consequently it is clear that

electrokinetic potentials [14].

Figure 1 shows the monomer and dimer mixture effective electrophoretic mobility  $\mu$  as a function of  $C_o$  in relation to the monomer and dimer analogues  $\mu_M$  and  $\mu_D$  when the PPCR phenomenon has been neglected. For increasing  $C_o$ 

it is clear that  $\mu \rightarrow \mu_D$  (at  $C_o \approx 3.25$  mM this peptide fragment is near the limit of solubility [13]) while the opposite is true for  $C_{\rm o}$   $\rightarrow 0\,$  giving  $\,\mu \!\rightarrow\! \mu_{\rm M}$  (for  $C_{\rm o}\!<\!0.5$ mM diffusion coefficient measurements presented low accuracy [13]). Thus the modified monomer and dimer in Ref. [13] are quite good analogues of the monomer and dimer associated with the A $\beta$ (12-28) peptide fragment. Further, in relation to thermodynamic calculations, Supporting Information Table 4 shows how the dimerization degree, equilibrium constant and effective free energy change vary with  $C_o$  validating the theoretical reasoning and analysis given above in Section 2.1. This table also shows that  $\Delta G^{\text{eff}}$  changes from positive to negative values (441.3 to -254.6 J/mol) at around  $C_0 \approx 1.25$ -1.5 mM yielding a favorable free energy change to dimerization of the A $\beta$ (12-28) peptide fragment when pH 2.9. Since  $\Delta G^{\text{eff}} \approx 0$  for  $C_o \approx 1.37$  mM, this initial monomer concentration value is a good estimate to indicate a significant nucleation for conversion to dimer of the peptide fragment giving  $X_D \approx 0.383$ . Further Supporting Information Figure 1 is equivalent to Figure 1 where  $\mu$ ,  $\mu_M$  and  $\mu_D$ are presented as a function of  $X_D$  where two extrapolating point are included at  $X_D = 0$  and 1 to show the consistency of calculations in terms of dimer molar fractions [9, 43].

## **3.3 Effects of the PPCR phenomenon for increasing initial monomer** concentration

Figure 2 shows the mixture effective electrophoretic mobility  $\mu$  as a function of C<sub>o</sub> in relation to the monomer and dimer analogues  $\mu_M$  and  $\mu_D$  when both IPCR and PPCR phenomena are considered in calculations (Eq. (2)). Once more for low and high C<sub>o</sub> the expected asymptotic tendencies of  $\mu$  are observed. Nevertheless the monomer and dimer analogues effective electrophoretic mobilities decrease with higher values of C<sub>o</sub> as a consequence of the inclusion of the PPCR phenomenon coupled also to particles hydration. The differences found in the comparison between Figure 1 and 2 demonstrate the relevance of the PPCR phenomenon when two peptides approach one another at higher C<sub>o</sub> values. To complement the illustration of the rapid interconversion rates of chains in the dimerization process, the change of the mixture effective charge number  $Z = Z_M X_M + Z_D X_D$  with C<sub>o</sub> is shown in the Supporting Information Figure 2.

At each increment of  $C_o$ , the PPCR phenomenon introduces an additional near molecule pH shifts in the calculation of the  $pH_{\alpha}^*$  previously shown in the Supporting Information Table 2. This effect is illustrated in Figure 3 where  $pH_{\alpha}^*$ as a function of  $C_o$  due to PPCR phenomenon is shown. In fact, at infinite dilution  $pH_M^* \rightarrow 3.98$  consistently with the theory of Section 3.1, while at the

Electrophoresis

higher  $C_o$ , the monomer analogue particles become closer and the  $pH_M^*$ increases by reaching a value around 4.31, which is the  $pH_D^*$  value when the PPCR phenomenon is neglected while the IPCR phenomenon is considered. In this regard the  ${}_{P}H_{D}^{\ast}$  shift of the dimer analogue due to the PPCR phenomenon changes from 4.31 at infinity dilution to 4.66 at around  $\,C_{\rm o}\,{\approx}3.25$  mM as shown in Figure 3, which cannot convert to higher oligomers at pH<4 as indicated above. Further for  $d \rightarrow 0$  the contact of two monomers may produce a dimer and the merged particles are not anymore kinematically independent. Figure 3 also illustrates the estimations of distances between pair of monomer analogue particles when  $C_o$  increases. For  $C_o \approx 3.25$  mM a distance of around three particle diameters are obtained. In this regard, these calculations are simpler for the dimer (merged particles) than for the two monomers in near contact at high  $C_{o}$  where Eq. (2) may start to be invalid [4]. Full sets of numerical calculations related to the inclusion of the PPCR phenomenon are available in Supporting Information Tables 5 and 6 for a better analysis of Figures 2 and 3. Results illustrate that the dimerization of the A $\beta$ (12-28) peptide fragment is induced when a substantial average approach among the hydrated particles is achieved by increasing  $C_o$ . In this regard the PPCR phenomenon is relevant in this process, in opposition to what happens at infinity dilution, by introducing a  $pH_a^*$ shift toward the pI additional to that caused by the IPCR phenomenon. From

these approximate calculations it is also clear that the PPCR phenomenon deserves special consideration in the interaction between proteins and peptides in general. Therefore further research is needed to completely elucidate this subtle phenomenon.

Accepted Article

Finally it is important to point out that the nucleation of the dimerization process via the central hydrophobic region LVFFA is facilitated by the  $pH_{\alpha}^{*}$ shifts toward the pI as  $C_{o}$  increases, which partially reduce the interparticle electrostatic repulsion. Also one has to observe that the wild effective charge number  $Z_{w\alpha}$  is higher than the regulated effective charge number Z as depicted in Supporting Information Tables 2, 3, 5 and 6 [3, 4, 25, 26].

## **4 Concluding Remarks**

The effective electrophoretic mobility is one of the relevant transport properties of amyloid-beta peptides to be measured at appropriate BGE compositions with the purposed of studying the amyloidogenic process. The  $pH_{\alpha}^{*}$  shifts toward the peptide pI is a consequence of both the IPCR and PPCR phenomena enhancing the oligomerization of the A $\beta$ (12-28) peptide fragment via the hydrophobic region LVFFA studied here at pH 2.9. These phenomena may be important mechanisms causing the amyloidogenic process of amyloid beta peptides leading to the neuronal plaque formations at higher pH values by reducing repulsion electrostatic forces between chains. To deepen this aspect, additional researches involving CE experimental studies of these peptides are This article is protected by copyright. All rights reserved. required. Our forthcoming manuscript will show that higher order oligomerization of amyloid-beta peptides is consistent with the results found here involving the IPCR and PPCR phenomena.

#### Acknowledgements

Authors wish to thank the financial aid received from Universidad Nacional del Litoral, Santa Fe, Argentina (CAI+D-2011) and CONICET (PIP-112-201101-00060).

The Authors have declared no conflict of interest.

### **5** References

[1] Hearn, M. T. W., Keah, H. H., Boysen, R. I., Messana, I., Misiti, F.,

Rossetti, D. V., Giardina, B., Castagnola, M., Anal. Chem. 2000, 72, 1964-1972.

[2] Chiti, F., Dobson, C. M., Annu. Rev. Biochem. 2006, 75, 333-366.

[3] Deiber, J. A., Piaggio, M. V., Peirotti, M. B., *Electrophoresis* 2014, *35*, 755-761.

[4] Deiber, J. A., Piaggio, M. V., Peirotti, M. B., J. Sep. Sci. 2014, 37, 2618-2624.

[5] Blennow, K., Hampel, H., Weiner, M., Zetterberg, H., *Nat. Rev. Neurol.*2010, *6*, 131-144.

[6] Masters, C. L., Selkoe, D. J., Cold Spring Harb. Perspect. Med. 2012, 2: a006262.

[7] Piaggio, M. V., Peirotti, M. B., Deiber, J. A., *Electrophoresis* 2005, 26, 3232-3246.

[8] Piaggio, M. V., Peirotti, M. B., Deiber, J. A., *Electrophoresis* 2006, 27, 4631-4647.

[9] Piaggio, M. V., Peirotti, M. B., Deiber, J. A., *Electrophoresis* 2007, 28, 2223-2234.

[10] Piaggio, M. V., Peirotti, M. B., Deiber, J. A., *Electrophoresis* 2007, 28, 3658-3673.

[11] Peirotti, M. B., Piaggio, M. V., Deiber, J. A., J. Sep. Sci. 2008, 31, 548-554.

[12] Piaggio, M. V., Peirotti, M. B., Deiber, J. A., *Electrophoresis* 2009, *30*, 2328-2336.

[13] Mansfield, S. L., Jayawickrama, D. A., Timmons, J. S., Larive, C. K., *Biochim. Biophys. Acta* 1998, *1382*, 257-265.

[14] Piaggio, M. V., Peirotti, M. B., Deiber, J. A., J. Sep. Sci. 2010, 33, 2423-2429.

[15] Deiber, J. A., Piaggio, M. V., Peirotti, M. B., *Electrophoresis* 2012, *33*, 990-999.

[16] Šolínová, V., Kašička, V., Koval, D., Hlaváček, J., *Electrophoresis* 2004, 25, 2299-2308.

[17] Simó, C., González, R., Barbas, C., Cifuentes, A., Anal. Chem. 2005, 77,
7709-7716.

[18] Benavente, F., Balaguer, E., Barbosa, J., Sanz-Nebot, V., J. Chromatogr. A 2006, 1117, 94-102.

[19] Šolínová, V., Kašička, V., Sázelová, P., Barth, T., Mikšík, I., J.*Chromatogr. A* 2007, *1155*, 146-153.

[20] Allison, S. A., Pei, H., Allen, M., Brown, J., Chang-II, K., Zhen, Y., *J. Sep. Sci.* 2010, *33*, 2439-2446.

[21] Aptisa, G., Benavente, F., Sanz-Nebot, V., Chirila, E., Barbosa, J., Anal.Bioanal. Chem. 2010, 396, 1571-1579.

[22] Deiber, J. A., Piaggio, M. V., Peirotti, M. B., *Electrophoresis* 2011, *32*, 2779-2787.

- [23] Allison, S. A., Perrin, C., Cottet, H., *Electrophoresis* 2011, *32*, 2788-2796.
  [24] Wu, H. F., Allison, S. A., Perrin, C., Cottet, H., *J. Sep. Sci.* 2012, *35*, 556-562.
- [25] Deiber, J. A., Piaggio, M. V., Peirotti, M. B., *Electrophoresis* 2013, *34*, 700-707.
- [26] Deiber, J. A., Piaggio, M. V., Peirotti, M. B., *Electrophoresis* 2013, *34*, 708-715.
- [27] Deiber, J. A., Piaggio, M. V., Peirotti, M. B., *Electrophoresis* 2013, 34, 2648-2654.
- [28] Šolínová, V., Kašička, V., *Electrophoresis* 2013, 34, 2655-2665.
- [29] Allison, S. A., Wu, H., Bui, T. M., Dang, L., Huynh, G. H., Nguyen, T.,
- Soegiarto, L., Truong, B. C., J. Sep. Sci. 2014, 37, 2403-2410.
- [30] Kašička, V., *Electrophoresis* 2012, *33*, 48-73.
- [31] Selvaraju, S., El Rassi, Z., *Electrophoresis* 2012, 33, 74-88.
- [32] Righetti, P. G., Sebastiano, R., Citterio, A., Proteomics 2013, 13, 325-340.
- [33] Kašička, V., *Electrophoresis* 2014, 35, 69-95.

Accepted Artic

- [34] Ali, I., Al-Othman, Z. A., Al-Warthan, A., Asnin, L., Chudinov, A., *J. Sep. Sci.* 2014, *37*, 2447-2466.
- [35] Verpillot, R., Otto, M., Klafki, H., Taverna, M., J. Chromatogr. A 2008, 1214, 157-164.
- [36] Sabella, S., Quaglia, M., Lanni, C., Racchi, M., Govoni, S., Caccialanza,
  G., Calligaro, A., Bellotti, V., De Lorenzi, E., *Electrophoresis* 2004, *25*, 3186-3194.
- [37] Kato, M., Kinoshita, H., Enokita, M., Hori, Y., Hashimoto, T., Iwatsubo,T., Toyo'oka, T., *Anal. Chem.* 2007, *79*, 4887-4891.
- [38] Colombo, R., Carotti, A., Catto, M., Racchi, M., Lanni, C., Verga, L.,
- Caccialanza, G., De Lorenzi, E., Electrophoresis 2009, 30, 1418-1429.
- [39] Picou, R. A., Moses, J. P., Wellman, A. D., Kheterpal, I., Gilman, S. D., *Analyst (Cambridge, United Kingdom)* 2010, 135, 1631-1635.
- [40] Picou, R. A., Kheterpal, I., Wellman, A. D., Minnamreddy, M., Ku, G.,
- Gilman, S. D., J. Chromatogr. B 2011, 879, 627-632.
- [41] Verpillot, R., Esselmann, H., Mohamadi, M. R., Klafki, H., Poirier, F.,
- Lehnert, S., Otto, M., Wiltfang, J., Viovy, J. L., Taverna, M., Anal. Chem. 2011, 83, 1696-1703.
- [42] Pryor, N. E., Moss, M. A., Hestekin, Ch. N., Int. J. Mol. Sci. 2012, 13, 3038-3072.

[43] Verzola, B., Fogolari, F., Righetti, P. G., *Electrophoresis* 2001, *22*, 3728-3735.

[44] Kumar, S., Nussinov, R., ChemBioChem 2002, 3, 604-617.

[45] Fraser, P. E., Nguyen, J. T., Surewicz, W. K., Kirschner, D. A., Biophys. J.

1991, 60, 1190-1201.

[46] Allison, S. A., Macromolecules 1996, 29, 7391-7401.

[47] Allison, S. A., Biophys. Chem. 2001, 93, 197-213.

**Figure 1**. Monomer  $\mu_M$  (**•**) and dimer  $\mu_D$  (**□**) analogues and monomer and dimer mixture  $\mu$  (\*) effective electrophoretic mobilities as a function of initial monomer concentration  $C_o$ . Results are from the PLLCEM at pH 2.9, I = 1 mM and 25°C when the hydrated particle sampled is spherical with BGE slip. The IPCR phenomenon is considered only.





This article is protected by copyright. All rights reserved.

**Figure 2**. Monomer  $\mu_M$  (**n**) and dimer  $\mu_D$  (**D**) analogues and monomer and dimer mixture  $\mu$  (\*) effective electrophoretic mobilities as a function of initial monomer concentration  $C_o$ . Results are from the PLLCEM at pH 2.9, I = 1 mM and 25°C when the hydrated particle sampled is spherical with BGE slip. The IPCR and PPCR phenomena are considered.



Figure 2



This article is protected by copyright. All rights reserved.

**Figure 3.**  $pH_{\alpha}^{*}$  shifts toward pI of monomer (**n**) and dimer (**D**) analogues and distance between particles d ( $\circ$ ) as a function of initial monomer concentration  $C_{o}$  when the IPCR and PPCR phenomena are considered. Results are from the PLLCEM at pH 2.9, I = 1 mM and 25°C when the hydrated particle sampled is spherical with BGE slip.





This article is protected by copyright. All rights reserved.