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Flexibility of the PDZ-binding motif in the micelle-bound form of Jagged-1 cytoplasmic tail

Matija Popovic a, Ventsislav Zlatev a, Vesna Hodnik b, Gregor Anderluh b,c, Isabella C. Felli d, Sándor Pongor a,⁎, Alessandro Pintar a,c

a Protein Structure and Bioinformatics Group, International Centre for Genetic Engineering and Biotechnology (ICGEB), AREA Science Park, Padriciano 99, I-34149 Trieste, Italy
b Laboratory for Biosynthesis and Biotransformation, National Institute of Chemistry, Hajdrihova 19, 1000 Ljubljana, Slovenia
c Magnetic Resonance Center and Department of Chemistry, University of Florence, Via Luigi Sacconi 6, I-50019 Sesto Fiorentino, Italy

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A B S T R A C T

Human Jagged-1, one of the ligands of Notch receptors, is a transmembrane protein composed of a large extracellular region and a 125-residue cytoplasmic tail which bears a C-terminal PDZ recognition motif. To investigate the interaction between Jagged-1 cytoplasmic tail and the inner leaflet of the plasma membrane we determined, by solution NMR, the secondary structure and dynamics of the recombinant protein corresponding to the intracellular region of Jagged-1, J1_tmic, bound to negatively charged lysophospholid micelles. NMR showed that the PDZ binding motif is preceded by four α-helical segments and that, despite the extensive interaction between J1_tmic and the micelle, the PDZ binding motif remains highly flexible. Binding of J1_tmic to negatively charged, but not to zwitterionic vesicles, was confirmed by surface plasmon resonance. To study the PDZ binding region in more detail, we prepared a peptide corresponding to the last 24 residues of Jagged-1, J1C24, and different phosphorylated variants of it. J1C24 displays a marked helical propensity and undergoes a coil-helix transition in the presence of negatively charged, but not zwitterionic, lysophospholipid micelles. Phosphorylation at different positions drastically decreases the helical propensity of the peptides and abolishes the coil-helix transition triggered by lysophospholipid micelles. We propose that phosphorylation of residues upstream of the PDZ binding motif may shift the equilibrium from an ordered, membrane-bound, interfacial form of Jagged-1 C-terminal region to a more disordered form with an increased accessibility of the PDZ recognition motif, thus playing an indirect role in the interaction between Jagged-1 and the PDZ-containing target protein.

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1. Introduction

Ligands to Notch receptors are type I membrane spanning proteins. They all share a poorly characterized N-terminal region, a DSL (Delta/Serrate/Lag-2) domain, a series of tandem epidermal growth factor-like repeats, a transmembrane segment, and a unique cytoplasmic tail of ~150 amino acids [1]. Five different ligands to Notch receptors have been identified in mammals, three orthologs (Delta-1, -3 and -4) of Drosophila Delta and two orthologs (Jagged-1 and -2) of Drosophila Serrate. Although the molecular mechanisms of ligand specificity are still unclear, evidence from in vivo studies suggest that each ligand exerts non-redundant effects. Gene knock-out of Jagged-1 [2] or Delta-1 [3], heterozygous deletion of Delta-4 [4], or homozygous mutants in Jagged-2 [5] all lead to severe developmental defects and embryonic lethality in mice. There is no significant sequence similarity shared among the intracellular region of the different ligands [6], apart from the identical PDZ binding motif (ATEV) found at the C-terminus of Delta-1 and Delta-4. Neither Delta-3 nor Jagged-2 presents a PDZ recognition motif whereas the cytoplasmic tail of Jagged-1 contains a different C-terminal PDZ interacting motif (EYIV). Jagged-1 was shown indeed to interact in a PDZ-dependent manner with afadin (AF6), a protein located at cell–cell adherens junctions [7,8]. The cytoplasmic tail of Notch ligands is also a target for ubiquitination and ligand endocytosis. Both Mind bomb 1 [9–11] and a myristoylated, membrane-anchored form of Neuralized-like 1 [12] have been proposed to be involved in...
mono-ubiquitylation of Jagged-1 in mice. Finally, there is compelling evidence that Notch ligands, much alike Notch receptors, undergo ectodomain shedding and regulated intramembrane proteolysis. The proteolytic processing of Jagged-1 is mediated by the ADAM17 metalloprotease [13,14] and by the presenilin-γ-secretase complex [13]. Although the fate of the cleaved intracellular region is still unclear, it is possible that it acts as a membrane-tethered transcriptional co-activator. The C-terminal fragment of Jagged-1 comprising part of the transmembrane segment and the intracellular region expressed in COS cells was shown to localize mainly in the nucleus, and to activate gene expression through the transcription factor activator protein 1 (AP1/p38/jun) enhancer element [13].

Depending on the environment, the intracellular region of Jagged-1 is then supposed to exist in at least two distinct forms, as a membrane-tethered protein located at the interface between the membrane and the cytoplasm and as a soluble nucleocytoplasmic protein. In a previous study [15], we showed that a recombinant protein corresponding to the cytoplasmic tail of human Jagged-1 (J1_tmic) is mainly disordered in solution, but has a distinct propensity to adopt a helical conformation in the presence of TFE. J1_tmic also gains secondary structure upon binding to the negatively charged surface of SDS micelles, or to vesicles made of negatively charged phospholipids such as DMPS and DMPG, which are prevalent components of the inner layer of the plasma membrane in eukaryotes. On the contrary, the structure of J1_tmic is not affected by the presence of vesicles formed by a zwitterionic phospholipid like DMPC, suggesting that the negative charge density at the surface of SDS micelles or lipid vesicles is required to promote binding and secondary structure formation. We also showed that helix formation is strongly pH-dependent, with a sharp increase in the helical content from pH 7 to 6. As J1_tmic contains six endogenous histidines, we suggested that protonation of one or more of the histidines is promoting helix formation or extension.

The partial folding of the cytoplasmic domain of Jagged-1 accompanied by its association with the inner side of the cell membrane may have relevant effects on the function of Jagged-1 in Notch signaling. For instance, it would selectively mask certain residues that are potential targets for post-translational modifications such as phosphorylation, ubiquitylation, or O-glycosylation with β-N-acetylglucosamine. In a similar way, it would mask or expose selected binding motifs with respect to binding partners. The partial folding and association of the intracellular region of Jagged-1 with the membrane is also expected to reduce the accessibility of the PDZ binding motif. To get more insight into the interaction between Jagged-1 cytoplasmic tail and the membrane, we studied a model system made of the recombinant protein corresponding to the cytoplasmic tail of Jagged-1 (J1_tmic) and negatively charged lysophospholipid micelles used to mimic the interface between the plasma membrane and the cytosol. We applied solution NMR methods to determine the secondary structure and dynamics of J1_tmic bound to micelles. Furthermore, we hypothesized that phosphorylation of selected residues close to the PDZ binding motif might alter the affinity for the negatively charged inner layer of the cytoplasmic membrane and possibly change the conformational properties of Jagged-1 C-terminal region. To test this hypothesis, a series of peptides corresponding to the C-terminal sequence of Jagged-1 (JAG1_HUMAN, residues 1195–1218) and phosphorylated at different positions was synthesized, and their conformational properties studied by far-UV circular dichroism.

2. Materials and methods

2.1. Protein expression and purification

The recombinant protein corresponding to the cytoplasmic tail of human Jagged-1, starting from the putative presenilin/γ-secretase cleavage site and with the transmembrane cysteine mutated to alanine (residues 1086–1218 of JAG1_HUMAN; C1092A; see Supporting Material, Fig. S1) was expressed in BL21(DE3) Escherichia coli (Novagen) cells from a codon-optimized synthetic gene and purified in two steps by immobilized metal ion affinity chromatography followed by RP-HPLC, as described [15]. For the preparation of the 15N- and of the 13C-labeled proteins, cells were grown in a minimal medium containing 6 g/L Na2HPO4, 3 g/L KH2PO4, 0.5 g/L NaCl, 0.12 g/L MgSO4, 0.01 g/L CaCl2, 1.7 g/L yeast nitrogen base without amino acids and ammonium sulfate (Difco), and 100 μg/mL ampicillin, supplemented with 0.5 g/L 15NH4Cl, 5 g/L d-glucose or 0.5 g/L 15NH4Cl, 5 g/L U-13C6-D-glucose for the single and doubly labelled proteins, respectively. Expression and purification of the labelled proteins were carried out as described above. The purified proteins were analyzed by LC–MS to confirm their identity.

2.2. NMR sample preparation

The protein stock solutions were prepared dissolving the freeze-dried labeled proteins in sodium phosphate buffer (20 mM), pH 6.1. Lysophospholipid stock solutions were prepared dissolving the 1-Myristoyl-2-Hydroxy-sn-Glycero-3-[phospho-rac-[1-glycerol]] sodium salt (LMPC) or 1-Myristoyl-2-Hydroxy-sn-Glycero-3-Phosphocholine (LMPC) (Avanti Polar Lipids Inc., Alabaster, AL, USA) in sodium phosphate buffer (20 mM, pH 6.1). The mixed micelle (LMPC/LMPC, 80/20 mol/mol) lysophospholipid stock solution was prepared dissolving the lysophospholipids in CH3OH/CHCl3, mixing the appropriate volumes of the two, gently drying the solution, and redissolving the mixture in sodium phosphate buffer, pH 6.1. Samples for NMR spectroscopy were prepared mixing the appropriate volumes of stock solutions, and adding D2O (10 vol.%).

The final protein concentration was ~0.1 mM for the reference sample dissolved in buffer only, ~0.5 mM for the 15N-labeled samples in the presence of LMPC (80 mM), LMPC (80 mM) or LMPC/LMPC (80 mM), and 0.9 mM for the 13C-labeled protein in the presence of LMPG (150 mM). Assuming a micelle aggregation number of ~120, the protein/micelle molar ratio was ~0.75. An additional sample with a protein concentration of ~0.33 mM and a LMPC/LMPC (80/20 mol/mol) concentration of 113 mM, which corresponds to a protein/micelle ratio of 0.35 was also prepared. From an average micelle aggregation number of ~120, a MW of 478.5 for each LMPG molecule, and a 1:1 stoichiometry, the calculated MW of the complex would be ~73 kDa.

2.3. NMR spectroscopy

NMR data were acquired at 298 K. Preliminary and 15N relaxation experiments were acquired on a Bruker Avance 500 MHz spectrometer equipped with a TXI triple resonance cryoprobe using the 15N-labeled protein. For assignment purposes, standard HNCA, CBCA(CO)NH, HNCCO, HNCO and HN(CA)NH experiments were acquired on Bruker Avance 700 and 800 MHz spectrometers equipped with cryogenic probeheads optimized for 1H detection. The protonless experiments CON (H/C)BCACON and (H/C)BCANCO [16–18] were acquired on a Bruker Avance 700 MHz equipped with a cryogenic probehead optimized for 13C direct detection. 1H−13C couplings, constant and exchange rates of backbone amides with solvent were estimated from HNHA [19] and CLEANEX-PM [20] experiments, respectively. Data were processed using XwinNMR and analyzed with the CCPN program ANALYSIS (http://www.ccpn.ac.uk). Chemical shifts were referenced to DSS. Deviations of Co, Cα, Hα, and Cβ chemical shifts from random coil values [21] were used to determine secondary structure regions. R1 and R2 relaxation rates were estimated from fitting a three parameter exponential decay to experimental intensities of HN cross peaks acquired with delays of 0.010, 0.150, 0.540, 1.00, 2.50 s for R1 and 17, 68, 153, 204, 271 ms for R2 experiments. 1H−15N steady-state
heteronuclear NOEs were obtained from the ratio of peak heights in paired spectra collected with and without proton saturation during the relaxation delay. 

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\text{J}_{\text{HN}} \text{coupling constants were calculated from the intensity ratio between the Hx cross peak and the diagonal HN peak using the formula } I_{\text{HN}}/I_{\text{HNx}} = -\tan^{-1}(2 \cdot \tau_1 \cdot f_\xi), \text{ where } J \text{ is the } J_{\text{HN}} \text{ coupling constant and } \xi = 13.05 \text{ ms. No correction for relaxation effects was applied. Solvent-HN exchange rates were estimated by semi-preparative RP-HPLC on a Zorbax 300SB-C18 column (9.4×250 mm, 5 \mu m, Agilent) and freeze-dried. The identity of the peptides was checked by LC-MS and the purity (>95%) estimated from RP-HPLC.}
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2.5. Circular dichroism

Samples for CD spectroscopy were prepared dissolving the freeze-dried J1_tmic protein or the J1C24 peptides in 5 mM Tris/HCl buffer, pH 7.4, to obtain stock solutions that were further diluted appropriately. Concentrations were estimated from the UV absorbance at 280 nm using the calculated extinction coefficients (16500 M⁻¹ cm⁻¹ and 6690 M⁻¹ cm⁻¹ for J1_tmic and the J1C24 peptides, respectively). Far-UV CD spectra were recorded for J1_tmic (7.5 \mu M) and for the J1C24 peptides (~40 \mu M) in buffer alone, in the presence of different concentrations of TFE, and in the presence of 1 mM for J1_tmic or 3 mM for the J1C24 peptides) LMPG or LMPC in 5 mM Tris·HCl buffer, and in the presence of 1 mM lysophospholipids (LMPG or LMPC) at pH 7.4 or pH 6.0.

Far-UV CD spectra of J1_tmic in buffer at neutral pH is typical of disordered proteins, with a minimum at 200 nm and ellipticity close to zero at 190 nm (Fig. 1). While the far-UV CD spectrum remains unchanged in the presence of zwitterionic LMPC micelles, the minimum shifts at 204 nm and the ellipticity at 190 nm becomes positive when J1_tmic is dissolved in the presence of negatively charged LMPG micelles (Fig. 1). At pH 6.0, the negative band is further shifted

2.6. Surface plasmon resonance

DMPG or DMPC in chloroform was dried under reduced pressure in round-bottom flasks to obtain a lipid film. The lipids were hydrated in 10 mM Hepes, 150 mM NaCl, pH 7.4 to form multilamellar vesicles.

The large unilamellar vesicles were then prepared by extrusion of multilamellar liposome suspension through polycarbonate filters with 100 nm pore size. The vesicles were diluted 10 times for DMPG and 40 times for DMPC prior the capturing onto the chip in the running buffer (10 mM Heps, 150 mM NaCl; the experiments were performed at pH 6.0 and 7.4).

The surface plasmon resonance (SPR) measurements were performed using Series S sensor chips L1 (Biacore, GE Healthcare) and the optical sensor Biacore T100 (Biacore, GE Healthcare) [23,24]. The system was primed twice with the running buffer and the chip was conditioned using 4 startup cycles. The vesicles were deposited at a low flow rate (2 \mu L/min) for 60 s for DMPG and 90 s for DMPC to reach the similar immobilization level (3500–4000 response units). After the capturing, the buffer was allowed to run 300 s over the surface to stabilize the vesicles on the chip. Then the samples were injected for 180 s and the dissociation was monitored for additional 180 s. The dilutions were prepared dissolving the freeze-dried J1_tmic in the running buffer and the concentrations used to monitor the binding of J1_tmic to DMPG were 0.195, 0.39, 0.78, 1.56, 3.13, 6.25, 12.5 and 25 \mu M. The surface was regenerated using two 30 s injections of 40 mM octyl-\beta-D-glucopyranoside and one 12 s injection of 0.05% sodium dodecyl sulfate (SDS). The sample and regeneration solutions were injected at flow rate 10 \mu L/min. All experiments were performed at 25 °C. The binding responses were evaluated with the T100 Evaluation Software. Dissociation constants (K_dis) were estimated from the equilibrium binding levels 4 s before the stop of the injection. The responses were fitted to the Steady State Affinity model. K_dis were obtained from two independent titrations.

3. Results

3.1. Secondary structure and dynamics of J1_tmic bound to lysophospholipid micelles

Far-UV CD spectra of J1_tmic in buffer at neutral pH is typical of disordered proteins, with a minimum at 200 nm and ellipticity close to zero at 190 nm (Fig. 1). While the far-UV CD spectrum remains unchanged in the presence of zwitterionic LMPC micelles, the minimum shifts at 204 nm and the ellipticity at 190 nm becomes positive when J1_tmic is dissolved in the presence of negatively charged LMPG micelles (Fig. 1). At pH 6.0, the negative band is further shifted.
at 208 nm, and the shoulder at 222 nm as well as the positive band at 190 nm become more evident. These changes suggest a transition from a mainly disordered state to a partially folded, helical conformation of J1_tmic in the presence of LMPG but not LMPC micelles and confirm secondary structure predictions (Supporting Material, Fig. S1) and the helical propensity of J1_tmic previously observed in the presence of TFE, SDS micelles, and DMPG liposomes [15]. The helical content estimated from deconvolution of CD spectra is ~20% for J1_tmic in the presence of LMPG micelles at neutral pH, and increases to ~30% at pH 6 (Supporting Material, Table S2).

In agreement with CD results, SPR measurements showed that the binding of J1_tmic to chip-immobilized DMPG liposomes was concentration dependant and stable at pH 6.0 or 7.4 (Fig. 2). The estimated $K_d$ for J1_tmic binding to lipid vesicles was evaluated to be $0.9 \pm 0.14$ and $3.5 \pm 0.7 \mu M$ for pH 6.0 and 7.4, respectively. J1_tmic did not bind to DMPC vesicles at 25 $\mu M$, the highest used concentration, at any pH (Fig. 2).

To map the helical segments onto the sequence, we used heteronuclear solution NMR techniques. From preliminary NMR spectra of J1_tmic dissolved in buffer, 115 backbone NH cross-peaks (~92% of the total) of the 125 expected could be detected in the HSQC spectrum, including those arising from the three glycines (Supporting Material, Fig. S2). The amide resonances display very limited chemical shift dispersion in the $^1H$ dimension (7.6–8.7 ppm), and several residues seem to have a major and a minor form. In the 7.2–7.3 ppm range in the $^1H$ dimension (~125 ppm in the folded $^{15}N$ dimension), three peaks tentatively assigned to the $\varepsilon NH$ of the R side chains are detectable, two of which are broad and overlapped. The $^{15}NH$ peaks of the two W side chains appear as distinct resonances, although very close in chemical shift (10.09 and 10.11 ppm). The amide resonances of the 21 N/Q side chains are all clustered in a narrow region.

The HSQC spectrum of $^{15}N$-J1_tmic in the presence of LMPC micelles (Supporting Material, Fig. S2) is very similar to that in buffer, with no or very little chemical shift variation for most of the resonances. The only significant differences are the doubling of W side chain $^{15}NH$s with two new peaks at 10.30 and 10.35 ppm, suggesting the presence of a major and a minor form (Supporting Material, Fig. S3) and the appearance of three peaks in the 7.5–7.7 ppm range (7.51/118.76; 7.58/117.28; 7.71/118.27).

On the contrary, the HSQC spectrum of $^{15}N$-J1_tmic in the presence of LMPG micelles (Fig. 3) is significantly different, with 119 detectable peaks, including those arising from the three glycines (~95% of the total), a new set of resonances appearing in the 7.5–7.8/116.3–118.5 ($^1H/^{15}N$) ppm region and a set of at least six distinct peaks possibly arising from the $\varepsilon NH$ of the arginine side chains. The amide resonances of the N/Q side chains show a larger chemical shift dispersion and many of them appear as distinct pairs of peaks. Increasing the temperature from 298 K to 308 K, or decreasing it to 288 K did not improve the appearance of the HSQC spectrum in a significant way.

The spectrum of $^{15}N$-J1_tmic in the presence of mixed LMPC/LMPG micelles seems to be intermediate between the two spectra recorded in the presence of either LMPC or LMPG micelles, and decreasing the protein/lipid ratio did not yield significant changes.

Sequential assignments of the backbone were achieved using standard triple resonance spectra, supported by NOESY and "proton-less" experiments. Assignments in the N-terminal half were particularly difficult because of the line broadening and low intensity – sometimes under detection limits – of HN resonances in the $^1H–^{15}N$ HSQC spectrum. Assignments of the C-terminal half residues, including the presence of five prolines, were more straightforward due to sharper HN cross-peaks, especially in the last 10–20 residues that

![Fig. 2](image-url) Surface plasmon resonance. Binding of J1_tmic to large unilamellar vesicles as measured by SPR. Black thin lines, binding to DMPG liposomes. Concentrations used were 0.195, 0.39, 0.78, 1.56, 3.13, 6.25, 12.5 and 25 $\mu M$ from bottom to top. Dashed line, binding of 25 $\mu M$ J1_tmic to DMPC liposomes.
includes the PDZ binding motif, which display very sharp resonances. Eventually, 113 of the 125 backbone HN amides could be assigned, and 6 of the 8 prolines.

Deviations from random coil values in the C\(\alpha\), C', Hx, and C\(\beta\) chemical shifts are plotted in Fig. 4 and were used to identify secondary structure regions. Four helical segments could be mapped, h1 (T1087-R1094), h2 (T1111-N1120) and h3 (N1136-N1149) in the N-terminal half of the sequence, and h4 (R1203-R1213) in the C-terminal region preceding the PDZ binding motif. The C-terminal boundary of h2 could not be established precisely as many of the resonances in this region could not be assigned. It is thus possible that h2 extends to K1123 or even to H1129. The location of h1, h2 and h4 is in good agreement with secondary structure predictions. Helix 3 is predicted by PsiPred but not by other methods, and an additional helix is predicted between residues M1161 and R1169. Secondary shifts in this region are however very small, and not as consistent as in the other helical regions. The strong negative secondary C\(\alpha\) shifts observed for some residues are associated with the presence of an adjacent proline, and are not likely to indicate stretches of extended structure. The total helical content estimated from chemical shifts is \(-40\%\), which is in good agreement with CD results taking into account that CD is very sensitive to even slight deviations from regular secondary structure, such as helix bending.

To confirm secondary structure assignments, we also acquired a 3D HNHA spectrum, from which \(3J_{HNH}\) were calculated (Supporting Material, Fig. S5). Because of the variations in the relaxation properties along the sequence, the accuracy of these values should be taken with care. In fact, due to the vanishing intensity of many of the HN and Hx cross peaks no quantitative information could be derived for h1, h2, and h3. For h4, an average \(3J_{HNH}\) of \(5.6 \pm 0.7\) Hz was calculated over residues 120–129, which is close to the mean value of \(5.7 \pm 0.9\) Hz obtained for all assigned residues with cross peaks of measurable intensities. Interestingly, \(3J_{HNH}\) values close to 8 Hz were measured for the last three residues of the PDZ binding motif, suggesting that these residues may be pre-organized in the extended conformation typical of many peptide/PDZ domain complexes.

\(15N\) longitudinal (\(R_1\)) and transverse (\(R_2\)) relaxation rates were measured on the \(15N\)-J1\_tmic sample in the presence of LMPG micelles (Fig. 5). \(R_1\) relaxation rates are in the range 0.62–2.12 s\(^{-1}\), with a mean value of 1.52 s\(^{-1}\) and a standard deviation (\(\sigma\)) of

Fig. 3. NMR. \(^1H\)–\(^15N\) HSQC spectrum of J1\_tmic (0.9 mM) in the presence of 150 mM lysophospholipid (LMPG) in 20 mM phosphate buffer, pH 6.1. For clarity, residue numbering starts with the initial methionine (M1); to obtain the actual numbering, add 1084.
0.27 s$^{-1}$. $R_2$ relaxation rates display a much wider range, varying between 2.53 and 30.6 s$^{-1}$, with a median of 10.2 s$^{-1}$ and $\sigma=5.7$ s$^{-1}$. For several peaks, $R_2$ values could not be determined due to vanishingly small values of the peak intensity at longer delays. The ratio $R_2/R_1$ was also calculated as a rough estimation of the local correlation time ($\tau_c$) assuming isotropic motion. Values are spread between 1.4 and ~30, with a median of 6.0 and a large dispersion ($\sigma=5.5$). There is a clear correlation between the line shape of the HN cross peaks in the $^1H$–$^{15}N$ HSQC spectrum and the $R_2$ values, with large $R_2$ values in N-terminal half of the sequence, particularly in h1, h2, and h3, and smaller $R_2$ values in the C-terminal half, in particular in h4 and in the PDZ binding motif, which is also displaying smaller $R_1$ values.

Heteronuclear $^1H$–$^{15}N$ NOEs can also provide an estimation of local backbone dynamics and are very useful in distinguishing flexible disordered regions from structured globular ones. Because of the limited dispersion in the HN chemical shifts, many regions of the HSQC spectrum suffer from severe peak overlap which, together with the very different line shapes, hamper the accurate determination of peak intensities, hence of NOE values. With these limitations in mind, a plot of $^1H$–$^{15}N$ NOEs (Fig. 5) shows a significant variability along the sequence, with a mean value of $0.37\pm0.22$. Values of $^1H$–$^{15}N$ NOEs in the N-terminal half of the sequence (mean: $0.51\pm0.19$), in particular in the h1 ($0.67\pm0.13$), h2 ($0.54\pm0.20$) and h3 ($0.54\pm0.10$) regions are higher than in the C-terminal half (mean: $0.27\pm0.19$).Remarkably, $^1H$–$^{15}N$ NOEs are below 0.2 in h4, with the last six residues (mean: $-0.11\pm0.28$) corresponding to the PDZ binding motif displaying values that become progressively more negative.

Exchange rates of backbone HN amides with solvent were evaluated using CLEANEX experiments. Although for most of the HNs the exchange rates could not be calculated in a reliable way, a plot of the residual HN intensity in the 60 ms mixing time was helpful in identifying three regions, the loop between h1 and h2, the central portion of J1_tmic, and the C-terminal part including h4, which display relatively higher solvent exchange rates while for all isolated peaks belonging to residues h1, h2, h3 no cross peaks arising from exchange processes with the solvent were identified (Supporting Material, Fig. S6).

### 3.2. Effects of phosphorylation on the C-terminal region

To study the C-terminal region of Jagged-1 cytoplasmic tail in more detail, we prepared, by solid phase peptide synthesis, a peptide corresponding to the last 24 residues of Jagged-1, J1C24 (JAG1_HUMAN, residues 1195–1218), and several variants of it. To test the potential effects of phosphorylation on the intrinsic structural properties of J1C24 and on the interaction with lysophospholipid micelles, three variants were designed to include the potential phosphorylation sites adjacent to (T1197) or included in (S1207, S1210) the helical region and the potential phosphorylation site within the PDZ binding motif (Y1216). Four phosphorylated peptides were synthesized, J1C24pS, J1C24pY, J1C24pSpY, and J1C24pTpS, to cover four possible phosphorylation patterns, pS1210, pY1216,
pS1210/pY1216, and pT1197/pS1207, respectively. An additional variant peptide, J1C24RQ, contains the R1213Q sporadic mutation found to be associated with extrahepatic biliary atresia, a congenital obstruction of the bile ducts. All peptides were designed to include (i) the C-terminal PDZ-recognition motif, (ii) the h4 helical region, (iii) W1196 for a more precise measurement of peptide concentration, and (iv) an acetylated N-terminus to avoid charge effects. Far-UV CD spectra were recorded on the peptides alone, in the presence of TFE, and in the presence of LMPC or LMPG micelles.

Far-UV CD spectra show that J1C24 is mainly disordered in buffer solution but display a significant intrinsic helical propensity in the presence of TFE and undergoes a coil–helix transition in the presence of LMPG, but not LMPC micelles (Fig. 6). The secondary structure formation is strongly pH-dependent, with an increase from 16 to 41% in the helical content in the presence of LMPG micelles when the pH is lowered from pH 7 to pH 6. Overall, CD spectra of J1C24 are consistent with CD and NMR spectra obtained for the full length J1_tmic.

The behavior of phosphorylated peptides is completely different. Phosphorylation upstream of the PDZ binding motif (pT1197, pS1207, pS1210) is sufficient to drastically reduce the intrinsic helical propensity of the peptides, as measured by far-UV CD in the presence of TFE, and to completely abolish the coil–helix transition observed for J1C24 in the presence of LMPG micelles. Phosphorylation (pY1216) or mutation (R1213Q) within the PDZ binding motif reduces the helical propensity of the peptides only partially but abolishes the conformational change observed in the presence of LMPG micelles. It can be remarked that, while J1_tmic is enriched in basic amino acids and has a quite high pI (calculated pI = 9.3), J1C24 and its variants have a negative total net charge which, taking into consideration the C-terminal carboxylate, varies from −2 for J1C24 to −6, depending on the peptide.

4. Discussion

4.1. Mimic of the membrane/cytoplasm interface

Proximal proteins and the cytoplasmic region of transmembrane proteins are peculiar, in that they “live” neither in the aqueous environment of the cytosol nor in the hydrophobic hydrocarbon core of the lipid bilayer, but rather at the interface between the two [25]. This interfacial layer is ~15 Å thick, is made by the polar groups of the fatty acid esters, the negatively charged group of the phosphate moiety, and by the phospholipid head group, which in turn determines the overall net charge of the phospholipid. The interaction of a proximal protein with the interfacial layer is mediated by charge–charge interactions, insertion of hydrophobic side chains into the hydrocarbon core, desolvation, and possibly reorganization of the lipid matrix itself. Several artificial systems have been employed to mimic the solvent/membrane interface, including SDS and DPC micelles, lysophospholipid micelles [26], bicontinuous [27], nanodisks [28], and large (LUV) or small (SLUV or liposomes) unilamellar vesicles. All NMR experiments presented in this study were carried out with LMPG and LMPC micelles. Lysophospholipid micelles are especially attractive because they are prepared in a straightforward manner, they are relatively stable in time in the temperature and pH conditions used for acquisition of NMR data, they have a well
defined composition, and they are biologically relevant, in that their head groups are identical to those found in natural phospholipids of eukaryotic plasma membranes. One of the potential drawbacks in the use of lysophospholipid micelles as models of the membrane/cytosol interface is the high curvature, which is dictated mainly by molecular size and shape. To our knowledge, structural parameters for LMPG micelles have not been yet determined precisely. A molecular mass of 35 kDa was derived by static light scattering measurements for LMPG micelles [29], which corresponds to an aggregation number of 73. This value is however significantly lower than that estimated from translational diffusion measured by NMR for LMPG-protein complexes [26] and for the related LPPG [30], LMPC [29,31], and LPPC [32] micelles, which all show aggregation numbers in the range 120–140. Small-angle X-ray scattering (SAXS) on LPPG micelles suggest an oblate, significantly flat disk shape rather than a spherical one, with a ratio between the two axes, a/b, of 0.65 and a gyration radius of ~37 Å [33]. Induction of secondary structure by lipid model systems in otherwise intrinsically disordered proteins has been observed in several cases, including the cytoplasmic tail of single-pass type I membrane proteins such as the ε [34,35] and ζ subunits [25] of the T cell receptor/CD3 complex. Contradictory results were however reported using different model lipid systems [36,37]. Partial folding of the T cell receptor ε chain was induced by LMPG micelles and DMPG SUVs, but not by POPG LUVs [36,37]. It was proposed that the protein can disrupt the lipid organization of LMPG micelles and DMPG SUVs, with helix formation triggered by the interaction between the hydrocarbon tail of the lipid and the tyrosine side chains of the ITAMs (immunoreceptor tyrosine-based activation motif) [37]. In more stable systems such as POPG LUVs, the organization of the lipid bilayer would be maintained and the interaction restricted to the interface, with no folding observed upon binding. Given that curvature effects are hardly predictable, and that we observed a consistent behavior of $J_{\text{1_tmic}}$ going from SDS micelles to LMPG or LMPC micelles, and to DMPG or DMPC liposomes, we reasoned that lysophospholipid micelles may represent a good model system at this stage. We have no evidence, so far, of hydrophobic interactions occurring between J1_tmic and the hydrocarbon portion of the micelles as the driving force for the observed conformational changes. As already discussed [15], the consistent behavior of J1_tmic in relation to the charge of the micelle surface, rather than its lipid composition, the pH dependence of the conformational change, and tryptophan

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**Fig. 6.** Effects of phosphorylation. Far-UV CD spectra of J1C24 (A, B) and J1C24pS (C, D) in 5 mM Tris·HCl buffer and in the presence of TFE (%, v/v) or in the presence of 3 mM lysophospholipids (LMPC or LMPG) at pH 7.4 and pH 6.0. Peptide concentration was 42μM for both J1C24 and J1C24pS.
fluorescence spectroscopy (data not shown) all point toward the interaction between J1_tmic and the interfacial layer of the micelle.

4.2. A model of Jagged-1 cytoplasmic tail at the membrane/cytoplasm interface

The lack of chemical shift dispersion in the backbone amide $^{1}H$ chemical shifts of J1_tmic in buffer is consistent with a mainly disordered state of the protein in solution and in agreement with previous studies [15]. In the presence of LMPG micelles, the appearance of a few new peaks and the doubling of W side chain $^{13}NH$ resonances suggest that, although remaining mainly unstructured, J1_tmic could interact, through the hydrophobic residues in its N-terminal region, with the hydrophobic core of the lysophospholipid micelle. This interaction is likely to be local, as most of the other resonances are not affected. This is consistent with the results obtained by circular dichroism (CD) where little or no difference was detected between the far-UV CD spectrum of J1_tmic in buffer and that of the protein in the presence of LMPG micelles [15]. On the contrary, the differences observed in the HSQC spectrum of J1_tmic in the presence of LMPG micelles show that a significant interaction is occurring between the protein and the micelle. These differences are in agreement with far-UV CD spectra, which show a coil–helix transition of J1_tmic in the presence of LMPG micelles [15]. For technical reasons, we were not able to accurately measure the affinity constants of J1_tmic for LMPG and LMPC micelles by SPR. In support of CD and NMR results, $K_d$ values measured by SPR for J1_tmic binding to DMPG liposomes, which bear the same head groups as LMPG, were in the micromolar range, while there was no binding to DMPC liposomes. Whereas LMPC is zwitterionic, with the external layer of the micelle dominated by the positive charge of the choline moiety, LMPG is negatively charged because of the phosphate moiety. It is thus possible that the interaction between J1_tmic, which has a calculated pl of 9.3, a total of 29 positively charged residues ($R + K + H$) and a net charge of +10, and the negatively charged LMPG micelle is favored by electrostatic interactions. Indeed, we initially proposed [15] that the increase in the helical content of J1_tmic observed when the pH is decreased from 7 to 6 in the presence of negatively charged SDS micelles or DMPG liposomes and confirmed here may be due to the protonation of the six endogenous histidines. Results obtained for J1C24 do not support however this hypothesis. Although both J1_tmic and J1C24 show a very similar pH-dependence in their conformational change, the J1C24 sequence does not contain any histidine. This suggests that protonation of other residues, most likely aspartate and glutamate side chains, as well as the carboxylate group at the C-terminus might be involved. Desolvation at the interface between the protein and the phospholipids could destabilize the carboxylate group in favor of its protonated form. Coordination of calcium ions could reduce or neutralize the negative charge of E/D residues. Charge–charge interactions, although important, are not likely to be the only driving force for binding. The small but measurable shifts in the positions of the W side chain $^{13}NH$s might be indicative of the insertion of J1_tmic hydrophobic N-terminal residues in the lipid core of the micelle. The fairly good chemical shift dispersion in the N/Q side chain NHs suggests a difference in the environment experienced and possibly specific hydrogen bonding. Even more interestingly, the appearance of relatively sharp, distinct resonances that can be tentatively assigned to $^{13}NH$s of the R side chain suggests the presence of hydrogen bonds between these atoms and possibly the negatively charged phosphate groups of the LMPG micelle.

A short segment of basic amino acids is present in the intracellular region of most type I membrane proteins, close to the transmembrane region. Lipid binding of the intracellular region of several multi-chain immune receptors was shown to be related to the presence of clusters of basic residues [36]. Targeting of proteins to the membrane through basic segments occurring in mainly disordered regions has been shown for src [38], K-ras4B [39], rac1 [40] and PLC-ζ [41]. Re-combinant fluorescent probes exploiting electrostatic interactions with the plasma membrane have been used to study the changes in charge distribution occurring during phagocytosis [42]. On the other hand, basic amino acids are also found in nuclear localization signals [43]. For Jagged-1, which undergoes regulated intramembrane proteolysis, basic segments may thus play a dual role, stabilizing the interaction with the inner leaflet of the plasma membrane and targeting the intracellular region to the nucleus, once the cytoplasmic tail has been proteolytically cleaved.

Backbone resonance assignments, mapping of the secondary structure, together with relaxation and solvent exchange studies, lead to a schematic model of J1_tmic bound to LMPG micelles (Fig. 7). The wide variability of $K_d$ values can be due very different local correlation times, to strong anisotropy, and to exchange processes and, given the complexity of this molecular system, all these mechanisms can contribute at the same time. A possible scenario consistent with the observed line widths, relaxation and solvent exchange data is given by the presence of regions that are partially or totally embedded in the micelle (h1, h2, and h3), regions that bind the surface of the micelle, probably in a dynamical way (h4), and other regions that do not significantly interact with the micelle and freely tumble in solution (the C-terminal PDZ binding motif).

4.3. Phosphorylation as a modulator of protein–protein interactions at the membrane/cytoplasm interface

If electrostatic interactions between the basic intracellular region of Jagged-1 and the negatively charged phospholipids of the membrane are important, it can be expected that phosphorylation, which is reducing the positive net charge, might significantly reduce the

![Fig. 7. Model of J1_tmic bound to micelles.](image-url)
affinity of J1_tmic for the membrane. In fact, all the C-terminal J1C24 peptides used have a negative total charge, ranging from $-2$ to $-6$ (including the C-terminus) depending on the phosphorylation state of peptide. The interaction of J1C24 with negatively charged LMPG micelles is then more likely to be due to conformation-dependent insertion of hydrophobic side chains into the lipid core, specific charge interactions and possibly to hydrogen bonding. Of the several phosphorylation sites predicted in the intracellular of residues within the PDZ recognition sequences has been shown to including $S_{1210}$) does have an effect on the conformation and binding phosphorylation of residues upstream to the PDZ binding motif (in-

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the PDZ binding motif, abolishing the intrinsic helical propensity of that region, also changes its affinity for the membrane thus contributing, in an indirect way, to the interaction with the target protein.

5. Conclusions

The cytoplasmic tail of transmembrane ligands and receptors play a functionally important role as final effectors in signal transduction from the extra- to the intracellular environment. Nevertheless, very few of these systems have been studied, partly because of the complexity of the physical milieu of residence, the interface between the plasma membrane and the cytosol. Here, we propose a model of the cytoplasmic tail of the Notch ligand Jagged-1 based on NMR and CD studies carried out on an artificial system comprising a recombinant protein, $J1_{tmc}$, and negatively charged lysophospholipid micelles used to mimic the interfacial region of the membrane. In our model, four different helical segments mediate the interaction of Jagged-1 with the membrane, while the C-terminal region that bears the PDZ interaction motif remains highly

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