Structural and biochemical differences between the Notch and the amyloid precursor protein transmembrane domains

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γ-secretase cleavage of the Notch receptor transmembrane domain is a critical signaling event for various cellular processes. Efforts to develop inhibitors of γ-secretase cleavage of the amyloid-β precursor C99 protein as potential Alzheimer’s disease therapeutics have been confounded by toxicity resulting from the inhibition of normal cleavage of Notch. We present biochemical and structural data for the combined transmembrane and juxta membrane Notch domains (Notch-TMD) that illuminate Notch signaling and that can be compared and contrasted with the corresponding traits of C99. The Notch-TMD and C99 have very different conformations, adapt differently to changes in model membrane hydrophobic span, and exhibit different cholesterol-binding properties. These differences may be exploited in the design of agents that inhibit cleavage of C99 while allowing Notch cleavage.

INTRODUCTION
The Notch receptor initiates one of the most important signaling pathways of mammalian cells. Notch signaling occurs in all vertebrate organisms and has been implicated in an array of developmental patterning choices (1–4). In the nervous system alone, Notch signaling plays key roles in neurogenesis, axon and dendrite growth, synapse plasticity, and neuronal death (5–10). The human Notch receptor is a type I membrane protein with more than 2500 amino acids and a modular domain composition (Fig. 1). Like the amyloid precursor protein (APP), Notch is subjected to a series of proteolytic steps that mature the protein, prime it for signaling, and trigger signal activation. The penultimate step of this process involves cleavage of Notch within its transmembrane domain (TMD) by γ-secretase to release a short peptide (NB) into the extracellular milieu and the large Notch intracellular domain (NICD) into the cytosol (11–14). The NICD translocates to the nucleus, where it forms transcriptional activator complexes (15–17) that control expression of many different genes (18, 19). The mechanism of Notch cleavage by γ-secretase cleavage is not well established.

There are experimentally determined structures for various soluble domains from Notch receptors. In previous studies, structures of the ligand-binding epidermal growth factor (EGF) repeats, the negative regulatory region (NRR), the ankyrin repeat domain, and the complete NICD–CSL (CBF-1/suppressor of hairless/Lag-1)–MAML (Mastermind-like) transcription factor complex were determined (20, 21). More recent structural studies have focused on the NRR, the complex of the EGF repeats with a Notch ligand, and a variety of other studies focusing on posttranslationally modified EGF repeats (22–25). Despite this remarkable body of work, there remains no structural information about the important Notch-TMD and its flanking juxta membrane segments.

The Notch receptor is also related to Alzheimer’s disease (AD), a devastating neurodegenerative disorder that affects tens of millions of people worldwide. The most prominent model for the etiology of AD holds that the production and oligomerization/aggregation of the amyloid-β (Aβ) polypeptide in the brain is the root cause of AD pathogenesis (26). Aβ is the end product of the amyloidogenic pathway, which is initiated by β-secretase release of the large ectodomain of the full-length APP. The remaining 99-residue transmembrane C-terminal domain of APP (C99) is then cleaved by γ-secretase to release the Aβ polypeptide into the extracellular matrix and the APP intracellular domain (AICD) into the cytosol (27). It is generally believed that long-term reduction of Aβ production would reduce the risk of AD. One potential way to lower Aβ production is inhibition of γ-secretase cleavage of C99, but this strategy has thus far proven clinically problematic because of the toxicity of γ-secretase inhibitors (28). One γ-secretase inhibitor, semagacestat, made it to phase 3 clinical trials before cancellation (29). Participants in these trials experienced a variety of physical issues, including skin cancers and reduced cognitive capabilities. Inhibition of Notch receptor processing was determined to be the underlying cause of these adverse effects. The inhibition of Notch cleavage by compounds intended to inhibit Aβ release from C99 is therefore a major obstacle to the exploration of γ-secretase inhibitors as potential Alzheimer’s therapeutic agents (28, 29). Given this problem and the broad significance of Notch signaling, a close examination of the structure and biochemical properties of the Notch-TMD seems well merited. These studies are also timely in light of the fact that the structure and ligand-binding properties of C99 have recently been characterized (30–38). The stage is thereby set for a compare-and-contrast study of the structures, membrane interactions, and biochemical properties of the Notch-TMD and C99 that may provide the basis for development of improved inhibitors of Aβ production.

RESULTS
Structure of the Notch-TMD
There are significant challenges to assigning side-chain resonances in membrane proteins, including (i) rapid T2 relaxation due to the relatively large size of the complexes formed with the protein and its membrane mimic (that is, bicelles) and (ii) the huge overlapping background 1H nuclear magnetic resonance (NMR) signals from the bicelles that

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will be present in spectra unless costly perdeuterated detergents/lipids are used (39). Here, using fully protonated bicelles as the model membrane system for our studies of the Notch-TMD, we were able to assign 95% of the side-chain resonances, complementing previous backbone assignments (40). Sensitivity increases from the use of a cryogenic probe and 3-mm NMR tubes, employment of pulse sequences optimized for a slow tumbling system, and effective data processing were key factors. With these approaches, lipid and detergent signals were minimized in the spectra used to complete NOTCH-TMD side-chain assignments.

The structure of the Notch-TMD in bicelles was determined using a variety of NMR-derived structural restraints combined with XPLOR-NIH calculations [see table S1 for summary of restraints and PROCHECK analysis (41)]. The 10 lowest energy structures were deposited in the Protein Data Bank (PDB; code 5KZO) and exhibited a fairly straight transmembrane helix that spans residues 1732 to 1757. The N-terminal juxtamembrane domain is largely disordered and is connected to the TMD by a tetraproline motif, which forms a type II polyproline helix (Fig. 2A) (42).

To gain further insight into the structure and membrane interactions of the Notch-TMD, each of the 10 NMR-determined XPLOR-NIH structures was solvated in a disordered phase dimyristoylphosphatidylcholine (DMPC) bilayer and subjected to restrained molecular dynamics (rMD) simulations. The resulting 10 representative structures from these trajectories are shown in Fig. 2 (B and C), with corresponding PROCHECK results in table S1. The key features of the XPLOR-NIH structures were maintained during the rMD simulations, but additional insight was gleaned. Although the rMD structures retain the \(\alpha\)-helicity of the TMD, it is observed that the helix does have a tendency to be transiently underwound at Leu\(^{1747}\), shifting the helical register for the entire lower end of the TMD. The reentrant loop in the C-terminal JMD was maintained throughout the simulations. It is also observed that the TMD has a tendency to tilt with respect to the DMPC bilayer normal. The rMD results also provide insight into residue-specific access by lipid chains and water (Fig. 3). There is, of course, a high level of contact between TMD (residues 1732 to 1756) sites and lipid chains in the TMD (Fig. 3A). A slight shoulder in acyl chain exposure is seen on the N-terminal side of the TMD, extending from residues Pro\(^{1731}\) to Phe\(^{1737}\), which indicates that these residues remain close to the membrane surface during the course of the simulation. As expected, this plot also shows increased contact with lipid acyl chains for the cytosolic JMD reentrant loop (Leu-Trp-Phe; residues 1767 to 1769). The results for water contact show that the TM segment (residues 1732 to 1754) is nearly totally devoid of water (Fig. 3B), with the exceptions of sites 1750 and 1753, which are near the initial \(\gamma\)-secretase cleavage site at 1754.
Impact of varying membrane thickness on the Notch-TMD structure and topology

Topological changes induced by changes in membrane thickness as a consequence of varying lipid composition may regulate γ-secretase cleavage of both C99 and Notch. We previously used bicelles containing lipids with different chain lengths and chemical structures (glycerolipids versus sphingolipids) to probe how the conformation and topology of C99 depend on bicelle composition (43). This led us to conduct analogous studies of the Notch-TMD. We examined the Notch-TMD in three sets of bicelles. The first condition was DMPC bicelles in which the lipid component has two relatively short 14-carbon chains; the second was egg sphingomyelin (ESM) bicelles, dominated by sphingomyelin that contains two 16-carbon chains; and the third was milk sphingomyelin (MSM) bicelles, in which the sphingomyelin has the usual C16 ceramide chain plus a fatty amide chain that is usually 22 to 24 carbons long.

Transverse relaxation-optimized spectroscopy (TROSY) NMR spectra of the Notch-TMD were collected in these three bicelle types in the presence and absence of water- or lipid-soluble paramagnetic probes, where peak intensity ratios were measured relative to the corresponding diamagnetic reference sample. We also measured backbone 15N chemical shifts for the protein in these three bicelle types.

The site-specific backbone amide 15N chemical shifts for the protein in these three bicelle types.

The paramagnetic probe experiments revealed modest differences in model membrane interactions for the Notch-TMD under the three sets of conditions (Fig. 5). There are no significant accessibility pattern differences at the C-terminal (intracellular) end of the TM helix in all three types of bicelles tested. As with C99, a cluster of basic residues (that is, Arg-Lys-Arg-Arg-Arg) terminates the C terminus of the Notch transmembrane segment. These residues serve as an abrupt TM-stop topological motif. Notably, only slight topological differences are seen under the three bicelle conditions for the N-terminal end of the TMD and the N-terminal juxtamembrane domain. This is in stark contrast to C99 (43), where the N-terminal end of the TMD exhibited significant variations in probe accessibility because the lipid component of the bicelles was changed from having relatively short (C14) acyl chains (DMPC) to having long (C22 to C24) chains (MSM).

The q = 0.33 dihexanoylphosphatidylcholine (DHPC)–DMPC, DHPC-ESM, and DHPC-MSM bicelles used in this work are rather detergent-rich (three DHPC detergent molecules for each lipid). The q = 0.33 DHPC-DMPC bicelles have been extensively characterized (44–46), particularly with respect to the question of whether they maintain an ideal “bicelle” morphology (DHPC edge-stabilized DMPC bilayer discs) or whether, at this high DHPC content, the assemblies are better described as mixed micelles in which there is little segregation of the lipid and detergent components. Compelling recent experimental evidence favors the bilayered disc morphology (47). For the DHPC-ESM and DHPC-MSM bicelles, there is little data to support one model or the other, although the longer lipid chains of ESM and MSM relative to DMPC might be expected to confer an even higher energetic preference for formation of bilayered discs rather than mixed micelles. Regardless
of the bicelle morphologies, the paramagnetic probe results for the Notch-TMD suggest that, when the hydrophobic span of the model membrane is varied, the identities of the residues located at its water-hydrophobic interfaces remain unchanged. This suggests that the Notch-TMD most likely adapts its topology in a real membrane to varying bilayer width by varying its tilt angle with respect to the bilayer normal to maintain the same transmembrane span of residues. This is consistent with the tilting seen for the Notch-TMD in most frames of the rMD DMPC simulations (Fig. 2, B and C). We postulate that the degree of TMD tilt is determined by the energetic imperative to maintain a roughly constant balance of membrane versus water exposure for N-terminal interfacial residues regardless of bilayer width.

Cholesterol titration of the Notch-TMD in bicelles

There is much evidence that cholesterol promotes AD. Whereas this phenomenon likely involves multiple mechanisms (48, 49), one seems to be the direct promotion of the amyloidogenic pathway (50, 51). The immediate precursor of Aβ, the transmembrane C99 domain of the APP, has been shown to form a 1:1 complex with cholesterol at physiologically relevant cholesterol concentrations (30, 52). The formation of this complex has been proposed to promote γ-secretase release of Aβ from C99, possibly by promoting partitioning of C99 into lipid rafts, where γ-secretase is enriched (53, 54). The role cholesterol plays in Notch biology/biochemistry has received little attention, possibly because any such role is limited. To test whether the Notch-TMD binds cholesterol, we monitored a cholesterol titration of the protein. Five bicelle samples were prepared, spanning a range of cholesterol concentrations ranging from 0 mole percent (mol %) to 15 mol % (mol % protein relative to moles of lipid). NMR spectra were collected at each titration point and compared to see whether there is any evidence for specific cholesterol binding. The spectral overlay is shown in Fig. 6. In contrast to previous results for C99 (30), only very small Notch-TMD resonance shifts were seen. After plotting the chemical shift changes relative to mole percent cholesterol, plots for the peaks undergoing minor shifts were usually linear, consistent only with very weak binding or completely nonspecific interaction of the Notch-TMD with cholesterol.

DISCUSSION

The structure of bicelle-associated Notch-TMD fills in an important gap in the formidable body of structural data for the Notch 1 receptor (20–25). Given that Notch shares with C99 the property of being subject to intramembrane cleavage by γ-secretase, the results of this work also enable the first compare-and-contrast study of γ-secretase substrates. Accordingly, this study may shed light on the broad substrate specificity of γ-secretase and may also reveal differences between these two proteins that can be exploited in the development of Notch-sparing modulators or inhibitors of C99 cleavage.

Implications of the Notch-TMD structure for Notch signaling and for recognition by γ-secretase

The structural properties of the Notch-TMD domain are likely well adapted for roles in the Notch signaling pathway. The extracellular...
The juxtamembrane region is largely disordered except for the unusual tetra-proline motif, leading to the TMD. It seems likely that the disordered linker (Val1721 to Glu1727) and the all-trans tetraproline motif (Pro1728 to Pro1731) serve to help absorb the strain that the TMD might otherwise experience during the forced extension of the extracellular NRR domain (Fig. 1) when the Notch ectodomain engages with a membrane-anchored ligand on an adjacent cell, followed by endocytosis of the bound ligand. The extracellular N terminus of C99 is similar to that of the Notch juxtamembrane ectodomain in its disorder, but the two proteins differ in how the N terminus is linked to the TMD. C99 has a short surface-associated amphipathic helix followed by a water-exposed connecting loop to the TMD, whereas the disordered N terminus of Notch is linked to the TMD through a tetraproline segment that adopts a type II polyproline helix. These observations are consistent with the notion that γ-secretase senses the distance of the N termini of potential substrates from the membrane surface, engaging only proteins with short extracellular domains.

Within the cytosolic juxtamembrane domain of Notch, membrane surface association of the Leu-Trp-Phe1769 segment is interesting in light of its overlap with the Trp-Phe-Pro1770 motif, which is known to be essential for the binding of the RBPJ-associated molecule (RAM) domain (Fig. 1) to a transcriptional activation partner CSL (55, 56). Our results suggest that the Trp-Phe-Pro1770 motif may normally be “tucked away” via membrane surface association, making it unavailable for interaction with CSL or other proteins until the entire NICD is released from the membrane by γ-secretase cleavage. In contrast, C99 lacks any membrane-associated segment immediately following its TMD; instead, it has a disordered 40-residue aqueous loop connecting the TMD to a surface-associated amphipathic helix at the C99 C terminus (30). The very different organization of the intracellular domains of the Notch-TMD and C99 is consistent with the notion that γ-secretase recognition of substrates is tolerant of the nature of the intracellular domains of single-span membrane proteins.

The Notch-TMD is not kinked and does not have a cholesterol-binding site

The TMD of Notch is a straight helix. This is in contrast to C99, for which NMR and electron paramagnetic resonance spectroscopy studies have indicated the presence of a kink associated with a di-Gly motif located near the center of the bilayer (30).

The Notch-TMD in bicelles does not bind cholesterol under the same conditions in which C99 is known to bind cholesterol with physiologically relevant affinity to form a 1:1 complex (30, 52). This biochemical difference highlights key structural features of C99 TMD that are missing in the Notch-TMD. First, as a result of the tandem GXXG “glycine zipper” sequence, C99 has a flat surface on the extracellular half of its TMD that is well suited for binding the cholesterol...
The Notch-TMD and cholesterol is not surprising because most previous studies of specific roles for lipids in Notch signaling and trafficking imply only indirect effects (62–64).

The fact that C99, but not Notch, has a cholesterol-binding site suggests that it may be possible to develop compounds that target the cholesterol-binding site of C99 to specifically reduce its cleavage by γ-secretase without affecting cleavage of the Notch-TMD. It has been reported that a class of compounds referred to as γ-secretase modulators (GSMs) act by binding to the same glycine zipper face of the C99 TMD that is central to cholesterol binding (65). It is now believed that GSMs do not associate with significant affinity to C99 alone but instead bind only as part of a ternary complex with γ-secretase (51, 66, 67). Our observation in this work of the lack of a flat recess on the Notch-TMD may explain why GSMs do not alter cleavage of Notch by γ-secretase: GSMs likely cannot participate in a ternary complex with Notch and γ-secretase because the Notch-TMD lacks the flat recess formed by the Gly zipper sequence motif at one face of the C99 TMD.

C99 and the Notch-TMD are conformationally robust but adjust topologically in different ways to changes in model membrane hydrophobic thickness

It is known that both the rate of total Aβ production and the ratio of Aβ42 to Aβ40 production are dependent on modulation of γ-secretase activity by membrane thickness (68, 69). Moreover, there are persistent reports that γ-secretase preferentially resides in lipid rafts (53, 54), which are generally thicker than corresponding disordered phase membranes. This work illuminates how two different γ-secretase substrates adapt to changes in membrane thickness and lipid type.

NMR was used to show that the Notch-TMD resembles C99 (43) by adopting a conformational state that is remarkably insensitive to changes in bimodal composition, even when the lipid component was changed from the relatively short C14-based phosphatidylcholine (DMPC) to a form of sphingomyelin (MSM), in which the fatty amide chain is usually 22 to 24 carbons long. This result matches corresponding results for C99 (43). This conformational “robustness” for C99 and the Notch-TMD may reflect a general structural property shared by many membrane proteins, which often must maintain a functional conformational state in the face of marked cellular changes in local membrane lipid compositions (51). We suggest that changes in substrate conformation as a function of lipid composition are unlikely to be a major mechanism regulating the rate by which these substrates are cleaved by γ-secretase.

On the other hand, our results suggest that C99 and the Notch-TMD adjust their membrane topologies differently in response to changes in membrane thickness. Both proteins have a membrane domain stop site composed of consecutive basic residues located just C-terminal to their TMD. However, the Notch-TMD adapts to bimodal containing long-chain lipids in a manner such that the residues comprising its buried transmembrane span remain invariant. Although it is not presently possible to state with certainty that the bimodules used in this work are actual bilayered discs rather than classical mixed micelles, this result suggests that the Notch-TMD may tilt in actual biological membranes so as to maintain a constant span of transmembrane residues in the face of varying bilayer thickness. C99 has very different properties. In bimodules of varying dimensions, the N-terminal start of the C99 TM helix seems to remain fixed in position such that when the span of the model membrane changes, the identity of the residue that starts the TMD adjusts—residues that are within the model membrane when it is thick (Asn698–Lys–Ala–Ile–Ile603) extend into the aqueous phase in thinner model membranes (43). This segment is evidently energetically tolerant of being located at different positions along the bilayer normal in the interfacial region.

The different ways Notch and C99 adjust their topologies in response to changes in membrane thickness may regulate how they initially engage with γ-secretase and the location of the subsequent substrate cleavage site. For C99, it is well established that, although γ-secretase often initially cleaves (“e-cleavage”) after Thr719, the substrate can also be cleaved at Leu720 (70). Following initial cleavage and release of the AICD, it is believed that the enzyme then processively clips tri- and tetrapeptides from the C-terminal end of the TMD until either Aβ40 or Aβ42 is released into the extracellular space, with Aβ40 being released from C99 that was initially cleaved at Thr719. In contrast, Aβ42 is released from C99 that was originally clipped at Leu720 (71). It has been shown that increasing lipid chain lengths to increase the bilayer width results in an increase in the Aβ40/Aβ42 ratio (68). It has also been shown that adding a pair of nonpolar residues to the C-terminal end of the C99 TMD leads to a decreased Aβ40/Aβ42 ratio, whereas adding nonpolar residues to the N-terminal end of C99 had the opposite effect (72). These results are consistent with the notion that the active site of γ-secretase cleavage is fixed with respect to the center of the membrane. In the case of C99, the fact that its TMD span adjusts asymmetrically with respect to the center of the bilayer as membrane thickness changes implies that the preferred γ-secretase cleavage sites in C99 will vary, consistent with the published results summarized above. In the case of Notch, cleavage is also processive, usually starting at Val1754 and proceeding up to Ala1755, and that mutating the C-terminal juxtamembrane lysine to arginine tends to favor cleavage at this site (73). This is consistent with the notion that the initial γ-secretase cut site in both Notch and C99 is determined, at least partly, by site position with respect to the center of the bilayer. A testable prediction of this model is that γ-secretase will tend to cleave the wild-type Notch-TMD at the same sites, regardless of bilayer thickness, if its TMD responds to changes in membrane thickness by altering its tilt—an adjustment that does not change relative residue positions with respect to the center of the bilayer.

CONCLUSIONS

The results of this work help fill in an important gap in the structural biology of the Notch receptor and also highlight differences between the
Notch-TMD and C99, which include very different conformations, ligand-binding properties, and modes of topological adaptation to varying model membrane thickness, as summarized in Fig. 7. These differences point to the versatility of γ-secretase in terms of being able to recognize a range of type I single-span membrane proteins with significantly different properties. These differences may also help inform the development of compounds that specifically target integrin receptor TMDs (74, 75), in part, by exploiting the unique structural features of the GXXG motifs found in integrin TMDs and those found in C99 but not in Notch.

**MATERIALS AND METHODS**

**Expression and purification of the N-terminally His₆-tagged Notch-TMD for NMR spectroscopy**

The Notch-TMD (residues 1721 to 1771; Fig. 1) was expressed and purified into bicelles, as previously described (40), with the solution containing 2% (w/v) DMPC/DHPC bicelles at q = 0.33 (where q equals the lipid-to-detergent mol/mol ratio) in 300 mM imidazole plus 10 mM dithiothreitol (DTT; pH 7.8). The protein-bicelle assemblies were concentrated by centrifugal ultrafiltration, including a 10x buffer exchange cycle, where 15 mM DHPC in 50 mM phosphate and 1 mM EDTA (pH 6.5) was added to a concentrated Notch-TMD solution. The sample was then concentrated 7.5 times to generate approximately 15% bicelles. D₂O (10%) was then added, and the pH was reduced to 5.5 using acetic acid, followed by transfer of 180 µL of the solution to a 3-mm NMR tube. Typical final NMR conditions were ~0.5 mM Notch-TMD, 15% DMPC/DHPC bicelles (q = 0.33), 20 mM phosphate, 65 mM imidazole, 2 mM DTT, and 1 mM EDTA (pH 5.5), with 10% D₂O. The concentration of the Notch-TMD was spectrophotometrically determined at 280 nm based on its extinction coefficient of 6990 M⁻¹ cm⁻¹.

**Side-chain NMR resonance assignments**

The U-¹⁵N,¹³C Notch-TMD was prepared for NMR, as previously published (40) and as described above. Backbone resonance assignments from our previous work (40) were also used. Experiments were carried out on a Bruker AV-III 600-MHz spectrometer equipped with inverse detection cryoprobes. Standard Bruker pulse sequences and their hard-coded delays were used for all experiments. Key experiments for the side-chain assignments were the TROSY- (H)(C)(CO)NH-TOCSY (trhccconhgp3d3) and H(C)(CO)NH (hccconhgp3d2) (76–78). Acquisition for both experiments used a 12-ms DIPSI-2 mixing time for the carbon TOCSY, waltz65 proton decoupling, and gap4 nitrogen decoupling using field strengths of 9.6, 3.6, and 1 kHz, respectively. Relaxation delays were 1.2 s for the TROSY and 1.0 s for the proton-proton experiments. The sweep widths were as follows: 7812 Hz for the direct proton dimension, 1460 Hz for the indirect nitrogen dimensions, and 12074 Hz for the indirect carbon or 5402 Hz for the indirect proton dimension with 2048, 48, and 126 complex points, respectively. Carriers were set to 4.7 parts per million (ppm) (¹H), 116.5 ppm (¹³N), and 40 ppm (¹³C). Echo/anti-echo detection was used for the nitrogen, and a States-TPPI scheme was used for proton or carbon in the indirect dimension. Water suppression was achieved through gradient coherence selection, with an additional 1-ms sinc-shaped flip-back pulse and a WATERGATE (79) sequence immediately before data acquisition in the carbon TOCSY. To minimize relaxation during acquisition, these elements were omitted in the proton TOCSY, still yielding excellent suppression of all unwanted signal. The carbon TOCSY was acquired with 32 scans per increments, whereas only 16 scans were used for the proton TOCSY, yielding in a total experiment time of 3 days for each experiment.

For the standard Bruker HCCH-TOCSY (hchdhpip3d2) (80, 81), a CPQCI probe was used. The parameters were as follows: Sweep width in the indirect proton dimension was 7812 Hz with 2048 complex data points, and both indirect carbon dimensions were at 9604 Hz and 96 complex points. A DIPSI3 spinlock of 22.6 ms at 9.3-kHz field strength and a 3.9-kHz carbon decoupling field during data acquisition was applied. A States-TPPI scheme was used in both indirect dimensions, using 16 transients with a 1.5-s recycle delay per increment. Water suppression was achieved through gradient coherence selection only. The total experiment time was 70 hours.

All NMR spectra were processed with NMRPipe (82) and analyzed with SPARKY (83). Residual water suppression consisted of a time domain polynomial subtraction. Indirect dimensions were linearly predicted to twice the data size before Gaussian window functions were applied. Polynomial baseline corrections were generally applied in the direct and indirect ¹⁵N dimensions. Residue-specific ¹H and ¹³C resonances from the H(CCO)NH- and C(CO)NH-TOCSY experiments were correlated via peaks observed in the HCCH-TOCSY spectrum. Side-chain resonances were assigned by correlations from existing sequential backbone assignments (40). In the few occurrences where ¹H-¹³C-correlated chemical shifts were observed but could not be unambiguously assigned, nuclear Overhauser enhancement spectroscopy (NOESY) data typically resolved ambiguity. The unassigned, although clearly distinguishable, proline chemical shifts in the HCCH-TOCSY spectrum were used to ascertain the presence of all-trans configurations for all prolines based on their C₅ and C₇ chemical shifts (84).

**Distance restraints from paramagnetic relaxation enhancement measurements**

The wild-type Notch-TMD has one cysteine residue (Cys¹⁷⁵₂; Fig. 1). For paramagnetic relaxation enhancement (PRE) measurements,
samples were prepared in which the native cysteine was spin-labeled. Samples were also prepared where the native Cys\textsuperscript{1752} was mutated to Ser, and Phe\textsuperscript{1744} was mutated to Cys and then spin-labeled. Both wild type and the single-Cys mutant were purified into 0.2% n-dodecylphosphocholine (DPC) and spin-labeled using an established method (30, 85). Briefly, each single-Cys form was concentrated to 0.5 mM, and the pH was lowered to 6.5 followed by addition of 2.5 mM DTT to ensure cysteine thiol reduction. The protein was then spin-labeled using the thiol-reactive probe S-(1-oxyl-2,2,5,5-tetramethyl-2,5-dihydro-1H-pyrrol-3-yl)methyl methanesulfonothioate (MTSL; Toronto Research Chemicals). A 20-fold excess of MTSL was added relative to ~0.5 mM Notch-TMD [buffer: 65 mM imidazole, 2 mM EDTA, 2.5 mM DTT, and 0.2% DPC (pH 6.5)]. The labeled sample was incubated overnight, followed by buffer exchange with 50 mM sodium phosphate buffer (pH 6.5) to remove the imidazole in preparation for metal ion affinity chromatography. The solution was then mixed with Ni-agarose gel, washed with 25 column volumes of 50 mM phosphate and 0.2% DPC (pH 7.8) to remove excess MTSL and MTSL-modified DTT. The resin was then rinsed with a 8 × 1 column volume of DHCPC in water, followed by 4 × 1 column volume exchange using a 2% (w/v) DMPC/DPHC bicelle mixture (q = 0.33) before the protein was eluted in the same 2% bicelles plus 300 mM imidazole at pH 7.8. The sample was then immediately exchanged into the 20 mM sodium phosphate and 65 mM imidazole NMR buffer, and the pH was reduced to 5.5 using glacial acetic acid. \textsuperscript{1}H-\textsuperscript{15}N TROSY NMR data were acquired at 900 MHz for the spin-labeled samples. In each case, a spectrum was collected, followed by removal of the paramagnet and reacquisition of a parameter-matched TROSY spectrum for the now-diamagnetic sample. Reduction of the spin label was carried out by adding ascorbic acid to 20 mM to remove excess MTSL and MTSL-modified DTT. The sample was then rinsed with a 8 × 1 column volume of DHCPC in water, followed by 4 × 1 column volume exchange using a 2% (w/v) DMPC/DPHC bicelle mixture (q = 0.33) before the protein was eluted in the same 2% bicelles plus 300 mM imidazole at pH 7.8. The sample was then immediately exchanged into the 20 mM sodium phosphate and 65 mM imidazole NMR buffer, and the pH was reduced to 5.5 using glacial acetic acid. \textsuperscript{1}H-\textsuperscript{15}N TROSY NMR data were acquired at 900 MHz for the spin-labeled samples. In each case, a spectrum was collected, followed by removal of the paramagnet and reacquisition of a parameter-matched TROSY spectrum for the now-diamagnetic sample. Reduction of the spin label was carried out by adding ascorbic acid to 20 mM to remove excess MTSL and MTSL-modified DTT. The sample was then rinsed with a 8 × 1 column volume of DHCPC in water, followed by 4 × 1 column volume exchange using a 2% (w/v) DMPC/DPHC bicelle mixture (q = 0.33) before the protein was eluted in the same 2% bicelles plus 300 mM imidazole at pH 7.8. The sample was then immediately exchanged into the 20 mM sodium phosphate and 65 mM imidazole NMR buffer, and the pH was reduced to 5.5 using glacial acetic acid. 

**Structure calculations**

Restraints used in structure calculations included using PRE-derived distances, backbone torsion angles derived from chemical shifts, and NOE distance restraints. Backbone C\textsubscript{α}, C\textsubscript{β}, CO, and N chemical shifts (40) were input into TALOS+ to generate dihedral angle restraints (85, 87). Chemical shift index analysis was also used to generate hydrogen bond restraints for the TM segment (88). These backbone dihedral angle and hydrogen bonding restraints were calculated from the backbone chemical shift values (89) and used in structure calculations. The backbone dihedral angles of the all-trans tetraproline motif were initially set to -79° and 149° (42), with generous errors allowed. A three-dimensional (\textsuperscript{1}H,\textsuperscript{1}H,\textsuperscript{15}N)-TROSY-NOESY experiment was carried out on the U-\textsuperscript{15}N Notch-TMD (100-ns mixing time) in DHCPC/DMPC bicelles under the standard NMR conditions indicated above to obtain 339 short- and medium-range NOE restraints used in structure calculations. The PRE restraints were very conservatively used, as previously described (39, 85). Briefly, each restraint was classified as “close in space” if the paramagnetic/diamagnetic intensity ratios were less than 0.15, in which case the paramagnetic probe and the target amide proton were assigned as being between 2 and 19 Å apart. Resonances with ratios between 0.15 and 0.85 were converted to explicit distances, as previously described (85, 86), and given generous uncertainties of ±7 Å. Resonances that exhibited little to no PRE effects (intensity ratios greater than 0.85) were loosely restrained to be between 19 and 100 Å apart. The PRE restraints were implemented as NOE-like restraints from the spin label to the backbone amide hydrogen (39, 90).

Structure calculations were conducted using XPLOR-NIH v2.24 (91), similar to previously published works (30, 85). Simulated annealing was carried out with 15,000 steps at 3500 K, with cooling to 100 K. During the temperature-cooling ramp, the van der Waals force constant was varied from 0.004 to 4 kcal mol\textsuperscript{-1} Å\textsuperscript{-2}. Similarly, force constants were increased for the NOE and PRE restraints from 1.0 to 30.0 kcal mol\textsuperscript{-1} Å\textsuperscript{-2}. High-temperature simulated annealing was followed by torsion angle and full-atom minimization steps. The lowest energy structures were selected, and the spin-labeled sites associated with the PRE distance restraints were mutated back to their wild-type residue types, followed by energy minimization to allow their side-chain conformations to optimize. Additional details regarding the structure calculation are given in table S1. The 10 XPLOR-NIH models (fig. S1) were deposited in the PDB (code 5KZO).

XPLOR-NIH–based structure determination was followed by AMBER14 rMD to refine the 10 top-scoring Notch-TMD XPLOR-NIH structures in a lipid bilayer. Structures were solvated in an explicit DMPC bilayer, and simulations included the use of NMR-derived distance restraints (85). Simulations used the Lipid14 AMBER lipid force field (92) and the f82SB AMBER force field. Refinement of each Notch-TMD structure was conducted via a 60-ns production rMD. For each of the 10 trajectories, the average Notch-TMD structure was calculated, and the frame that best matches the average structure was selected as the “representative model” from that trajectory using CPPTRAJ (93). Both the initial X-PLOR structures and the AMBER refinement structures were validated by PROCHECK (see table S1) (41). We also used CPPTRAJ (93) to determine water penetration and protein-lipid interactions. Additional technical details regarding the rMD calculations are provided in the Supplementary Materials.

**Dependence of protein structure and topology on varying bilayer thickness**

We have previously described methods to determine paramagnetic probe accessibility to membrane protein sites in bicelles containing lipids of varying chain lengths (43). Briefly, DMPC (C\textsubscript{14} chains), ESM (mostly C\textsubscript{16}), or MSM (mostly C\textsubscript{22} to C\textsubscript{24}) was weighed as a solid into a glass vial. A 2% bicelle solution was generated by adding the appropriate amount of DHCPC and water to form a q = 0.33 bicelle mixture that was then used to equilibrate and elute the Notch-TMD from a Ni(II)-resin column. NMR samples of the U-\textsuperscript{15}N Notch-TMD were prepared as described above, concentrated, and split into three parts. The first sample was a paramagnet-free reference sample. The second contained a water-soluble paramagnet, Gd(III)–diethylenetriamine pentaacetic acid [Gd-DTPA; added to 2 mM from a 0.1 M stock solution (pH 5.5)]. The third sample contained the lipophilic paramagnetic probe, 16-doxylstearic acid (16-DSA; 2 mM; Santa Cruz Biotechnology). The 16-DSA sample was prepared by adding an aliquot of 16-DSA stock (2.5 mg/ml) in methanol into a tube and then removing the methanol in a speed vacuum. The protein sample was then added to the dried 16-DSA and gently mixed (to a concentration of 2 mM) before it was transferred into a 3-mm NMR tube. TROSY spectra (900 MHz) were collected for each bicelle condition using a delay time of 4 s between scans to ensure complete relaxation between scans. TROSY
peak intensities were measured for the two paramagnet-containing samples and compared to the corresponding peak intensities from the parameter-matched reference sample. The ratios of the peak intensities were plotted as an indicator of site access to the paramagnetic probes.

### Cholesterol titration of the Notch-TMD

Samples were prepared essentially as previously described for C99 (30). Briefly, cholesterol-containing bicelles were prepared by dissolving DMPC and cholesterol in chloroform at the desired molar ratio and then removing all solvent by high vacuum, followed by mixing the cholesterol/phospholipid film with DHPC and water to form a 2% (w/v) bicelle solution containing 15 mol% cholesterol (relative to total moles of DHPC + DMPC + cholesterol). This mixture was subjected to freeze-thaw cycles until a clear solution was formed, aliquots of which were then mixed with cholesterol-free bicelles to make a series of q = 0.33 bicelles at 15% (w/v) that contained 0, 2.5, 5, or 10 mol% cholesterol in water. Each of these solutions was used to equilibrate (with no imidazole) and then elute (with 300 mM imidazole) the Notch-TMD from the resin. Samples were then prepared for NMR, as described above, and contained 0.35 mM protein. A \(^{15}\)N-\(^{1}H\) TROSY was collected at 318 K for each cholesterol titration point on a Bruker AV-III 900-MHz spectrometer equipped with a CPTCI cryogenic probe.

### SUPPLEMENTARY MATERIALS

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/3/4/e1602704/DC1

Supplementary Methods

fig. S1. Top 20 NMR-determined XPLOR-NIH structures of the Notch-TMD.

table S1. Statistics for Notch-TMD structure determination and structure quality for the 10 final XPLOR-NIH NMR structures and for the 10 representative restrained MD structures.

References (94–96)

### REFERENCES AND NOTES


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Structural and biochemical differences between the Notch and the amyloid precursor protein transmembrane domains

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