Endogenous modulators of neurotrophin signaling: Landscape of the transient ATP-NGF interactions

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1. Introduction

The NGF is the first discovered member of the NT family [1]. It is essential for the development and phenotypic maintenance of neurons in central and peripheral nervous systems. It has a crucial role in activation of the immune and endocrine systems and in the pain pathway [2]. Mature NGF is obtained by furin protease cleavage from the precursor proNGF [3]. NGF binds with high affinity to the tyrosine kinase TrkA receptor, determining cell survival and differentiation, as well as to p75NTreceptors triggering apoptosis [4,5]. proNGF creates a signaling complex by simultaneously interacting with p75NT and a member of the family of Vps10p-domain receptors, Sortilin [6].

Translational and clinical research highlighted the spectrum of diseases that could benefit from NGF treatment (e.g. Alzheimer’s Disease, diabetic skin wound healing, corneal pathologies, etc.) [7]. Its therapeutic application has been limited owing to poor plasma stability, restricted nervous system penetration and, importantly, the pleiotropic actions deriving from its concomitant binding to multiple receptors. One strategy to overcome these limitations is to target NGF and/or individual NGF receptors with small molecule ligands. The latter might also modulate various aspects of the involved signaling pathways in distinct ways from the native NTs. These ligands might provide novel therapeutic options for a broad range of neurological indications [2,8–12].

There are very few reports, which explored the role of essential endogenous ligands as modulators of NGF biological activity [13–19]. The ability of DNA, RNA, oligonucleotides, lipids and glycosaminoglycans to gate the facilitatory action of NGF on its signaling pathways has been seldom considered [13–19]. Thus, a deeper understanding of the molecular determinants of the ligands...
binding mode is required to better appreciate and potentially exploit the therapeutic potential of NGF.

Recently, ATP binding has been reported for protein families, which lack the ATP-binding domain. It has been proven that NGF binds to ATP and that the formed complex protects hippocampal neural cells from death [20–23]. The ligand signature and the physiological role of the extracellular ATP-NGF interrelationship remain however to be elucidated [21,22]. Similar observations were reported for another member of the NT family, BDNF [24], as well as for other growth factors (FGF [25], Vascular Endothelial Growth Factor [26]). The ATP-NGF association and the role of the concomitant presence of divalent ions have been analyzed by MS [21]. It was suggested by analogy to other ATP-binding proteins that ATP might bind NGF at a putative heparin-binding site [22].

ATP metabolism, in different cellular contexts, is regulated by divalent cations, among which, the role of Mg2+ and Zn2+ is of particular interest [27–29]. Zn2+ binding is characterized by a tightly regulated homeostasis and its link to cytotoxicity has been proven [27–29]. Zn2+ interaction with NTs has been reported [30,31].

It is well known that the release of ATP into the brain is finely tuned [32,33]. Different extracellular ATP levels in brain have been detected in health or disease conditions, and related to the activation of specific subclasses of P2 receptors [32,33]. A set of crucial cellular studies, investigating the ATP neurotrophic role in synergy with NGF highlighted a crosstalk between neurotrophins signaling and the purinergic receptors system, in different disease states [34,35]. The relative extracellular levels of ATP and NGF were reported to be involved in the interaction with their specific receptors. The imbalance in the extracellular levels of both ATP and NGF would be a key factor into the cellular response to cell damage. ATP was reported to have a role into PC12 cells activation, synergistically with NGF [35]. However, the lack of knowledge on the structural determinants and of the thermodynamic fingerprints of the ATP-NGF interactions, represent a major drawback towards the understanding of the underlying functional and biological properties of the complex.

We carried out a thorough structural/biophysical study including state-of-the-art solution NMR, integrated by extensive MD simulations aimed at investigating the interaction of ATP with rhNGF. For the characterization of ATP-rhNGF binding, we used 2D HSQC experiments, the most robust NMR method for detecting and evaluating weak ligand–protein binding. This technique, which was originally exploited for detecting weak small fragments binders to protein targets by using the so-called SAR by NMR approach [36]. Since then it is one of the most important fragment screening methods, not only because of its sensitivity in detecting weak interactions (Kd in the μM to mM range), but also it is frequently used as validation tool to confirm specific binding [37–39]. The cartography of the interactions between ATP and rhNGF unveiled the binding sites on the molecular surface and the ligand’s orientation. An SPR analysis allowed investigating the functional role of this interaction in the context of TrkA and p75NTR receptors. The functional role of divalent ions in conjunction with ATP was also exploited. The present structural analysis pinpoints ATP as a genuine endogenous modulator of NGF signaling in vivo, in health and disease conditions.

2. Materials and methods

2.1. Expression, refolding and purification of rhNGF

rhNGF was recombinantly expressed according to well-established protocols which exploit the in vitro refolding and subsequent proteolytic cleavage from its precursor rh-proNGF. rhNGF was expressed in Rosetta (DE3) E. Coli cells, transformed with the pET11a plasmid containing the cDNA of human proNGF VSAR. The mutation RSKR – VSAR at furin cleavage site allowed the introduction of a specific trypsin cleavage for the preparation of mature rhNGF (US Patent 2015/0087020). Solubilization and refolding of rh-proNGF from inclusion bodies was carried out according to published protocols [40]. Mature rhNGF was obtained by controlled trypsin cleavage of rh-proNGF [41]. Optimization of the expression of rh-proNGF VSAR in minimal medium with 15N- or 13C,15N-labeling was needed, starting from the conditions described for the recombinant mouse proNGF [41]. The best expression conditions were obtained by supplementing M9 medium with 0.4% glucose and upon induction with IPTG for 18 h at 25 °C. The integrity of rhNGF was confirmed by MALDI-MS analysis. Protein fold was characterized by biophysical techniques (DSF, FT-IR, NMR).

2.2. NMR experiments

The high-resolution spectra were recorded on Agilent Technologies DD2 600 MHz and VNMRS 800 MHz Spectrometers, using cryoprobes, at 30 °C.

2.2.1. NMR experiments for rhNGF spectral assignment

The HNCA, 15N NOESY-HSQC, 13C NOESY-HSQC spectra were acquired at 800 MHz on samples containing 15N- or 15N,13C-labelled rhNGF (0.1 mM) in 50 mM Hepes at pH 7 and 10% D2O. HNCA experiment was employed to obtain sequence specific 13Cα backbone assignments [42]. 15N NOESY-HSQC, 13C NOESY-HSQC were used for side chain aliphatic assignments. All spectra were processed using NMRPipe/NMR-Draw [43] and analyzed using CARA [44]. Intra-molecular proton distance restraints were derived from 15N- and 13C-NOESY-HSQC spectra (mixing time 100 ms). Backbone assignment was guided by the identification of the expected set of connectivities within and inter-strands, while side chains assignment was complicated by the severe overlap of resonances from highly abundant Leu, Ile, Lys, Arg residues. Backbone and side chain assignment was completed at 82% and 74%, respectively, and deposited in the BMRB database (ID: 34515).

2.2.2. Structure determination of rhNGF

Automated NOE cross-peak assignments and structure determination were performed using ARIA 2.3 software [45,46]. The NMR structural bundle was generated by ARIA using as input the ambiguous and unambiguous intra-protemer NOE distance restraints (both manually and automatically assigned) derived from 15N- and 13C-NOE-HSQC aliphatic experiments at 30 °C, in addition to a set of manually assigned unambiguous inter-protemer restraints. The total number of unambiguous restraints used in the calculations was 1584 (per protomer), of which 1528 intramolecular and 56 intermolecular. Structure-derived H-bonds (25/protomer) were also used as restraints, the latter only being introduced when present in 50% of the preliminary calculations. In the final ARIA run, 200 structures were generated in the last iteration. After refinement of the 60 lowest global energy structures by MD simulation with explicit water, 20 structures ranked on global energy were selected as representative of the structure and used for statistical analysis. Structure quality was evaluated with PROCHECK at the final stage of the calculations by ARIA. The ensemble obtained at the end of the refinement process converged, with an average RMSD of 1.05 Å from the medoid structure as calculated for statistical analysis. Structure quality was evaluated with PROCHECK at the final stage of the calculations by ARIA. The ensemble obtained at the end of the refinement process converged, with an average RMSD of 1.05 Å from the medoid structure as calculated for statistical analysis.
and 50 mM NaCl in D2O, pH 7.3 buffer. The residual water was suppressed using excitation sculpting with 5 ms selective pulse. T1ρ filter of 30 ms was used to eliminate the background protein resonances. The STD spectra were recorded at an ATP-rhNGF ratio of 100:1. To achieve the excess of the cations (50-fold for Mg2+ or 25-fold for Zn2+), 1 mM MgCl2 or 0.25 mM ZnCl2 were added to the samples, with a protein concentration of 20 μM (sample with Mg2+) or 10 μM (sample with Zn2+). Notably, the presence of Zn2+ required strict controlling of the experimental conditions. Zn2+ itself affects the sample, resulting in protein precipitation outside an optimized ATP-Zn2+-rhNGF stoichiometric ratio.

The 1H STD ligand epitope mapping experiments [47] were performed under quantitative conditions considering the nonuniform relaxation properties of ATP. The inversion recovery T1 experiments showed that the 1H T1 relaxation times of ATP range from 3.9 s in the adenine moiety to 0.43 s in the sugar moiety. Therefore, STD amplification factors were determined with a short saturation delay of 0.21 s to avoid the effect of the longitudinal relaxation rate on the signal intensities [48]. Spectra were recorded with a 6578 Hz spectral width, 16,384 data points, a relaxation delay of 5 s and 4500–10600 scans. Selective saturation was achieved by a train of 50 ms long Gauss-shaped pulses separated by 1 ms delay. The on-resonance selective saturation of rhNGF was applied at 0.32 ppm at transmitter offset referenced to 4.64 ppm. The off-resonance irradiation was applied at 30 ppm for the reference spectrum. Subtraction of the on and off-resonance spectra was performed internally via phase cycling. Spectra were zero-filled twice and apodized by an exponential line-broadening function of 3 Hz. Errors in the STD amplification factor were estimated according to the formula: STD amplification factor absolute error = STD amplification factor × (NSTD/ISTD)2 + (NREF/IREF)2)1/2 [49]. NSTD and NREF are noise levels in STD and reference spectra, respectively. ISTD and IREF are signal intensities in STD and reference spectra, respectively.

The 1H-15N NOESY-HSQC spectrum of ATP-rhNGF + Mg2+ complex was collected. All spectra were processed using NMRPipe/NMR-Draw [43] and analyzed using CARA [44]. The combined 1H-15N CSP (Δδ) were calculated from 1H and 15N CSP using the equation: Δδ = ((Δδ 1H)2 + (0.04 × Δδ 15N))1/2 [50].

2.4. FT-IR

The infrared spectra were recorded with a FT-IR Bruker Vertex 80 spectrometers with a nominal resolution of 2 cm−1 and an averaging of 128 scans and apodized with a Blackman-Harris 3-term function. All spectra were recorded at 25 °C with a Golden Gate ATR temperature-controlled cell (Specac). rhNGF at 0.15 mM concentration in 50 mM Hepes buffer, pH 7, was used. The visibility of the amide I and amide II bands was achieved by subtracting the original protein solution spectra with the spectrum of the Hepes buffer alone.

2.5. ITC

ITC experiments were carried out using an ITC 200 microcalorimeter (Malvern Panalytical, UK). Experiments were performed in 50 mM Hepes pH 7.2. rhNGF concentration was determined using a Nanodrop LITE Spectrophotometer (Thermo Fisher Scientific, US). 100 mM ATP stock solution was prepared in the same buffer. Both rhNGF and ATP samples were centrifuged for 5 min at 13000 rpm and degassed. Titrations were performed at 30.0 ± 0.1 °C with a stirring of 750 rpm to ensure rapid mixing. A first set of 19 injections of 2 μL, 2 mM ATP, were performed into a measurement cell filled with 200 μL of 40 μM dimeric rhNGF. The first set of titrations was followed by a set of 19 injections of ATP at the same concentration, to approach the plateau, for a grand total of 38 injections. The two measurements files were merged by using the software concar (Malvern). The same protocol was repeated two more times with 5 mM ATP while keeping rhNGF at 40 μM, to ensure a larger ATP-rhNGF final molar ratio. Negative control titrations were performed by running a set of 19 injections of 2 or 5 mM ATP into the cell filled with buffer only. Negative controls allowed for the estimation of ATP dilution heats.

Integration of the raw thermograms has been done with the program NITPIC [52] that allowed for a precise and unbiased extraction of the heats from the experimental curves.

Global fitting of the three ATP-rhNGF titration experiments by nonlinear least squares was performed to analyze the ATP-rhNGF binding isotherms by means of SEDPHAT software [53]. Due to the complexity of the binding interaction, the fitting procedure has been carried out by assuming one ATP binding site per monomer of rhNGF, as previously reported by mass spectrometry in the absence of divalent cations [22]. The binding sites on the two protomers were assumed strictly symmetric, as suggested from NMR data.

2.6. MD Simulations and Analysis

All atom MD simulations were performed using NAMD [54] with the CHARMM36 [55] force field and TIP3P water model on GPU server. The PME method was used for electrostatics with a real space cutoff of 12 Å. All bonds involving hydrogen atoms were constrained to their equilibrium values with the SHAKE algorithm. An integration time step of 2 fs was used. Simulations were carried out in the NPT ensemble with p = 1 atm and T = 300 K using Nose-Hoover algorithm and Langevin piston pressure control. NMR structure rhNGF was modeled at pKa = 7.0 without applying the two-fold symmetry relationship between the protomers. In order to obtain insight into dynamical fluctuations in apo-form of rmNGF, we evaluated average RMSD per residue from the two separate 300 ns MD simulations performed for both dimers which were solvated by the neutral aqueous solution enclosed in the simulation box of 80 × 72 × 72 Å3. Neutrality was achieved by the addition of the proper number of Cl− counter-ions to the simulation box. Next, four different models of ATP-rhNGF + Mg2+ complex were generated in which different ATP poses, selected according to

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to the information obtained from the NOE analysis, were taken into account. All models were solvated by 50 mM NaCl. Each system was simulated for 600 ns under the same conditions.

An algorithm was developed to detect the most stable poses of ATP in contact with rhNGF. When analyzing the distances between given ATP and rhNGF hydrogens, we first selected ATP poses where the distances between the hydrogen atoms were shorter than 4.5 Å. The search was further substantiated by calculating the duration of each continuous contact during the MD simulation. The final selection was made under the requirement of a contact duration of more than 10 ns.

To quantitatively assess binding preferences of ATP for Site1 or Site2 from MD simulations, the potential energy differences between bound, \( '_B' \), (to site1/2) and unbound, \( '_U' \), states of ATP were calculated. The internal (self-) energy of the ATP molecule, \( E_{AA} \), and its interaction energy with the surroundings, which include protein and solvent molecules, \( E_{AS} \), were considered according to the following formula:

\[
\Delta E_{1/2} = E_{AS,B} + E_{AA,B} > _{1/2} > E_{AS,U} + E_{AA,U} \quad (1)
\]

Each average \(< > \) was taken over 2000 frames of MD trajectory pre-selected according to the position of ATP molecule relative to Site1/2 or its unbound state. Omitting the change in internal energy of protein and solvent in (1), which is estimated to be comparable for both sites, and neglecting the change in molecular volume, the difference \( \Delta E_{1-\Delta E_{2}} \) can be called the relative binding enthalpy of ATP.

2.7. SPR

Experiments were performed with a Biacore T100 equipment (GE Healthcare, UK), equipped with Biacore T100 Control software 2.0.1. The amine coupling reaction was performed according to manufacturer’s instructions (GE Healthcare, UK), on CM5 chips.

2.7.1. ATP-rhNGF interaction

rhNGF was dissolved in 10 mM Sodium Acetate pH 5.5 at 100 µg/mL and immobilized at 8700 RU surface concentration on CM5 chip. Due to the very low binding response of ATP on rhNGF when injected as analyte, several attempts were made to increase the binding, by changing association times, coupled with different flows. No improvement was observed.

2.7.2. Kinetics and affinity experiments

TrkA (R&D, TrkA Fc Chimera, 175-TK-050) and p75<sup>NTR</sup> (R&D, R/TNFRSF16 Fc Chimera, 367-NR-050) receptors extracellular domains were used as ligands, dissolved in 10 mM Sodium Acetate pH 5 at 20 µg/mL and immobilized at a 3600 RU (TrkA) and 2600 RU (p75<sup>NTR</sup>) surface concentration on CM5 chip. The analyte proteins were injected in Hepes Buffer Saline (20 mM Hepes, 150 mM NaCl, 0.005% P20, pH 7.2), at a flow rate of 30 µL/min, 180 s association time and 600 s dissociation time. Samples for the kinetics measurements were prepared by 1:1 serial dilution, starting from a mixture of rhNGF 50 nM and/or Mg<sup>2+</sup> 500 nM, Zn<sup>2+</sup> 500 nM, ATP 500 nM. Concentration of rhNGF for the kinetic analysis was in the range 0.39 nM – 12.5 nM. Six different concentrations obtained by 1:1 dilution have been used. Chip regeneration was performed with 10 µL of 10 mM NaOH. Every dataset was repeated 3 times. The measurement of rhNGF alone was only carried out once, being the resulting kinetic analysis in agreement with the previously reported literature data [56]. Data analysis was carried out using Biacore T100 Evaluation 2.0.1 and BIAevaluation 3.2 Softwares. A 1:1 Langmuir fitting model was assumed for the evaluation of the kinetic and equilibrium constants.

3. Results

3.1. 3D NMR structure of rhNGF in solution

Using NMR we determined the solution structure of unbound rhNGF required for our ATP binding studies. The 2D [1H<sup>3</sup>H<sup>3</sup>] HSQC spectrum of rhNGF, showing a good spectral dispersion (Supp. Fig. S1) was assigned almost to completeness (BMRB: 34515). The NOEs obtained from the 3D [1H-13C] NOESY-HSQC spectra were used for the 3D structural model construction and validation (Fig. 1a, b – PDB: 6YW8; Supp. Table S1). The solution structure closely resembles those determined by X-ray crystallography for rhNGF-ligand complexes (PDB: 1WWW, 1SG1, 2IJ2, 4EDW, 4EDX, 4ZBN, 5JZ7), especially as to the topology of the Cys-knot, whereas it exhibits notable differences within the flexible loop regions (Fig. 1c).

Loop III (residues 60–67), which is missing in all the X-ray crystal structures of bound rhNGF, showed to be flexible also in the apo rhNGF in solution, where residues 60–67 are absent or weak in the 2D [1H<sup>13</sup>C] HSQC spectrum (Supp. Fig. S1). The presence of side-chain NOEs in the [13C-NOESY-HSQC spectra, as well as the inter-residues NOEs in the 13C-NOESY-HSQC spectrum, allowed to partially define this region (Fig. 1b). A wider-open conformation of Loop II (residues 39–48), if compared with the crystallographic structures has also been identified. Furthermore, in the present NMR structure of unbound rhNGF, the N-terminus is flexible and lacks a specific secondary structure propensity, whereas in the X-ray crystal structures of the rhNGF-TrkA complex, it is involved in the interacting surface and adopts a helicoidal conformation (Fig. 1c).

The conformational plasticity of the rhNGF solution structure was also exploited by MD simulations and compared to the only other solution structure of a NT, namely that of rmNGF (Fig. 2, Supp. Figs. S1, S2). The analysis of RMSDs per residue in MD highlights regions with higher conformational plasticity, like Loop II (residues 39–48) in rmNGF as compared to rhNGF (Fig. 2a, b). Indeed, an opening of Loop II has been reported [17,59] in small ligands binding to mNGF (PDB: 4EAX, 4XPJ) or NGF from cobra venom (PDB: 4EC7). Looking at the N-terminus, earlier solution NMR studies on rmNGF prompted to the propensity of the N-terminal residues (1–10) to adopt an helicoidal structure [41]. On the contrary, in rhNGF, the N-terminal residues are poorly defined (Supp. Fig. S2). This was further confirmed by the analysis of evolution, in the MD trajectories, of the intra H-bonds (NH–CO distances) formation engaging the N-terminal residues of either rh- or rmNGF (Fig. 2c–h). These observed differences well agree with previous experiments aimed at the thorough characterization of the biochemical and biophysical properties of m- and hNGFs [60].

3.2. ATP interaction with rhNGF

Hunting for the structural and functional characterization of new endogenous modulators of hNGF biological activity, we deeply investigated its binding properties to ATP. FT-IR measurements (Supp. Fig. S3) ruled out changes in the secondary structure of rhNGF due to the interaction with ATP. SPR, ITC and 1D [1H] STD NMR pinpoint to low affinity (mM range) ATP-rhNGF binding. ITC titration data (Supp. Fig. S4) returned a \( K_D \) of 1.38 mM. As early defined in the 80’s [61] and underlined in the recent literature, transient interactions with weak affinities in the mM range, also named “quinary interactions” [61] were often neglected due to technical challenges in the isolation of molecular complexes to be described [62–64]. Still, these transient and weak quinary interactions are important in specific subcellular compartments and in specific cellular states, for the regulation of dynamic biological processes needed for cell survival [62,65,66]. Therefore, our data prompt to
ATP as a modulator of NTs activity. DSF explored the effects of different divalent cations on the binding interactions between ATP and rhNGF. A major effect of ATP on rhNGF thermal stability occurred in the presence of Zn$^{2+}$ (Supp. Fig. S5). SPR analysis confirmed that ATP binding to rhNGF was markedly affected by Zn$^{2+}$ (Supp. Fig. S6). 1D $^{1}$H STD NMR experiments were used to determine the binding epitope of ATP when bound to rhNGF in the presence of either Mg$^{2+}$ or Zn$^{2+}$ ions. In either case, the strongest interaction was observed for H$_{2}$ on the adenine and H$_{4}'$ on the sugar moieties of ATP, respectively (Supp. Fig. S7). Our 1D $^{1}$H STD NMR experiments complete and extend the previous MS findings by highlighting the ATP functional groups involved in the binding to rhNGF.

3.3. ATP-rhNGF binding: Identification of binding surface

2D $^{1}$H-$^{15}$N HSQC spectra following a titration with increasing amounts of ATP (Supp. Fig. S8) allowed to characterize the rhNGF residues engaged in the interactions with ATP. The analysis of the combined $^{1}$H/$^{15}$N CSP ($\Delta \delta$) showed residues that are likely involved directly in ATP binding with $^{1}$H/$^{15}$N CSP ($\Delta \delta$) larger than 0.04 ppm (Fig. 3).

These residues, localized in Loops I, II and V, showed to be clustered in close spatial proximity on the NMR solution structure of rhNGF (Supp. Fig. S9) and their close neighbors in space show a change in the NOE pattern, consistent with changes in their chemical environment due to the presence of ATP (Table 1, lower row). An earlier study [22] based on MS and mutagenesis studies reported the involvement of the C-terminus of rhNGF in ATP binding. Our data instead clearly show no involvement of this region in rhNGF binding. The 3D $^{15}$N-NOESY-HSQC spectrum, collected at the titration endpoint, allowed the identification of definite newly originated NOEs upon ATP binding, which were unambiguously attributed to ATP protons (Table 1; Supp. Figure S10, Table S2). This analysis points towards the presence of two previously unknown binding sites for ATP on each rhNGF protomer, with a 4:2 (ATP:rhNGF) ratio.
rhNGF) stoichiometry. We named the site encompassing residues V20 and E55 as “Site 1” and that defined around F49 as “Site 2” (Table 1).

The new NOEs attributable to ATP were included to drive MD simulations aimed at mapping the likely binding orientation of ATP on rhNGF. During unbiased and constraint-free simulations of 2.4 μs, the ATP molecule was found migrating over few different protein sites without an expressed tendency to remain trapped in any specific pose.

In a subsequent analysis, we identified all ATP positions explored during MD simulations that correspond to a continuous contact between selected ATP and rhNGF hydrogens lasting longer than 10 ns. These poses are depicted in Fig. 4 (a, b). Interestingly, most of them nicely match the ATP binding sites resulting from
the NOE and CSP analysis. As clearly visible in Fig. 4 (a), ATP binding Site 1 partially overlaps with the binding cleft of rhNGF in the TrkA-d5 domain complex (PDB: 1WWW, 2IFG). Site 2 (Fig. 3(b)) nicely overlaps instead with the rhNGF binding site in the complex with extracellular domain of the p75NTR receptor (PDB: 1SG1). Two representative poses for the two sites are shown in Fig. 4(c, d). In both poses, the distances between HN protons of residues of rhNGF engaged with ATP protons are shorter than 5 Å, in agreement with the identified NOE. These poses also well agree with the STD NMR experiments, showing an involvement of both adenine and sugar moieties of ATP in the rhNGF interaction (Supp. Fig. S7).

Table 1
Observations in the 3D 15N-NOESY-HSQC spectrum collected at ATP-rhNGF titration endpoint indicating ATP binding to two sites.

| New NOEs between HN of rhNGF and ATP protons | Site 1 | Site 2 |
|---------------------------------------------|-------|-------|
| V20 HN – ATP H2                            |       | F49 HN – ATP H3' |
| E55 HN – ATP H2                            |       | Loop 1 (T29-K34) |
| V20, W21, W22, E55                         |       | Loop 2 (S47, V48) |

3.4. ATP-rhNGF binding: Effects on TrkA and p75NTR receptors

SPR data on ATP-rhNGF binding to TrkA and p75NTR receptors are in line with our NMR studies. We investigated ATP binding to the two receptors alone, as well as in the presence of divalent ions, namely Mg2+ (physiologically relevant) and Zn2+ (dysregulated in pathological conditions). The kinetics and affinity data highlight that ATP-Mg2+ weakly affect TrkA or p75NTR affinity if compared to rhNGF alone (Table 2, Supp. Fig. S11). In the case of ATP-Zn2+, there is a clear decrease in the affinity versus the TrkA receptor, mediated by either Zn2+ alone or ATP-Zn2+, as compared to rhNGF alone (Table 2, Supp. Fig. S11). The latter effect is less marked for the p75NTR receptor (Table 2, Supp. Fig. S11).

These effect clearly depend on the relative stoichiometry of the ATP-ion-rhNGF mixture. By careful comparison of the response level for the ATP-ion-rhNGF samples with the reference rhNGF, it emerges that the different ligands induce a change in the maximum response level, on either receptor. This might indicate a change in the fraction of the analyte binding to the receptors. Since
ATP binds with low affinity to rhNGF, this is an indirect evidence of the formation of the ATP-ion-rhNGF complex, which transiently forms during the SPR experiment in small percentage. This effect is stoichiometry-dependent (ATP-ion-rhNGF) and differs when considering Mg\(^{2+}\) or Zn\(^{2+}\) (Supp. Fig. S12). In the case of a large excess of Zn\(^{2+}\) and ATP a dramatic change in the curve shape is observed, reflecting a change in the binding mode. This is not surprising, having been early reported that Zn\(^{2+}\) binds to NGF\(^{[67]}\). To investigate the active role of the ions in the ATP-rhNGF receptors' interaction, a qualitatively control experiment was carried out by pre-incubating the ATP-ion-rhNGF prior to SPR injection. In the case of Mg\(^{2+}\), no effect of the pre-mixing was observed on the binding affinity (TrkA (KD = 0.27 nM); p75\(^{NTR}\) (KD = 1.07 nM) with respect to rhNGF alone (TrkA KD = 0.16 nM; p75\(^{NTR}\) KD = 0.83 nM). The preincubated ATP-Zn\(^{2+}\)-rhNGF mixture resulted in a clear trend towards a decrease of the affinity to either TrkA (KD = 0.89 nM) or p75\(^{NTR}\) (KD = 3.20 nM) with respect to rhNGF alone (TrkA KD = 0.16 nM; p75\(^{NTR}\) KD = 0.83 nM) and confirmed a more pronounced effect on TrkA than on p75\(^{NTR}\) receptor's binding.

### 4. Discussion

The present study by integrative structural biology, combining multiple experimental techniques, unveiled the binding cartography of ATP to rhNGF, aiming at the structural and functional characterization of new endogenous modulators of NGF biological activity.

The ATP-rhNGF binding is characterized by low affinity (mM range), as confirmed by SPR, ITC and STD-NMR. It can thus be counted among transient interactions with weak affinities, named “quinary interactions”, which were recently highlighted\(^{[61]}\) to be important for the regulation of dynamic biological processes needed for cell survival. Therefore, our data prompt to ATP as a modulator of neurotrophins activity.

Many biological data confirmed the presence of low extracellular ATP levels in physiological conditions, while increased extracellular ATP levels up to mM concentration have been reported to cause cell damage or inflammation. The ATP modulating purinergic signaling suggests that any alteration of the fine tuning may affect several human diseases, including neurodegenerative disorders as...
well as inflammatory dysfunction and tumors [33]. High levels of Zn$^{2+}$ were found in cell damage or neurodegenerative conditions [28]. What is more, ATP in the 0.1–10 mM concentration range have been recently reported to kinetically modulate pathogenic tau fibrillation [68]. Furthermore, millimolar concentrations of ATP were found to be linked to a reduction of the fibrillation of neuronal proteins [69] and ATP was found to specifically bind the RNA-binding domain of the nucleocapsid protein of SARS-CoV-2 with a millimolar $K_D$ [70].

We investigated the concomitant presence of ATP and Mg$^{2+}$ or Zn$^{2+}$ and conclude that a tight regulation of the relative stoichiometry of ATP-ion-rhNGF is needed to modulate the endogenous biological response. Indeed, it is well established that NTs and the purinergic receptors systems are in functional interplay, in physiological as well as in pathological conditions [71,72]. The direct ATP and rhNGF interaction was however only recently described [20]. Our data support this finding. Based on the solution NMR titration of ATP to rhNGF we identified the residues in contact with ATP, localized in Loops I, II and V. The solution NMR analysis in the presence of Mg$^{2+}$ allowed identifying new NOEs attributable to ATP, which pointed to the existence of two binding sites of ATP on rhNGF, identified as Site 1 and Site 2. These data were functionally used to guide an MD study to map the likely orientation of ATP in binding the rhNGF. Site 1 cover regions involved into the binding of rhNGF to the TrkA-d5 domain, while residues involved in the Site 2 belong instead to the region of rhNGF known to be involved into the binding with p75NTR receptor extracellular domain.

It is worthy of mention that König et al. [21] suggested one of the two ATP-binding sites to be located close to the cysteine knot in rhNGF. Furthermore, time-dependent proteolytic digestion followed by EI and MALDI-TOF MS analyses prompted to the stability of the ATP-rhNGF complex to be unaffected by the loss of the N-terminal (1–9) peptide SSSHPIFHR and of the Loop II (35–50) peptide EVMVLG EVNINNSVFK. As depicted in Fig. 5, our combined CSP NMR data and MD simulations weakly support these findings. Based on MS experiments, Rose et al. [73] prompted ATP binding site of the FGF2 growth factor to overlap with its HBD. Moreover, it has been previously hypothesized that NGF also possesses a HBD (amino acids 113–120) that binds heparin very weakly [74]. Since evidences have been provided, that residues (113–115) within the HBD play an important role for bioactivity of NGF [75] Hasche et al. [22] explored several rhNGF mutants. They tested whether the mutant proteins could still bind to ATP as well as whether the ATP-rhNGF conjugates could still act as neuropeptidic ligands. Interestingly all mutants (rhNGF[K115S] and rhNGF[S113T/R114S/K115S]) but one (rhNGF[L112A]) lost their ability to bind ATP as well as their ATP mediated neuroprotection. It is tempting to hypothesize that the loss of ATP binding and neuropeptidic activity could likely be ascribed to misfolding of the rhNGF mutants, as being already suggested for rhNGF mutants at these very same positions reported to have lost their bioactivity in several bioassays (PC12 cells neurite outgrowth, TrkA phosphorylation and binding) [75]. It is worthy of note that none of the above-mentioned rhNGF mutants, located at the C-termnus of rhNGF and proposed to be part of the rhNGF HBD, overlap with the residues encompassing either of the two ATP-rhNGF binding sites unveiled in the present study. In addition, the proposed C-terminal residues of rhNGF do not overlap either with those rhNGF residues, elucidated by using carbohydrate microarray and computational approaches, to be involved in interactions with glycosaminoglycans [18]. Intriguingly, one of the two ATP-rhNGF binding sites unveiled in the present study, namely Site 2, instead, fully overlap with that proposed by Rogers et al. [18] for the CS-E, a glycosaminoglycan homolog of heparin, to bind to rhNGF (Fig. 5).

**Fig. 5.** rhNGF ligands binding sites. rhNGF residues in contacts with ATP during MD simulations (green filled diamonds). rhNGF residues affected in NMR CSP upon ATP titration (red filled circle). rhNGF residues predicted by computational docking [18] to be involved (within 5 Å of the oligosaccharide) in CS-E binding (light blue filled “stars”); rhNGF residues within 5 Å from ATP binding Site 1 or Site 2 according to the best MD poses (see Fig. 4) are highlighted by magenta or yellow upper lines, respectively. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
Furthermore, the binding of lipids to NGF have been shown to possibly result in shifting the equilibrium between forming the NGF-p75\textsuperscript{NTR} complex or forming the NGF-TrkA complex thus shaping the signal transduction pattern of NGF. Interestingly crystallographic studies of mouse and cobra NGF-lipids complexes indicated that lysophosphatidyl serine (PDB ID 4EAX), lysophosphatidylinositol (PDB ID 4XPJ) and docosanoic (PDB ID 4EC7) acid bind in a NGF pocket-lining residues mostly overlapping Site 2 [17,59].

Thus, our findings clearly highlight Site 1 as a likely specific ATP-rgNGF binding site, whereas Site 2 represents a likely promiscuous rhNGF binding site for endogenous ligands such as ATP, glycosaminoglycans and lipids. Indeed, this hypothesis is substantiated by the significantly difference in the potential energy of NGF-ATP interactions of Site1 versus Site2. The relative binding enthalpy of ATP for Site 1 is indeed –12.2 kcal/mol lower than for Site 2. This observation opens new potential investigational avenues for the biological implication of these small molecules binding into NTs biology.

The present results further support the suggested role of ATP as a functional endogenous modulator of NGF bioactivity. SPR data on rhNGF binding to the TrkA and p75\textsuperscript{NTR} receptors well agree with our NMR studies. The K\textsubscript{D} affinity values of rhNGF binding to its receptors showed a weak effect of either Mg\textsuperscript{2+} or ATP-Mg\textsuperscript{2+}. In the presence of Zn\textsuperscript{2+} or ATP-Zn\textsuperscript{2+} we detected a higher effect (lower rhNGF binding affinity) on TrkA than on the p75\textsuperscript{NTR} receptor. This clearly depends on the relative stoichiometry of the ATP-ion-rhNGF mixture and is more evident in the presence of a large excess of ATP-Zn\textsuperscript{2+}.

As demonstrated previously by MALDI-TOF MS analysis, the stoichiometry of the ATP-rhNGF complex depends on the type of divalent ions [21,22]. It is tempting to hypothesize that in physiologic conditions, ATP might amplify hNGF signaling through TrkA receptor, as reported [23]. In neurodegeneration or cellular stress, massive release of extracellular ATP is reported [33] while a reduction of TrkA signaling has been observed [76]. In line with this, our data suggest a reduction of hNGF affinity for the TrkA receptor in the presence of Zn\textsuperscript{2+} and ATP. Moreover, the activation of P2X receptors by mM ATP concentration in trauma conditions triggers the prevention of a large inflammatory response [32,71]. The direct binding of ATP to hNGF – albeit of low affinity - might also serve as a molecular switch for the relative activation of NTs versus purinergic receptors systems to meet better the cellular needs.

5. Conclusion

The present 3D binding cartography and the functional description of the ATP-rhNGF interaction represents an important step towards the understanding of the complex puzzle underlying the molecular determinants of NTs binding to its receptors in health and disease. Furthermore, it paves the way towards to the characterization of other multifunctional endogenous ligands (e.g. lipids, lysophospholipids, fatty acids and glycosaminoglycan oligosaccharides [15,18,19]) as novel transient modulators of NGF, as well as of the other members of the NT family [77]. The outcomes of the present structural study shall pave the way for further extending the research on this important biological topic aiming at the precise identification of the molecular mechanism underlying neurotrophins signaling by endogenous small molecules and the cross-talk with the purinergic pathways.

CRediT authorship contribution statement

Francesca Paoletti: Conceptualization, Formal analysis, Investigation, Writing, Visualization. Franci Merzel: Formal analysis, Investigation, Visualization, Funding acquisition. Alberto Cassetta: Formal analysis, Investigation, Visualization. Iza Ogris: Formal analysis, Investigation. Sonia Covacevich: Formal analysis, Investigation, Visualization. Jožo Grždalenik: Formal analysis, Investigation, Visualization, Funding acquisition. Doriano Lamba: Conceptualization, Formal analysis, Investigation, Writing, Visualization. Simona Golič Grždalenik: Conceptualization, Formal analysis, Investigation, Writing, Visualization, Funding acquisition.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.csbj.2021.05.009.

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