Effect of Src Kinase Phosphorylation on Disordered C-terminal Domain of N-Methyl-D-aspartic Acid (NMDA) Receptor Subunit GluN2B Protein*

Received for publication, May 6, 2011, and in revised form, June 17, 2011 Published, JBC Papers in Press, June 28, 2011, DOI 10.1074/jbc.M111.258897

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NMDA receptors are ligand-gated ion channels with a regulatory intracellular C-terminal domain (CTD). In GluN2B, the CTD is the largest domain in the protein but is intrinsically disordered. The GluN2B subunit is the major tyrosine-phosphorylated protein in synapses. Src kinase phosphorylates the GluN2B CTD, but it is unknown how this affects channel activity. In disordered proteins, phosphorylation can tip the balance between order and disorder. Transitions can occur in both directions, so it is not currently possible to predict the effects of phosphorylation. We used single molecule fluorescence to characterize the effects of Src phosphorylation on GluN2B. Scanning fluorescent labeling sites throughout the domain showed no positional dependence of the energy transfer. Instead, efficiency only scaled with the separation between labeling sites suggestive of a relatively featureless conformational energy landscape. Src phosphorylation led to a general expansion of the polypeptide, which would result in greater exposure of known protein-binding sites and increase the physical separation between contiguous sites. Phosphorylation makes the CTD more like a random coil leaving open the question of how Src exerts its effects on the NMDA receptor.

Ionotropic glutamate receptors are central to mediating the plasticity that underlies higher brain functions like learning and memory and development of neural circuits (1). NMDA-sensitive glutamate receptors (NMDAR)² are ligand-gated ion channels with modular functional components that include extracellular N-terminal and ligand-binding domains, a transmembrane ion channel, and a regulatory intracellular C-terminal domain (CTD) (Fig. 1A) (2). Although one can measure ligand-gated channel activity without the CTD (3), it is central to NMDAR regulation in neurons. The CTD regulates membrane expression and synaptic clustering (1, 4), alters gating and ion selectivity of the channel pore (2), and acts as a scaffold for intracellular signaling molecules (e.g. Ca²⁺/calmodulin-dependent kinase 2 (CAMKII)). The activity of the CTD in these different functional roles depends on the state of post-translational modifications (2), but we lack a mechanism to describe how the CTD influences the ligand-gated channel.

NMDARs are obligate heterotetramers being composed of the ubiquitous GluN1 subunit and some combination of GluN2 and/or GluN3 subunits. In the cortex, NMDARs typically contain GluN2A or GluN2B subunits (5). In GluN2A and GluN2B, the CTD is the largest single domain in the protein (Fig. 1A) (6). Genetic deletion of the GluN2B CTD is lethal (7), which is equivalent to deleting the entire receptor (8).

The CTD allows many signaling pathways to modulate GluN2B. A central palmitoylation motif divides the CTD into two “domains” (CTD1 and CTD2, Fig. 1B) (9). This lipid modification would drive the N-terminal membrane attachment of CTD2 (10), and it affects trafficking and synaptic localization of NMDARs (9). The CTD also interacts with scaffold proteins that direct trafficking and synaptic retention (11–13).

The GluN2B CTD is the target of multiple kinases (14) and the major tyrosine-phosphorylated protein in the postsynaptic density (15). Somehow phosphorylation near the distal C terminus affects the ion channel. For example, phosphorylation by Src family tyrosine kinases enhances NMDAR currents (16, 17) and reduces Zn²⁺ inhibition (18, 19).

The CTD of GluN2B is predicted to be intrinsically disordered by both PONDR (20) and IUPRED (Fig. 1B) (21). Intrinsically disordered proteins lack fixed secondary or tertiary structure and instead dynamically sample an ensemble of different conformations (22). Phosphorylation sites are frequently located in disordered regions, even in proteins with structured domains (23). Phosphorylation can promote structural transitions between order and disorder (24). Structure can be induced in disordered regions (25) or structure can be lost (26). Phosphorylation can reduce tertiary contacts (27) or increase intramolecular dynamics (26) within the native state ensemble. Increasing the charge of disordered peptides can favor more extended states (28–30). However, some proteins show no global expansion upon phosphorylation (31).

Recently, we provided the first experimental confirmation of disorder in CTD2 (Fig. 1B) from GluN2B (32). However, CTD2 does not show the rapid intramolecular dynamics of a random coil but rather undergoes conformational switching on the second time scale (32). If the structure dictates function then the function would be changing every second. Switching would...
allow “conformational selection,” wherein a productive interaction could only occur if the active conformation is present when the ligand arrives (33).

In this study, we describe the effect of Src phosphorylation on CTD2 from the GluN2B subunit, which contains both physiological Src sites (Tyr-1336 and Tyr-1472). Single molecule fluorescence measurements showed no positional dependence of the fluorescence resonance energy transfer (FRET) efficiency. Instead, FRET only scaled with the separation between labeling sites suggesting a lack of local structure.

Phosphorylation led to a general expansion of the polypeptide rather than inducing stable structure. Expansion would result in greater exposure of potential protein-binding sites, which could change interactions within the receptor or with other synaptic proteins. Phosphorylation did not stabilize an “active” conformation instead making the CTD more random coil-like. How the CTD affects channel function remains an open question.

**EXPERIMENTAL PROCEDURES**

**Protein Expression and Purification**—The C-terminal domain 2 of GluN2B (residues 1259–1482, Fig. 1B) was cloned into pPROEX HTB (Invitrogen) and expressed in the Rosetta strain of *Escherichia coli* (EMD Biosciences, San Diego). The His<sub>6</sub> fusion was purified under reducing and denaturing conditions as described (32). The protein was exchanged to non-denaturing conditions by extensive washing while bound to nickel-nitriiotriacetic acid-agarose. The His<sub>6</sub> tag was removed with tobacco etch virus protease, and the protein was further purified with cation exchange and size exclusion (GE Healthcare). Chicken c-Src kinase (residues 83–534) was kindly provided by Dr. Markus Seeliger. Proteins were phosphorylated in the presence of 1 mM ATP and 1 mM MgCl<sub>2</sub> followed by cation exchange to remove Src.

**Limited Proteolysis**—Wild-type CTD2 was purified and phosphorylated as described above. Sequencing grade trypsin and chymotrypsin were added at 0.002 mg/ml (Promega, Madison, WI). Replicate samples were prepared for each time point and inhibited at the indicated time with 0.5 mm phenylmethylsulfonyl fluoride (PMSF).

**Circular Dichroism Spectroscopy**—Circular dichroism (CD) spectra were collected using a Chirascan CD spectrometer (Applied Photophysics, Leatherhead, UK at 0.2 mg/ml protein concentration in PBS (10 mm sodium phosphate, 50 mm NaCl, 0.5 mm EDTA, pH 7.4) at 20 °C. The step size was 1 nm, and the signal averaging time was 0.3 s.

**Analytical Size Exclusion Chromatography**—Purified proteins were injected onto a Shodex KW-802.5 column in 20 mM Tris, 150 mm NaCl, 0.5 mm EDTA, pH 7.5, at 4 °C as described (32). Radius of hydration was determined using a calibration equation of 1 mM ATP and 1 mM MgCl<sub>2</sub> followed by cation exchange and size exclusion (GE Healthcare). The Cys cluster II, residues Cys-1215 to Cys-1245) divides the CTD into two halves, which we have termed CTD1 and CTD2 (Fig. 1B). CTD2 could be purified under non-denaturing conditions indicating that the protein is not expressed in inclusion bodies (data not shown). CTD2 was prone to degradation during the purification, which could be removed by size exclusion. We lysed the cells in denaturants to avoid this complication and “refolded” the protein by exchanging to physiological buffer while bound to beads. Measurements on denaturant and native
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**FIGURE 1. Disorder in the C-terminal domain of GluN2B.** A, known and unknown portions of the glutamate receptor structure. A schematic representation of the x-ray structure of an AMPA-sensitive glutamate receptor subunit (3KGO) (38) is depicted within a membrane (gray). The disordered C-terminal domain is drawn in approximate scale (circle). The CTD volume was calculated based on the assumption of a compact disordered globule (22), which is supported by our data. B, prediction of intrinsic disorder in the CTD. Probability of disorder is plotted for residues 838–1482. We analyzed the CTD with two major disorder predictors PONDR (dashed line) (20) and IUPRED (solid line) (21). Plot shows probability of disorder for each residue. The location of the palmitoylation sites is indicated by the gray vertical bars crossing the panel. The second palmitoylation site defines an intervening membrane binding region that partitions the CTD into two domains that we term CTD1 and CTD2. The location of the phosphorylation sites are inset at the bottom of the panel as black vertical bars.

purified proteins were identical indicating that the use of denaturants is not causing the intrinsic disorder (data not shown). All measurements were made in biological buffers at physiological pH with sodium chloride. Although these are not the physiological counter ions for intracellular proteins, measurements made in a range of counter ions showed little difference at physiological ionic strengths (data not shown).

*Ensemble Characterization of CTD2 Phosphorylation—* Nakazawa et al. (35) demonstrated in 2001 that Src robustly phosphorylates recombinant GluN2B CTD constructs in vitro. To phosphorylate CTD2, we used purified recombinant c-Src (residues 83–534), which contains Src homology 2 and 3 and kinase domains. SDS-PAGE was used to confirm phosphorylation of all constructs. CTD2 is 24.9 kDa but runs anomalously on SDS-PAGE, in coincidence with the 30-kDa marker (Fig. 2A). Src phosphorylation of CTD2 induced a discernible decrease in the electrophoretic mobility on SDS-PAGE (*lanes marked +*, Fig. 2A). The shift from phosphorylation was dependent on the presence of Mg2+ as Src in the presence of ATP alone had no effect. To confirm the sites of Src phosphorylation, we mutared Tyr-1336 and Tyr-1472 to threonine (*ΔY*), which eliminated the mobility shift induced by Src (*right lanes*, Fig. 2A).

Limited trypsin proteolysis of CTD2 with 2 μg/ml trypsin resulted in complete degradation in less than 30 min as expected for a disordered protein (Fig. 2B). High molecular weight intermediate bands are visible at early time points indicating that some cleavage sites are partially protected. CTD2 is degraded more slowly by chymotrypsin with clear protected intermediates lasting close to 2 h (Fig. 2C). This suggests differential exposure of lysine and arginine residues (trypsin) relative to aromatic residues (chymotrypsin).

Phosphorylation did not alter the kinetics of digestion of CTD2 by either protease. However, the structural changes induced by phosphorylation altered the pattern of digestion. This is seen in the loss of most of the intermediate bands in the trypsin digestion, indicating greater exposure of cleavage sites (Fig. 2B). Similarly, phosphorylation also changed the digestion pattern for chymotrypsin (Fig. 2C). In particular, the ~14 kDa band is destabilized after phosphorylation, and a new higher molecular weight band appears ~17 kDa. All intermediate
digestion products are shifted toward higher molecular weight indicating that the phosphorylation sites are present in these products.

To identify any changes in secondary structure, we used circular dichroism (Fig. 3). The spectrum for CTD2 shows no double minima at 208 and 222 nm indicating a lack of stable α-helices (open circles, Fig. 3). The negative peak at 200 nm is consistent with intrinsic disorder, but the slightly negative ellipticity above 210 nm suggests some β-sheet character. Phosphorylation had little effect on the CD spectrum (filled circles, Fig. 3) indicating that Src phosphorylation does not induce secondary structure in CTD2.

To measure the ensemble radius of hydration ($R_H$) we used analytical size exclusion chromatography (SEC). As in SDS-PAGE, CTD2 elutes from SEC with a much larger apparent molecular weight (78.8 ± 1 kDa). The calculated $R_H$ value of 3.5 ± 0.02 nm for wild type CTD2 was consistent with a compact globular form of disorder rather than an extended random coil (22). Src phosphorylation increased the $R_H$ of wild type CTD2 by 10% (to 3.9 ± 0.02 nm) suggesting that the negatively charged phosphate moieties are causing the polypeptide to expand. This is surprising because, based on the amino acid sequence, the net charge of CTD2 is +9, so phosphorylation actually reduces the net charge.

A 10% increase in $R_H$ corresponds to a 30% increase in the effective volume occupied by the same amount of polypeptide, which implies much greater solvent exposure of the core. A folded protein of this size would have a $R_H$ of only ~2.3 nm, which results in a much higher packing density. After phosphorylation, the measured $R_H$ for CTD2 is close to that predicted for a random coil of the same molecular weight (4.1 nm) (22).

Single Molecule FRET Characterization of CTD2 Phosphorylation—We created a series of CTD2 constructs containing two unique cysteines for fluorescent labeling (Fig. 4A). The natural cysteine residues were used in some FRET pairs and are denoted by a capital C (e.g. C1394 in Table 1). Six other sites encoding for serine or threonine were chosen and mutated to cysteine (Table 1). To measure the conformation with FRET, we created eight double cysteine mutants based on these nine single labeling sites (Fig. 4A). We used SEC to confirm that neither the mutations nor fluorescent labeling affected the ensemble $R_H$. The mean $R_H$ for the labeling mutants was 3.3 ± 0.1 nm without dyes and 3.2 ± 0.1 nm after labeling. After phosphorylation, the mean $R_H$ for the labeling mutants was 3.9 ± 0.1 nm without dyes and 3.8 ± 0.1 nm after labeling. In contrast to the large increase in $R_H$ induced by phosphorylation, the introduction of dyes has comparatively little effect.

To confirm that selected labeling sites did not affect physical properties of the dyes, we made ensemble measurements of donor anisotropy, which showed little difference from free dye (data not shown). Similarly, the donor quantum yield varied by less than 20% between sites (data not shown). Despite this finding, we are hesitant to interpret the FRET changes a distance given the following: 1) the uncertainty in relationship between the dye and the protein backbone and 2) the uncertainty of an appropriate model for the underlying dynamics.

Overlapping labeling site pairs were created to fully sample all of CTD2 in an effort to reveal any local structure (Fig. 4A). Samples were numbered in order with the increasing separation between labeling sites (Table 1). Data from single molecules containing a validated single donor and single acceptor dye pair were recorded at 100 ms/frame (Fig. 4B). The background corrected FRET was calculated for each frame of the movie (i.e. point wise) until photobleaching occurred (10–100 s). Histograms contain hundreds of single molecules and were replicated in separate experiments. In contrast to the narrow gaussian smFRET distribution of a random coil (32), all the labeling combinations showed broad distributions spanning the range of middle FRET values (Fig. 4C). Although the constructs sample different regions of CTD2, there was little dispersion in the FRET peaks suggesting a lack of local structure.

After Src phosphorylation (of fluorescently labeled protein), all labeling constructs showed a trend toward lower FRET values in the smFRET distributions (Fig. 5A, solid curves). Only three of the samples (NRB-3, -4, and -8) showed a statistically significant decrease in median FRET (Table 2). The different constructs are variably located relative to the phosphotyrosines (Fig. 4A) but show the same effect. The dyes did not affect Src phosphorylation as an identical electrophoretic mobility shift was induced in labeled protein (data not shown).

We plotted the median FRET value for the predominant peak in each histogram against the number of amino acids separating the labeling sites (Fig. 5B). FRET in CTD2 showed a simple linear dependence on the separation (Fig. 5B, open circles). The different constructs sampled regions throughout the protein but revealed this simple dependence rather than positional differences in the conformation of CTD2. After phosphorylation, the median FRET remained a linear function of the separation with a nearly identical slope to unphosphorylated CTD2 (Fig. 5B, solid squares). However, the intercept of the line had shifted toward a lower FRET. Even after phosphorylation, the median FRET values were much higher than expected for an extended random coil (32).

Phosphorylation did not have any effect on slow conformational dynamics in CTD2 (supplemental Fig. S1). All labeling
constructs showed stochastic changes in FRET efficiency. As done previously (32), we characterized the FRET transitions using a Canny edge detection method (36). All constructs had a similar transition frequency in the range of 0.5 to 0.8 s\(^{-1}\) (data not shown). The transitioning was independent of the fluorophore separation and not affected by phosphorylation (Table 3).

The one exception was NRB-1, which showed a narrow FRET peak that was not shifted by phosphorylation, and it did not show stochastic changes in FRET efficiency. NRB-1 has

![Diagram showing single molecule FRET measurements of the GluN2B CTD2 conformation. A, constructs used in this experiment. Eight constructs of CTD2 were created that contained two unique cysteines for fluorescent labeling. Constructs were numbered in order of their fluorophore separation with NRB-1 being the shortest at 15 residues and NRB-8 being the longest at 172 residues (Table 1). The length and position of each construct are depicted as a bar above a scale representation of the primary sequence. The constructs overlap to sample the length of the polypeptide. The positions of cysteine residues used for labeling are indicated above the scale bar. Natural cysteines are indicated by capitalization. Phosphorylation sites are depicted below the scale bar. B, representative single molecule intensity time traces for unphosphorylated CTD2 (top row) and phosphorylated CTD2 (bottom row). Acceptor emission intensity is colored red. Donor emission intensity is colored green. Panels in the left column show representative single molecule intensity time traces for the control sample N2B-1 measured using the encapsulation method. Panels in the middle and right columns show representative single molecule intensity time traces of N2B-3 using the encapsulation and surface immobilization method, respectively. Additional representative data for the remaining samples are shown in supplemental Fig. S1. C, histograms of single molecule FRET for the eight labeling site combinations in CTD2. The histogram contains FRET values calculated for each 100 ms frame of observation. Histograms contain hundreds of molecules and were replicated in separate experiments. FRET is presented as normalized probability. Histograms are colored from red to violet in order of increasing fluorophore separation. N2B-1, red; N2B-2, orange; N2B-3, yellow; N2B-4, dark green; NRB-5, light green; N2B-6, cyan; N2B-7, blue; and N2B-8, violet.
only 15 residues between labeling sites so high FRET is expected for such a short dye separation. This short polypeptide segment should not be capable of large conformational changes and is N-terminal to both phosphotyrosines. That we do not see FRET transitions or shifts confirms that these phenomena arise from CTD2 structural dynamics and not photophysics of the dye pair.

Effect of Surface Tethering on smFRET in CTD2—In the GluN2B subunit, CTD2 would be anchored to a lipid bilayer via an N-terminal palmitoylation motif. Having one end fixed could provide a significant restriction on the accessible conformations because the N and C termini can no longer cross paths. To examine the effect of surface attachment on conformational dynamics, we replaced the N-terminal palmitoylation motif with a biotinylation site. The resultant biotinylated protein was attached directly to a passivated microscope slide using streptavidin. To prevent interactions with the slide, we used an inert proteolipid surface composed of biotinylated bovine serum albumin followed by the application of a dilute solution of protein-free liposomes. The resulting surface shows extremely low levels of nonspecific binding (supplemental Fig. S2).

We measured smFRET from the same labeling site combinations used in encapsulation studies (Fig. 6). Src phosphorylation was carried out in solution after fluorescent labeling but before immobilization. Similar effects on smFRET were observed when Src (along with ATP and Mg\(^{2+}\)) was added to immobilized protein in the sample chamber (data not shown). When CTD2 is surface-immobilized, the shift induced by phosphorylation is much more apparent (solid curves, Fig. 6A). The entire distribution shifts visibly toward low FRET. In contrast to the encapsulated samples, all constructs showed a statistically significant shift in median FRET efficiency (Table 4). Immobilization of CTD2 appeared to magnify the effectiveness of phosphorylation as a signal by further separating the two states. As expected, NRB-1 remained the exception with a narrow FRET peak, with no FRET shift upon phosphorylation and no FRET transitions.

Transitions were unaffected by tethering (data not shown). Surprisingly, it is the unphosphorylated state that is most affected by N-terminal tethering (Tables 2 and 4). Surface immobilization had no effect on median smFRET after phosphorylation although smFRET before phosphorylation shows a trend toward higher FRET values. This suggests that the immobilized protein is initially more compact but extends to the

### TABLE 1

| Construct | Labeling sites | Separation (amino acids) |
|-----------|----------------|-------------------------|
| N2B-1     | s1273c, s1288c | 15                      |
| N2B-2     | C1394, s1477c  | 83                      |
| N2B-3     | s1353c, C1453  | 100                     |
| N2B-4     | s1373c, C1394  | 121                     |
| N2B-5     | s1353c, s1477c | 124                     |
| N2B-6     | s1323c, C1453  | 130                     |
| N2B-7     | s1273c, s1413c | 140                     |
| N2B-8     | s1273c, C1445  | 172                     |

### TABLE 2

| Construct | Unphosphorylated | Phosphorylated | p value |
|-----------|-----------------|---------------|---------|
| N2B-1     | 0.90 ± 0.075    | 0.88 ± 0.003  | NA*     |
| N2B-2     | 0.64 ± 0.016    | 0.59 ± 0.042  | 0.299   |
| N2B-3     | 0.55 ± 0.034    | 0.48 ± 0.013  | 0.084   |
| N2B-4     | 0.45 ± 0.008    | 0.36 ± 0.016  | 0.019   |
| N2B-5     | 0.49 ± 0.061    | 0.43 ± 0.013  | 0.286   |
| N2B-6     | 0.47 ± 0.051    | 0.43 ± 0.008  | 0.319   |
| N2B-7     | 0.43 ± 0.032    | 0.40 ± 0.038  | 0.497   |
| N2B-8     | 0.38 ± 0.027    | 0.30 ± 0.015  | 0.065   |

* NA means not applicable.
Phosphorylation of GluN2B

same extent after phosphorylation. The linear relationship between FRET efficiency and fluorophore separation in the primary sequence was preserved in immobilized samples indicating a lack of local structure (Fig. 6B).

DISCUSSION

The intracellular domain of GluN2B is so essential to receptor function that genetic deletion of the C terminus is lethal (5). Although the CTD is intrinsically disordered, it exerts allosteric effects on receptor gating and trafficking that are modulated through Src phosphorylation, which alters both channel currents and receptor trafficking (14). To understand how tyrosine phosphorylation regulates NMDAR, we characterized the effect of Src phosphorylation on the conformation and dynamics of the C-terminal domain from the NMDA receptor subunit GluN2B.

This study used a fragment (CTD2) that contains the two major Src phosphorylation sites (Fig. 1). All ensemble methods confirm that CTD2 is intrinsically disordered under physiological buffer conditions. This agrees with predictions based on amino acid sequence (Fig. 1B). The protein is rapidly degraded during proteolysis experiments (Fig. 2). The appearance of higher molecular weight intermediates during digestion suggests the existence of a protein “core” that contains aromatic residues. This agrees with the SEC data, which show that CTD2 is more compact than a random coil. CTD2 is a compact disordered globule with no stable secondary structure (Fig. 3).

To probe the conformation of CTD2, we used single molecule FRET. This necessitates the use of covalently attached fluorophores that are charged and aromatic. It is inconceivable that the dyes do not participate in the conformational equilibrium, an unavoidable problem associated with using exogenous probes to study biomolecules. Fluorescent labeling had minimal effects on the radius of hydration particularly when compared with the much smaller phosphate moiety. This speaks to the powerful specific activity of this post-translational modification.

We placed FRET pairs throughout the protein to probe for local structure. If local structure existed, we would have expected a dispersion of FRET values (37). Instead, the results

| TABLE 3 |
| Effect of Src phosphorylation on the percent of molecules showing stochastic FRET transitions |

| Construct | Unphosphorylated | Phosphorylated |
|-----------|------------------|----------------|
| N2B-1     | 3 ± 4            | 1 ± 1          |
| N2B-2     | 52 ± 18          | 62 ± 10        |
| N2B-3     | 75 ± 6           | 80 ± 3         |
| N2B-4     | 87 ± 1           | 85 ± 0         |
| N2B-5     | 81 ± 3           | 93 ± 4         |
| N2B-6     | 81 ± 5           | 88 ± 6         |
| N2B-7     | 81 ± 7           | 91 ± 1         |
| N2B-8     | 84 ± 1           | 83 ± 9         |

| TABLE 4 |
| Effect of Src phosphorylation on median smFRET efficiency in surface immobilized CTD2 |

| Construct | Unphosphorylated | Phosphorylated | p value |
|-----------|------------------|----------------|---------|
| N2B-1     | 0.92 ± 0.011     | 0.90 ± 0.013   | NA²     |
| N2B-2     | 0.65 ± 0.006     | 0.55 ± 0.005   | 0.003   |
| N2B-3     | 0.64 ± 0.015     | 0.45 ± 0.023   | 0.010   |
| N2B-4     | 0.50 ± 0.010     | 0.37 ± 0.038   | 0.045   |
| N2B-5     | 0.57 ± 0.036     | 0.43 ± 0.005   | 0.033   |
| N2B-6     | 0.57 ± 0.045     | 0.42 ± 0.013   | 0.044   |
| N2B-7     | 0.45 ± 0.030     | 0.35 ± 0.0002  | 0.041   |
| N2B-8     | 0.40 ± 0.029     | 0.27 ± 0.030   | 0.047   |

² NA means not applicable.

FIGURE 6. Effect of surface tethering on phosphorylation-induced conformational changes in the GluN2B CTD2. For these experiments, the palmitoylation sites in the N terminus of CTD2 were replaced with a biotinylation sequence, which allows for directional tethering of CTD2 to a passivated microscope slide. All labeling sites for FRET were identical to the previous experiments. A, single molecule FRET was measured for each FRET construct before and after Src phosphorylation. Constructs are shown in order of the fluorophore separation as indicated in each panel. The histogram contains FRET values calculated for each 100-ms frame of observation. Unphosphorylated proteins are shown as dashed lines. Phosphorylated proteins are shown as solid lines. B, median FRET for each distribution above plotted against the fluorophore separation in residues. Data were fit to a line using nonlinear least squares fitting. Unphosphorylated proteins are shown as open circles fit to a dashed line. Src-phosphorylated proteins are shown as filled squares fit to a solid line. Error bars are from replicate measurement of median FRET in separate experiments.
showed minimal positional dependence and instead the FRET scaled linearly with the fluorophore separation. Such statistical scaling is a general property of random coils and suggests a lack of discernible, stable conformations or local structure in the native state ensemble. The protein is truly “molten” sampling a random distribution of states albeit with some component on a slow time scale.

The smFRET distributions were irregularly shaped with some positional variability that was reproducible. The shape of the point wise smFRET distribution depends on both the FRET states visited and the duration of each FRET state. Thus, changes in conformation or the lifetime of individual states would change the shape. It is difficult to interpret changes in the distribution shape, so we only extracted the median FRET because the distributions had a clear predominant peak.

Both the ensemble and single molecule data suggest that phosphorylation results in a general expansion of the polypeptide chain. Median FRET shifted for all constructs regardless of their positioning relative to the phosphorylation sites. Regardless of the phosphorylation state, the FRET scales linearly with amino acid separation. This along with the CD spectra suggest a featureless energy landscape without favored conformations or a stable local structure.

Because GluN2B is palmitoylated in neurons, we examined the effect of surface tethering on the conformation of CTD2. Disordered domains are frequently linked to transmembrane proteins, with unknown effects on their conformation. The surface should reduce the number of accessible conformations through steric repulsion. Such effects are difficult, if not impossible, to measure in solution but can be observed in immobilized single molecules.

The conformation of CTD2 was affected by immobilization with a trend toward higher FRET values (Tables 2 and 4). This suggests that CTD2 becomes more compact when the N terminus is immobilized. In contrast, the FRET values after phosphorylation were unaffected by immobilization so the expanded state appears the same. The net result is a greater net shift upon phosphorylation. Thus, in these experiments, surface tethering amplified the signal induced by phosphorylation by magnifying the difference between states.

Phosphorylation does not induce structure in CTD2, so the effects on NMDA receptor activity must arise from the general expansion. If anything, CTD2 becomes more like a random coil. The volume change could affect steric interactions with the membrane and change the orientation of this domain relative to the ion channel or exert tension on the final transmembrane domain. The proteolysis experiments suggest that the expansion would result in greater surface exposure of potential sites of post-translational modification and protein binding. Such a global rearrangement could change the affinity for protein interactions either with the other NMDA receptor subunits or synaptic proteins like CAMKII or PSD-95.

CTD2 may interact with other portions of GluN2B (e.g. CTD1 or intracellular loops) or CTD from the other subunits within a tetrameric receptor. Such interactions could easily change the structure of this “domain” away from what we observe in these experiments. CTD2 in isolation represents a “ground state” at equilibrium under physiological pH and ionic strengths. The effect of such interactions remains to be investigated.

Intrinsically disordered intracellular domains are a common feature of cell surface receptors and channels. Many are similarly regulated by phosphorylation. How such disordered CTD might modulate the activity of their associated proteins is a complete unknown. Our findings here suggest that in GluN2B disorder actually increases, which only deepens the mystery of how the allosteric affects are mediated.

Acknowledgments—We thank Ben Sondergord and Markus Seeliger for providing purified Src kinase, Daniel P. Raleigh for assistance with circular dichroism, and Suzanne Scarlata for assistance with fluorometry. We benefited from conversations with Nicolas Nassar, Todd Miller, and Rasheek Kazi.

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