Formulation matters! A Spectroscopic and Molecular Dynamics Investigation on the Peptide CIGB552 as Itself and in its Therapeutical Formulation.

Marco Savioli,^a Lorenzo Antonelli,^a Gianfranco Bocchinfuso,^a Francesca Cavalieri,^{a,d} Rita Cimino,^a Emanuela Gatto,^a Ernesto Placidi,^b Julio Raul Fernandez Masso,^c Hilda Garay Perez,^c Hector Santana,^c Maribel Guerra-Vallespi,^c Mariano Venanzi^{a,*}

^a Dept. of Chemical Sciences and Technology, University of Rome Tor Vergata, Via della Ricerca Scientifica 1, 00133, Rome, Italy.

- ^b Dept. of Physics, University of Rome 'Sapienza', P.le A. Moro 5, 00185 Rome, Italy.
- ^c Center for Genetic Engineering and Biotechnology, PO Box 6162, 10600, Havana, Cuba.
- ^d School of Science, RMIT University, Melbourne, VIC 3001, Australia.

*<u>venanzi@uniroma2.it</u>

Abstract

Synthetic therapeutic peptides (STP) are intensively studied as new-generation drugs, characterized by high purity, biocompatibility, selectivity and stereochemical control. However, most of the studies are focussed on the bioactivity of STP without considering how the formulation actually used for therapy administration could alter the physico-chemical properties of the active principle. The aggregation properties of a 20-mer STP (Ac-His-Ala-Arg-Ile-Lys-D-Pro-Thr-Phe-Arg-Arg-D-Leu-Lys-Trp-Lys-Tyr-Lys-Gly-Lys-Phe-Trp-NH₂), showing antitumor activity, were investigated by optical spectroscopy and Atomic Force Microscopy imaging, as itself (CIGB552) and in its therapeutic formulation (CIGB552TF). It has found that the therapeutic formulation deeply affects the aggregation properties of the investigated peptide and the morphology of the aggregates formed on mica by deposition of CIGB552 and CIGB552 aggregation under physiological ionic strength conditions (NaCl 150 mM), showing that peptide oligomers, from dimers to tetramers, are preferentially formed in this environment. Interestingly, cell viability assays performed on H-460 cell lines indicate a major antiproliferative activity of the peptide in its therapeutic formulation with respect to the peptide aqueous solution.

1. INTRODUCTION

Despite the relevant progress in the prevention and therapy of tumors, cancer diseases remain one of the most important causes of death. Chemotherapy drugs currently in use target quite successfully tumor cells, but, unfortunately, are biased by deleterious side effects on healthy cells and tissues. This dramatic situation urges for the search of a new generation of drugs, and those based on synthetic therapeutic peptides (STP) represent one of the most promising sources.¹ STP possess the distinct advantage of being biocompatible, highly selective, and obtainable with good enantiomeric purity and stereochemical control applying well-established synthetic procedures.² Design of new STP aims to optimize their pharmacokinetics in terms of bioavailability, biodegradability and affinity for receptors or target cells.^{3,4}

Among the many promising applications for biotherapy, STP have been proposed as potential antitumor agents for their fast diffusion through biological tissues and enhanced cell permeabilization properties. Furthermore, peptide drugs have shown potent antineoplastic activity interfering with several molecular mechanisms involved in the carcinogenesis, inhibiting angiogenesis or blocking protein-protein interactions.⁵ Peptide receptor antagonists featuring pro-apoptosis properties have also been investigated for limiting the growth of tumors.⁶

Recently, a new therapeutic peptide, derived from *Limulus sp*, and based on the amphipathic cyclic pentapeptide LALF₃₂₋₅₁, denoted as L2, was investigated as a novel antitumor peptide.⁷ With the aim to improve the antitumor activity of L2, some of us from the Center of Genetic Engineering and Biotechnology (CIGB) of Havana (Cuba) synthesized a new linear peptide analog of L2, denoted in the following as CIGB552.⁸ L2 was modified by including a proline (P(6)) and a leucine (L(11)) residues in the peptide sequence as D-amino acids, and an acetyl (Ac) group at the N-terminus.

These modifications were shown to improve the antitumor activity of L2, increasing its apoptosis efficiency and its capacity to inhibit cell proliferation.⁸ CIGB552 has also proven to be effective in reducing the size of tumor and increasing lifespan in tumor-bearing mice,⁹ and in dogs with naturally-occurring cancer.¹⁰ It has also been found that CIGB552 decreased the microvessel density in the HT-29 xenograft human, suggesting potent anti-angiogenesis properties.⁹

The unique capabilities of CIGB552 arise from its cell-penetrating properties, which allow it to enter cells and elicit a pro-apoptotic effect through its major mediator, the COMMD1 protein,^{11,12} a pleiotropic factor involved in the regulation of many cellular and physiological processes that include oxidative stress, protein trafficking, NF- κ B-mediated transcription and oncogenesis.

Recently, it has been demonstrated that endocytosis is a preferred mechanism for CIGB552 internalization, which in turn favours the interaction between the peptide and COMMD1 in the

endosomal compartment, as proved by proteomics and genomics experiments.¹³⁻¹⁵ It has also been shown that the ability of CIGB552 to negatively modulate NF-kB and HIF-1 pathways is impaired in the COMMD1 Knock-out NCI-H460 cell line, confirming that COMMD1 is essential for the peptide mechanism of action.

It should be noted that in the final pharmaceutical formulation, the active substance, currently in clinical phase 2, is administrated with excipients, that usually do not explicate a direct therapeutic activity, but protects the active substance from chemical effectors and facilitates its pharmacokinetics.

In this work, we compared the physico-chemical properties of CIGB552 in water with those of the same peptide in a formulation mimicking drug administration conditions, containing α , α -trehalose and (L)-tartaric acid as excipients. The latter system will be denoted in the following as CIGB552TF for brevity, where the suffix TF denotes CIGB552 in its therapeutic formulation. The molecular formula of CIGB552 (Ac-His-Ala-Arg-Ile-Lys-D-Pro-Thr-Phe-Arg-Arg-D-Leu-Lys-Trp-Lys-Tyr-Lys-Gly-Lys-Phe-Trp-NH₂) is reported in Scheme 1. The presence in the peptide chain of three intrinsically fluorescent residues, i.e. W(13), Y(15) and W(20), allowed us to apply optical spectroscopy techniques to characterize the aggregation process of CIGB552 in water and in the therapeutic formulation. Atomic Force Microscopy imaging experiments and Molecular Dynamics simulations were also carried out to investigate on the morphology of the aggregates and on the structure of the small peptide clusters nucleating the growth of large peptide aggregates, respectively.



Scheme 1. Molecular formula of the therapeutic peptide investigated (CIGB552)

2. MATERIALS AND METHODS

2.1 Materials. CIGB552 was synthesized at the Center for Genetic Engineering and Biotechnology (Havana, Cuba). Details of the synthesis has already been reported elsewhere.⁹ Briefly, CIGB552 was obtained by solid-state synthesis using the Fmoc strategy, and purified by RP-HPLC inverse

(purity>95%) under an acetonitrile/trifluoroacetic acid gradient. The peptide was characterized by electrospray mass spectrometry (Micromass, UK) and NMR spectroscopy. A CIGB552 formulation (CIGB552TF), containing 32.1% (w/w) CIGB552, 50.7% α , α -trehalose, and 17.2% (L)-tartaric acid, was also prepared for mimicking drug administration conditions. All the solutions for spectroscopy and microscopy experiments were prepared from a 0.28 mM peptide stock solution at pH 7.2.

2.2 Spectroscopy. UV-Vis absorption measurements were carried out at room temperature by a Cary100 Scan spectrophotometer, using 3x3 mm quartz cuvettes (Hellma). Steady-state fluorescence measurements were carried out by a Fluoromax-4 (HORIBA, Jobin Yvon), using a 3x3 mm quartz cuvette at room temperature. Fluorescence emission and excitation spectra were measured at the excitation wavelengths of 280 and 295 nm and emission wavelengths of 360 and 420 nm, respectively.

2.3 Atomic Force Microscopy (AFM). Atomic Force Microscopy experiments were carried out in air at room temperature on a Veeco Multiprobe IIIa instrument (Santa Barbara, CA). The AFM measurements were carried out in tapping mode on a film obtained by drop casting 1 mM aqueous solutions of CIGB552 or CIGB552TF on mica, and incubating for 18 hours in a dryer. For the AFM measurements, a Si super sharp tip, functionalized by carbon flake was used (curvature 1 nm, elastic constant 5 N/m, resonance frequency 150 kHz).

2.4 Molecular Dynamics (MD) simulations. MD simulations were performed using the GROMACS package version 2019.6¹⁶ with a modified GROMOS96 53a6-FF ¹⁷ force field including parameters for D-Pro and D-Leu residues. In order to analyse the peptide conformational landscape and aggregation properties, MD simulations with 1 and 4 CIGB552 molecules were carried out in 150 mM NaCl aqueous solution (MD1A, MD4A) and in water (MD1B, MD4B) (Table 1). Time steps of 2 fs were used for equilibration and production runs. The equilibrium temperature (300 K) was controlled by velocity rescaling ¹⁸ with a coupling constant of 0.1 ps.

Pressure was controlled by an isotropic Parrinello-Rahman barostat ¹⁹ with a coupling constant of 2 ps and a reference external pressure of 1 atm. Lennard-Jones long range interactions were treated with a cut-off radius of 1.2 nm. Coulombic interactions were calculated by the Particle Mesh Ewald method.²⁰ 200 ns-long simulations have been performed for three replicas of each system having different starting configurations. MD simulation analyses were carried out using either built-in GROMACS tool and handmade Tcl script in VMD.²¹ Relative solvent accessibility (RSA) was calculated as the ratio between the solvent accessibility surface area (SASA) value and the maximum solvent accessible surface area for each residue.²² Secondary structure content was assessed through the DSSP (Dictionary Secondary Structure of Proteins) algorithm.²³ SASA and

gyration radii were used to evaluate the equilibration time of each MD. RSMD-based cluster analysis was carried out applying similarity conditions with a 0.2 nm cut-off.²⁴

2.5 Cell viability assay. Cell viability assay was carried out on the H-460 (ATCC, HTB-117) cell line, maintained at 37 °C and 5% CO₂ in RPMI 1640, and supplemented with 10% fetal bovine serum (HyClone, Logan, UT) plus 30 μ g/ml of gentamycin (Gibco). Proliferative assay was performed as previously described.⁷ Peptide concentrations ranging from 0 to 300 μ M were added to 10,000 cells/well and incubated for 48 h at 37 °C in 5% CO₂. Subsequently, 50 μ l of 80% trichloroacetic acid solution were added to the cells and incubated for one hour at 4 °C. Finally, 100 μ l of the sulforhodamine B solution were added to the wells, and absorbance was measured at λ =492 nm. Each sample point was carried out in triplicate, and experiments were performed twice. IC50 values were obtained from the respective growth curves.

3. RESULTS AND DISCUSSION

3.1 UV-Vis Absorption

The absorption spectrum of CIGB522 in the near UV region is determined by the overlap of the $n \rightarrow \pi^*$ transition of the Trp [λ_{max} = 283 nm; $\epsilon(283)$ = 5800 M⁻¹cm⁻¹] and Tyr residues [λ_{max} = 274 nm; $\epsilon(274)$ = 1400 M⁻¹cm⁻¹] comprised in the peptide sequence. For comparison, the absorption spectra of CIGB522TF in the concentration range between 5.2 and 280 µM are reported in Figure 1A. From these data, we determined the molar extinction coefficient of CIGB52FF at 280 nm [$\epsilon(280)$ = 12500 M⁻¹cm⁻¹], in good agreement with the sum of the molar extinction coefficients of the aromatic residues reported above. This finding indicates that strong (short-range) ground-state interactions between the aromatic moieties of CIGB552 in its therapeutic formulation can be neglected. It should be noted that the shapes of the absorption spectra of CIGB552 in its therapeutic formulation and in solution are strictly the same (data not shown). On the contrary, while for the former the measured absorbances were found to closely follow the linear trend forecast by the Lambert-Beer law, for the latter a marked deviation from the linear behavior is observed at the higher concentrations (Figure 1B), suggesting that CIGB552 in water may give rise to the formation of aggregates at micromolar (> 50 µM) concentrations.



Figure 1. UV absorption experiment on CIGB552. A) Absorption spectra of CIGB552TF (therapeutic formulation) at different concentrations (from 5.2 to 280 μ M); B) Absorbance at λ =280 nm as a function of the peptide concentration. Red dots: CIGB552; black squares: CIGB552TF.

3.2 Fluorescence

Fluorescence emission spectra of CIGB552 and CIGB552TF were measured at $\lambda_{ex}=280$ (both Tyr and Trp photoexcited) and at $\lambda_{ex}=298$ nm (selective photoexcitation of Trp). For both the excitation wavelengths, the fluorescence emission spectra of CIGB552 in water and in the therapeutic formulation, show a relatively broad emission band centred at $\lambda_{em}=359$ nm, typical of the indole emission in a polar environment (Supplementary Information, Figure SI1). The close similarity of the emission spectra measured at $\lambda_{exc}=280$ and 298 nm indicates that an almost total Tyr \rightarrow Trp energy transfer takes place for both the CIGB552 formulations.

Emission spectra for different concentrations of CIGB552TF at λ_{ex} = 298 nm are shown in Figure 2A. Where the emission intensities of CIGB552 and CIGB552TF at λ_{em} = 359 nm are reported as a function of the peptide concentration, the same intensity vs. concentration trend observed by UV absorption measurements is obtained. In particular, in the case of CIGB552TF, it has found a linear dependence of the emission intensity on the peptide concentration, while for the higher concentrations (> 100 µM) of the peptide water solutions an asymptotic trend is obtained (Figure 2B). The same trend was observed for λ_{exc} = 280 nm (data not shown). These results, in parallel with the UV absorption findings, strongly suggest that CIGB552, but not CIGB552TF, gives rise to the formation of aggregated species at the higher concentrations investigated.



Figure 2. Fluorescence intensity (λ_{ex} =298 nm; λ_{em} =359 nm) as a function of the peptide concentration (from 5.2 to 280 μ M). Red dots: CIGB552; black squares: CIGB552TF (therapeutic formulation).

3.3 Atomic Force Microscopy (AFM)

AFM experiments carried out on peptide films obtained by deposition and overnight drying of CIGB522 and CIGB522TF millimolar solutions on mica revealed interesting differences in the morphologies of the aggregates formed on the solid substrate. In particular, AFM imaging of CIGB552TF showed nanostructures of globular morphology typical of an aggregation process driven by hydrophobic effects (Figure 3A).^{25,26} These globular structures are characterized by a relatively large size distribution, with diameters ranging from 10 to 50 nm. The importance of hydrophobic effects is emphasized by the AFM image reported in Figure 3B, in which the peptide was seen to form, in some limited regions of the mica surface, micrometric toroidal structures, characterized by an outer diameter of 990 \pm 10 nm and an inner diameter of 56 \pm 9 nm.



Figure 3. AFM images of 1 mM CIGB552TF (therapeutic formulation) aqueous solutions deposited on mica.

Interestingly, AFM imaging of CIGB552, obtained from deposition of peptide aqueous solutions, (Figure 4) revealed the formation of fibril structures, characterized by nanometric lengths $(377\pm112 \text{ nm})$, and thicknesses $(26 \pm 6 \text{ nm})$.



Figure 4. AFM images of 1 mM CIGB552 aqueous solutions deposited on mica. Figure B clearly shows the alignment of the transient globular structures nucleating the growth of peptide fibrils.

The structures reported in Figure 4B highlights the mechanism of formation of fibril-like aggregates, involving the coalescence and directional growth of nanosized elongated peptide clusters. It has been shown that the formation of such structures is favoured when directional interactions, like hydrogen bonding and π - π stacking of aromatic groups, predominate with respect

to nonspecific driving forces, like, entropically-driven hydrophobic effects.²⁷ The AFM images reported in Fig. 4 clearly spotted nanometric globules coalescing in fibrillar structures of micrometric length, in close analogy with the formation mechanism of peptide amyloids responsible for several neurodegenerative diseases.²⁸

These results clearly agree with the idea that in the therapeutic formulation CIGB552 is well dispersed within the α,α -trehalose and (L)-tartaric acid matrix, in agreement with spectroscopic results. In contrast, the fibril structures imaged by AFM experiments on CIGB552 deposited on mica from aqueous solutions could be formed only by closely interacting peptide chains, that was only possible in the aqueous peptide formulation.

3.4 Molecular Dynamics

MD simulations have been carried out to investigate on the conformational and aggregation properties of CIGB552 mimicking physiological conditions. In particular, MD simulations were carried out for 1 and 4 CIGB552 molecules in the presence of NaCl, corresponding to 150 mM NaCl aqueous solutions (MD1A, MD4A), and in water (under low ionic strength conditions) (MD1B, MD4B). The aim is to analyze the conformational landscape of a single peptide chain and the formation of the small peptide clusters characterizing the first steps of peptide aggregation, experimentally observed in the absence of the excipients present in the therapeutic formulation.

RMSD-based cluster analyses (CA) on MD1A provided for the CIGB552 monomer two predominant cluster-averaged structures, the most representative of which are shown in Figures 5. These clusters account for 51.5% and 28.6% of all sampled conformations, respectively. Interestingly, in the most abundant conformer (Figure 5A), the 20-mer peptide folds into a rather compact structure, characterized by the presence of several antiparallel β -sheet segments.

Centre-of-mass distance distribution analysis revealed that for more than 30% of the sampled conformations, W13 indole and R9 guanidinium are located less than 5 Å away, suggesting the occurrence of a cation- π interaction between the two sidechain groups (Figure SI2).²⁹ This finding is confirmed by the analysis of the relative solvent accessibility area (RSA),²² which drastically decreases to 40% of the maximum allowed accessible surface for the aforementioned residues. This interaction may contribute to stabilize the rather compact structure of conformer A (Fig. 5), and it could explain the observed Trp red-shift emission typical of a polar environment.³⁰



Figure 5. Most representative structures of CIGB552 from cluster analysis (0.2 nm cut-off) of MD1A simulations in NaCl aqueous solution. Abundances: 51.5% (A); 28.6% (B). The backbone is reported as a ribbon, with the β -sheet tracts coloured in yellow. The side chains of Arg, Trp and Tyr residues are reported as sticks coloured in blue, lime and orange, respectively. A parallel-shaped cation- π interaction between R9 and W13 can be clearly spotted in the most abundant conformer (A).

Analysis of the distances between the Y15-W13 and Y15-W20 sidechains in MD1A (Figure SI3) reveals that for 80% of the former and 70% of the latter pair, the phenol/indole separation distances are definitely shorter than their characteristic Förster distance (9-16 Å). This finding is in close agreement with the fluorescence results reported above, showing an almost complete Tyr \rightarrow Trp Energy Transfer.

Using the same procedure applied for MD1A, also the three 200 ns-long trajectories of MD4A were joined and considered as a whole. The obtained ensemble was first analyzed to investigate on the CIGB552 aggregation propensity. In Table 2, the frequency distribution of CIGB552 oligomers is reported, in terms of both the oligomer occurrence and the probability of a single peptide unit to be part of a given oligomer. For comparison, the results of MD simulation in water (MD4B) are also reported in Table 2.

From these data, it appears that the formation of peptide oligomers in NaCl is predominant with respect to the presence of monomeric species, indicating a net propensity of CIGB552 to form small peptide clusters in this environment.

Interestingly, while in MD4A the monomer/oligomers ratio amounts to 1:2, with a remarkable presence of tetramers, MD4B (low ionic strength conditions), shows a comparable monomer/oligomer proportion, and a negligible content of tetrameric structures (Figure SI4). This finding suggests that hydrophobic effects, notoriously enhanced with increasing ionic strength, are

important for CIGB552 aggregation. In agreement with these considerations, RSA analysis showed that, in the tetramer, the solvent accessibility sensibly decreased in the peptide regions rich of aromatic residues.

The secondary structure content of the CIGB552 monomer (MD1A, MD1B) and tetramer (extracted from MD4A and MD4B) was analyzed by the DSSP algorithm.²³ In all MD simulations CIGB552 populates a rich variety of secondary structures, the abundances of which are summarized in Table 3. From these data, it appears that the conformational landscape of CIGB552 in NaCl is essentially the same in the monomeric and tetrameric structures, with a slight increase of coil and β -sheet conformations at expenses of decreasing bend and turn conformations in the tetrameric structures. As regards the secondary structure content, no significant differences were found for both monomeric and tetrameric species between MD simulations carried out in NaCl and in water.

It should be noted that, although helical conformations are sometimes evocated as intermediates in the self-assembly early stages of amyloid fibrils,³¹ our analysis does not reveal the presence of helical structures in both CIGB552 monomers and tetramers, suggesting that these conformations do not play a significant role in the aggregation mechanism of this peptide.

Cluster analysis of the tetrameric structures extracted from the whole MD4A sampling time, applying the same similarity conditions used for MD1A (0.2 nm cut-off), provided three predominant cluster-averaged structures of comparable abundance, i.e. 34.5%, 28.7% and 22.2%, respectively, the most representative of which are shown in Figure 6.



Figure 6. Cluster average structures of CIGB552 tetramers from MD4 simulation in 150 mM NaCl. Abundances: 34.5% (A); 28.7% (B); 22.2% (C). Backbone: ribbon (β -sheet tracts in yellow.); Arg: blue; Trp: lime; Tyr: orange.

Figure 6 clearly reveals the presence of stretched peptide aggregates, even if more compact configurations also appear with comparable proportion (Figure 6B). At a glance, the interaction between the β -sheet tracts (an intrinsically directional interaction) seem to play an important role in the stabilization of the aggregate.

In this regard, one might ask about the role of π - π interactions among the four Tyr and eight Trp residues in the stabilization of CIGB552 tetramers. In Figure 7 we reported the separation distance distribution obtained by summing over the 32 Tyr-Trp pair distances for all the tetramers, showing up during the MD4A simulation time.

Similar to MD1A (Figure SI3), two well-defined peaks at 6 and 9 Å, respectively, fall well below the Tyr-Trp Förster distance, accounting for the very efficient energy transfer observed at all the peptide concentrations investigated. Interestingly, the maxima of the Tyr-Trp distance distribution are regularly spaced by 4 Å, a typical distance between aromatic groups stabilized by stacking interactions.³² The observed regularity of the separation patterns extends over distances longer than 2 nm, a clear indication of the stability of tetrameric structures.

In Figures 7 we partitioned the distance distribution between intra- and inter- peptide Tyr-Trp pairs in the tetrameric structures. This partition shows that distances comprised within 1 nm are mostly accounted for by intra-peptide interactions, and are almost absent in the inter-strand component of the distance distribution. These data strongly suggest that π - π interactions stabilize the single chain conformation in the oligomers but do not contribute to the inter-peptide interactions.

This conclusion is strengthened by taking into account the 6 Tyr-Tyr (Figure SI5) and 56 Trp-Trp (Figure SI6) distance distributions. The former represents a topological parameter that depends only on inter-peptide interactions. In this case, a regular pattern of distances was observed (Figure SI5), characterized by two distribution peaks centered at around 1.3 and 1.8 nm, and the absence of short distances contributions. The same pattern was observed for both the intra- and inter-strand contributions to the Trp-Trp distance distribution.,



Figure 7. Distance distribution of Tyr-Trp pairs in the tetrameric structures from MD4A simulation. A) Black: sum over all 32 Tyr-Trp distances; blue: Intra-peptide distance distribution; red: Interpeptide distance distribution.

To highlight the inter-chain interactions stabilizing the aggregate, we have analyzed the hydrogen bonding (HB) network in the CIGB552 tetramer. Where the HB connectivity map is considered, three very stable (30-50% of the simulation time) and a dozen of quite stable (10-30% of the simulation time) inter-peptide HB contacts have found to stabilize the peptide tetramer (Figure SI7). Overall, these findings suggest that aromatic-aromatic interactions are important to stabilize the secondary structure ordering of the single peptide units, that form relatively stable oligomers, maintained by hydrophobic interactions and by a long-lasting inter-peptide HB network.

Interestingly, in all the structures reported in Figures 5 and 6, most of the charged peptide sidechains (i.e., the four Lys and three Arg residues) protrude outwards into the solvent, providing the peptide surface of a net positive charge character. This finding may explain the strong cellpenetrating properties of CIGB552 in its monomeric form, as well as small peptide oligomers.

3.5 Cell viability assay

In order to study the cytotoxic activity of the investigated peptide in aqueous solution and in its therapeutical formulation, a cell viability assay was performed on H-460 cell line. IC₅₀ values were obtained by fitting the experimental data, i.e., the optical densities (OD) of sulforhodamine B at λ =492 nm at different peptide concentration, by the Hill equation:³³

$$OD(x) = OD_{max} - \frac{OD_{max} - OD_{min}}{1 + \left(\frac{x}{IC_{50}}\right)^H}$$

Where x represents the peptide concentration and H the phenomenological Hill number. Figure 8 shows the result of a typical experiment of cell viability assay for the two peptide formulations investigated.

Interestingly, CIGB552TF shows IC_{50} values definitely lower than the aqueous peptide formulation (Table 4). This result strengthens the idea that the formulation of the peptide, and therefore its aggregation state, is relevant for its biological activity.



Figure 8. Cell viability assay of CIGB522 (red dots and line) and CIGB522TF (black square and line) on H-460 cell line. The optical density of sulforhodamine B at λ =492 nm was reported as a function of the peptide micromolar concentration (log scale).

4. CONCLUSIONS

The aggregation properties of CIGB552 in water and in its therapeutic formulation were investigated by optical spectroscopy and microscopy technique with nanometric resolution. UV-Vis and fluorescence experiments showed that the excipients, i.e. 50.7% α , α -trehalose, and 17.2% (L)-tartaric acid, strongly deplete the peptide aggregation propensity in the therapeutic formulation, maintaining the peptide in its monomeric form. This conclusion is strengthened by AFM experiments that showed that the peptide in its therapeutic formulation formed almost exclusively globular aggregates, a morphology typical of aggregation processes driven by hydrophobic effect.

The role of hydrophobic effects was highlighted by the comparison of MD simulations carried out in water and under ionic strength conditions mimicking physiological conditions

On the contrary, AFM imaging revealed the fibrillization of aggregates formed from deposition of aqueous CIGB552 solution. In this regard, it is important to stress that formation of peptide fibrils necessitates of specific peptide-peptide interaction.^{34,35} MD simulations suggest that an inter-strand HB network concurs to stabilize peptide clusters, and these interactions could be the basis of the clear directionality observed in the aggregates.

All these findings strongly suggest that trehalose-CIGB552 interactions may take place in the therapeutic formulation, inhibiting peptide fibrillogenesis. Trehalose-peptide conjugates have already been shown to act as inhibitors of amyloid fibrillation,³⁶ enhancers of resistance toward proteolytic degradation,³⁷ protectants against thermal denaturation of proteins.³⁸

These ideas pave the way for explaining the higher cytotoxic activity of CIGB552 in its therapeutic formulation, suggesting that peptide fibrillation may inhibit its therapeutic activity. On the other hand, control of the morphology of peptide nanostructures could represent a suitable approach for modulating the efficiency of therapeutic peptides through fine tuning of their pharmacokinetics.

Supporting Information. Fluorescence spectra of CIGB552 in water and in its therapeutic formulation at λ_{ex} =280 and 298 nm; W13-R9, Y15-W13, Y15-W20 distance distributions from MD1A simulations; distribution of CIGB552 aggregates, Y-W, Y-Y and W-W distance distributions from MD4A simulations.

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| MD simulation | Box dimensions (nm) ³ | Number of CIGB552 molecules | Number of water molecules | Number of Na ⁺ ions | Number of Cl ⁻ ions |
|------------------|--|-----------------------------------|---------------------------------|-----------------------------------|-----------------------------------|
| MD1A | 5.44x5.44x5.44 | 1 | 5005 | 15 | 23 |
| MD1B | 5.44x5.44x5.44 | 1 | 5035 | - | 8 |
| MD4A | 4.98x4.98x7.96 | 4 | 5964 | 18 | 50 |
| MD4B | 4.98x4.98x7.96 | 4 | 5996 | - | 32 |

Table 1. Experimental conditions for MD simulations at T=300 K with 1 and 4 CIGB552 molecules in 150 mM NaCl aqueous solution (MD1A and MD4A) and in water (MD1B, MD4B).

| Oligomer | Oligomer occurrence (%) | | Peptide probability (%) | |
|----------|-------------------------|------|-------------------------|------|
| | MD4A | MD4B | MD4A | MD4B |
| Monomer | 34.8 | 47.1 | 16.3 | 25.6 |
| Dimer | 30.1 | 23.2 | 28.2 | 25.3 |
| Trimer | 22.1 | 28.7 | 31.3 | 46.9 |
| Tetramer | 13.0 | 1.0 | 24.4 | 2.2 |

Table 2. CIGB552 oligomer and peptide distribution from MD4 simulation in NaCl 150 mM (MD4A) and in water (low ionic strength) (MD4B).

| Secondary Structure | MD1A(%) | MD1B (%) | MD4A (%) | MD4B (%) |
|---------------------|---------|----------|----------|----------|
| Coil | 39.5 | 41.9 | 43.5 | 42.2 |
| β-Sheet | 19.9 | 15.8 | 21.4 | 20.0 |
| β-Bridge | 4.5 | 3.8 | 6.0 | 3.2 |
| Bend | 22.8 | 28.2 | 19.5 | 22.1 |
| Turn | 13.4 | 10.3 | 9.5 | 12.5 |

Table 3. Percentage of secondary structures adopted by CIGB552 for 600 ns MD simulations in 150mM NaCl (MD1A, MD4A) and in water (MD1B, MD4B).

Table 4. Effect of the peptide therapeutic formulation (CIGB552TF) and aqueous solution (CIGB552) on the cell viability in H-460 tumor cell line.

| | CIGB552TF | CIGB552 |
|----------------------------------|-----------|---------|
| H-460 IC50 (µMol/l) ^a | 58 ±3 | 94±11 |
| | | |

^aMean \pm SD of three determinations on three different peptide lots.