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Effects of pseudophosphorylation mutants on the structural dynamics of smooth muscle myosin regulatory light chain

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Abstract

We have performed 50 independent molecular dynamics (MD) simulations to determine the effect of pseudophosphorylation mutants on the structural dynamics of smooth muscle myosin (SMM) regulatory light chain (RLC). We previously showed that the N-terminal phosphorylation domain of RLC simultaneously populates two structural states in equilibrium, closed and open, and that phosphorylation at S19 induces a modest shift toward the open state, which is sufficient to activate smooth muscle. However, it remains unknown why pseudophosphorylation mutants poorly mimic phosphorylation-induced activation of SMM. We performed MD simulations of unphosphorylated, phosphorylated, and three pseudophosphorylated RLC mutants: S19E, T18D/S19D and T18E/S19E. We found that the S19E mutation does not shift the equilibrium toward the open state, indicating that simple charge replacement at position S19 does not mimic the activating effect of phosphorylation, providing a structural explanation for previously published functional data. In contrast, mutants T18D/S19D and T18E/S19E shift the equilibrium toward the open structure and partially activate *in vitro* motility, further supporting the model that an increase in the mol fraction of the open state is coupled to SMM motility. Structural analyses of the doubly-charged pseudophosphorylation mutants suggest that alterations in an interdomain salt bridge between residues R4 and D100 results in impaired signal transmission from RLC to the catalytic domain of SMM, which explains the low ATPase activity of these mutants. Our results demonstrate that phosphorylation produces a unique structural balance in the RLC. These observations have important implications for our understanding of the structural aspects of activation and force potentiation in smooth and striated muscle.

Keywords: Smooth muscle myosin; striated muscle; regulatory light chain; phosphorylation; pseudophosphorylation; molecular dynamics simulations.

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Introduction

Smooth muscle myosin (SMM) is a member of the myosin superfamily of motor proteins, which use chemical energy from ATP hydrolysis to perform mechanical work on actin. Activation of smooth muscle requires phosphorylation of the regulatory light chain (RLC), which is distant from the myosin active site on the catalytic domain (Fig. 1A). Unphosphorylated

SMM is auto-inhibited by interactions between the two catalytic domains ^{1, 2} that are relieved by RLC phosphorylation ^{3, 4}, through a mechanism that is proposed to involve interactions among the heavy chain, the essential light chain (ELC), and the RLC ^{5, 6}.

Our previous EPR experiments⁷ and Fig. 1. (A) Crystal structure of myosin S1, with heavy chain molecular dynamics (MD) simulations ^{8, 9} showed that upon phosphorylation of S19, of RLC undergoes a disorder-to-order



(HC, blue), essential light chain (ELC, green), and regulatory light chain (RLC, red). (B) Structural model of phosphorylation-induced structural changes within the RLC in solution, based on FRET experiments and MD simulations (Ref. 10). Left (unphosphorylated RLC): the closed (top) and open (bottom) states are in equilibrium, with the more helical open state of PD (purple) being more the 25-residue phosphorylation domain (PD) populated. Right: Upon phosphorylation, the structure shifts toward the open state, activating smooth muscle.

transition accompanied by an increase in helical structure. Complementary FRET and MD simulations showed that the biochemical and structural states of the RLC are loosely coupled: the PD of the RLC simultaneously populates two structural states, closed and open, in both unphosphorylated and phosphorylated biochemical states¹⁰. This complementary approach revealed that unphosphorylated RLC preferably populates a closed conformation, and that phosphorylation shifts the conformational equilibrium toward the open state, increasing its mol fraction by 23% ¹⁰. Based on these observations, we proposed a model in which a modest change

in the conformational equilibrium of RLC is sufficient and necessary to activate smooth muscle (**Fig. 1B**).

These complementary studies suggest that incorporation of negative charge into the phosphorylation sites of the RLC should be sufficient increase the mol fraction of the open structural state, resulting in SMM activation. However, experiments have shown that simple charge replacement or deletion of PD causes partial or no activation ^{11, 12}, suggesting that full activation requires specific structural changes within PD. In particular, functional assays have shown that pseudophosphorylation mutants have minimal or no effect on SMM activation ¹². For example, mutant S19E was studied to determine the effect of simple charge replacement in the PD; functional assays performed on this mutant showed that S19E does not activate ATPase SMM^{12} . motility of Biochemical experiments activity or also showed that pseudophosphorylation mutants T18D/S19D and T18E/S19E poorly mimic phosphorylationinduced ATPase activation, although they retain in vitro motility¹². Only in the case of T18E/S19E was there both a slight activation of ATPase activity and a moderate activation of motility ¹².

These observations suggest that a structural shift of the RLC toward the open state is required for smooth muscle activation, and that pseudo-phosphorylation mutants do not induce this structural shift. To test this hypothesis, we have performed 50 independent 0.2- μ s MD simulations to determine the effect of three pseudophosphorylation mutants (S19E, T18D/S19D and T18E/S19E) on the structural dynamics of the RLC.

Methods

Preparation of the systems

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The structure of the RLC bound to the IQ motif of myosin heavy chain was obtained by homology modeling using the procedure described previously¹⁰. Unphosphorylated, phosphorylated (at position S19) and pseudophosphorylation mutants S19E, T18D/S19D and T18E/S19E were modeled using the PSFGEN utility of NAMD 2.9¹³. Phosphate was assigned a charge of -2, based on $pK_a = 6.5$ for phosphoserine and pH > 7 in smooth muscle¹⁴, and previous time-resolved FRET experiments¹⁰. We adjusted the side-chain ionization states of the RLC to pH = 7.0 with PROPKA¹⁵. The system was embedded in TIP3P water boxes with a minimum distance of 2 nm between the protein and the edges of the periodic box. Na⁺ and Cl⁻ ions were added to neutralize charge and produce an ion concentration of approximately 150 mM, as in experiments. The final systems contained ~100,000 atoms. Topologies and parameters were used according to the CHARMM22 force field with CMAP correction ^{16, 17}.

Molecular dynamics protocol

MD simulations were performed by using the program NAMD 2.9 ¹³. Periodic boundary conditions ¹⁸, particle mesh Ewald ^{19, 20}, a nonbonded cutoff of 9 Å and a 2 fs time step were used. The NPT ensemble was maintained with a Langevin thermostat (310K) and a Langevin piston barostat (1 atm). The system was first subjected to energy minimization for 1000 steps, followed by a warming up period for 200 ps. This procedure was followed by equilibration for 0.01 μ s with backbone atoms harmonically restrained using a force constant of 100 kcal mol⁻¹ nm⁻². To improve conformational sampling, we performed ten independent MD simulations for each system, with randomly selected initial atomic velocities ²¹. Ten independent simulations of unphosphorylated, phosphorylated and pseudophosphorylation mutants were continued for 0.2 μ s, for a total of 50 MD simulations and a cumulative simulation time of 10 μ s.

Analysis and visualization

VMD ²² was used for analysis, visualization, and rendering of the structures. STRIDE ²³ was used to analyze the evolution of the secondary structure of the PD in unphosphorylated, phosphorylated and four pseudophosphorylation mutants. STRIDE recognizes secondary structural elements in proteins from their atomic coordinates. It utilizes both hydrogen-bond energy and main chain dihedral angles to define the secondary structure pattern, relying on database-derived recognition parameters with the crystallographers' secondary structure definitions as a standard-of-truth.

We calculated C_{α} - C_{α} distances between residues S2, K3 and A7 of PD and M129 of the Cterminal lobe of the RLC core (**Fig. 1B**). The distances for each set of ten independent MD simulations were then combined into a single histogram. Finally, each histogram was fitted to one-Gaussian and two-Gaussian distance distributions $\rho(R)$. We found that the best fits, indicated by a lower χ^2 (**Table S1, Supplementary material**), were consistently obtained for the two-Gaussian model:

$$\rho(R) = \sum_{j=1}^{2} X_j \sigma_j^{-1} (2\pi)^{-1/2} \exp(-[(R-R_j)/(2\sigma_j)]^2 \sigma_j = FWHM_j/[2^*(2\ln 2)^{1/2}].$$

Each fit yielded six independent parameters of $\rho(R)$: centers for open (*j*=1) and closed (*j*=2) structural states, R_1 and R_2 , widths *FWHM*₁ and *FWHM*₂, and mol fractions X_1 and X_2 .

Results and discussion

We previously showed that phosphorylation of PD at S19 does not induce noticeable global structural changes on the RLC; instead, phosphorylation induces structural changes only around

residue K11 of the PD ¹⁰. To determine whether pseudophosphorylation mutants alter the structural stability of the RLC, PD, or both, we calculated the C_{α} root-mean square fluctuations (RMSF) averaged over all individual MD simulations (**Fig. 2**). Analysis of the RMSF plots revealed negligible differences between unphosphosphorylated, phosphorylated and pseudophosphorylated RLC. For example, variable residue mobility is observed in the N- and C-termini of the RLC. Despite these small differences, the RMSF patterns are similar for phosphorylated and pseudophosphorylated RLC, indicating that phosphorylation and pseudophosphorylation mutants do not alter the global structural stability of the RLC. Furthermore, RMSF plots indicate that the 24-residue PD is very flexible in phosphorylated and unphosphorylated RLC (i.e., maximum RMSF values of 2.4 nm), in agreement with a previous study ¹⁰. This suggests that the effects of pseudophosphorylation mutants on the structural dynamics of the RLC are localized to the

PD.

Because RMSF plots do not directly indicate secondary structure, we calculated the percentage of helical content for each individual residue in PD (**Fig. 3**). In agreement with previous MD simulations, we found that helical periodicity of unphosphorylated PD is disrupted at residue K11, and phosphorylation



s disrupted at **Fig. 2.** Root-mean-square fluctuations (RMSF) of the RLC. RMSF values were calculated about the average position of C_{α} atoms. Each plot represents the average RMSF of ten independent MD trajectories.

increases the helical content of this residue by $\sim 60\%$ ⁸⁻¹⁰. Analysis of mutant S19E revealed a secondary structure pattern was similar to that of unphosphorylated RLC, indicating that this





Fig. 3. Percentage of time that each residue of the PD spends an in α -helical conformation. (A) unphosphorylated, (B) phosphorylated, (C) T18D/S19D,(D) T18E/S19E, and (E) S19E.

T18D/S19D and T18E/S19E restore the helical periodicity of PD at position K11 in a similar manner as phosphorylation. These observations indicate that, except for S19E, pseudophosphorylation mimics the local disorder-to-order transition of PD. These observations also indicate that, if present, the structural effect of pseudophosphorylation is localized around residue K11, which suggests that the inability of pseudophosphorylation to optimally activate SMM does not result from global structural alterations in the RLC.

We calculated the distributions of distances from residues in PD (S2, K3 and A7) to M129 in the C-terminal lobe of the RLC (**Fig. 4**), because these are the sites previously labeled in FRET experiments ¹⁰. As shown in Table S1 (Supporting Material), a two-Gaussian distribution best fits the data in all cases, indicating two clearly resolved structural states: with a short distance assigned to the closed state and a longer distance assigned to the open state. **Table 1** shows the mol fraction in the open state, compared with previous experimental results. Phosphorylation shifts the distribution by 23% toward the open state, in excellent agreement with previous timeresolved FRET experiments¹⁰. These findings indicate that although MD simulations were not performed on the entire regulatory domain and were performed in the submicrosecond time scale, the MD simulations capture the structural dynamics of RLC observed in experiments ¹⁰. These observations are also consistent with a previous study showing that 5–10 independent MD



Fig. 4. Comparison of the distance distributions of unphosphorylated (blue) and phosphorylated (red, pS19) RLC with those of pseudophosphorylation mutants. Each distance distribution, determined for each Cα-Cα distance between S2, K3, A7 and M129, is the best 2-Gaussian fit to the average of histograms from 10 trajectories.

simulations are sufficient to capture the structural properties of a protein observed in experiments ²¹. In contrast, pseudophosphorylation gave results substantially different from that of phosphorylation: S19E showed no shift at all toward the open state, while the other mutants showed shifts much greater than that of phosphorylation (**Fig. 4, Table 1**). Thus simple charge replacement does not explain the structural effects of phosphorylation, and this is consistent with the observation that the functional effects are also not equivalent (**Table 1**).

Functional experiments for S19E. have shown that that, except mutant pseudophosphorylation partially restores SMM motility *in vitro*¹². Our simulations showed that only mutant S19E is unable to induce a shift toward the open state (Table 1); this evidence suggests a relationship between an increase in the mol fraction and motility. Indeed, we found a relationship between the mol fraction of the open state of the RLC and in vitro motility of SMM. This linear relationship indicates that an increase in mol fraction of the open state and motility are coupled. These observations suggest that increasing the mol fraction of the open state is sufficient to partially activate motility of SMM.

RLC species	Mol fraction in open state ¹	Increase of mol fraction in open state ²	Relative motility ³	Relative actin- activated ATPase activity ³
Unphosphorylated	0.25 (0.33) ¹	-	0.00 ± .02	0.02 ± .01
Phosphorylated	0.48 (0.56) ¹	0.23 (0.23) ¹	1.00	1.00
S19E	0.25	0	0.00 ± .02	0.00 ± .01
T18E/S19E	0.88	0.63	0.45 ± .03	0.16 ± .01
T18D/S19D	0.79	0.54	0.59 ± .02	0.0 ± .01

Table 1. Summary of	structural and	functional	data on RLC
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¹Values in parentheses are from TR-FRET experiments ¹⁰.

²Increase in mol fraction relative to unphosphorylated RLC.

³Experimental functional data ¹².

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We found that the inability of mutant S19E to shift the structure of the RLC toward the open state correlate with *in vitro* assays showing that this mutant does not activate ATPase activity ¹². Mutants T18D/S19D and T18E/S19E substantially increase the mol fraction of the open structural state, sometimes even more effectively than phosphorylation. However, functional assays showed that these mutants poorly mimic phosphorylation-induced actin-activated ATPase activity of SMM. If T18D/S19D and T18E/S19E can effectively shift the equilibrium toward the open state and increase motility, why are they unable to optimally regulate ATPase activation? We have previously shown that an interdomain salt bridge between R4 (PD) and D100 (C-lobe) plays a role in the stabilization the closed state in both unphosphorylated and phosphorylated RLC¹⁰. The presence of this PD-to-C-lobe salt bridge in both phosphorylated and unphosphorylated RLC indicates that the R4-D100 salt bridge serves as an anchor to properly position PD onto the C-terminal lobe in the closed state. To evaluate the presence of this salt bridge, we calculated the distance distribution between atoms N_{ζ} of R4 and C_{γ} of D100 (**Fig. 5**). We found that unphosphorylated RLC features three peaks, which correspond to the closed state stabilized by salt bridge R4-D100 (closed_(SB), R_{mean} =0.4 nm), a closed state not stabilized by a salt bridge (closed_(noSB), R_{mean}=1.1 nm) and an open state (Open, R_{mean}=3.3 nm) (Fig. 5, right). In phosphorylated RLC, the peak representing $closed_{(SB)}$ is still present, but the $closed_{(noSB)}$ state is no longer present. The distance distribution of S19E is very similar to that of unphosphorylated RLC, explaining the absence of ATPase activity ¹². Among the mutants that increase the fraction of the open state, only T18E/S19E retains features of both unphosphorylated and phosphorylated RLC: two small peaks corresponding to closed_(SB) and closed_(noSB), and broad open states, characteristic of dynamic tertiary structure (Fig. 5). This pattern is completely disrupted in mutant T18D/S19D.



Fig. 5. Distance distribution between atoms N_{ζ} **of R4 and** C_{γ} **of D100.** Distance distributions are the average of histograms from the 10 trajectories combined for the five RLC species. Distance distributions were constructed by fitting distance histograms to a 3-Gaussian model. Right: Typical structural details. Closed_(SB) is the closed state stabilized by a salt bridge between R4 and D100. Closed_(no SB) is the closed state that is not stabilized by this salt bridge. Open is the open state, which also lacks the salt bridge. PD, C-terminal lobe and HC are colored in purple, red and blue, respectively. The shaded areas in the distribution plots indicate the presence of the closed_(SB) (cyan), closed_(noSB) (grey) and open (white) states.

These results suggest that upon phosphorylation, only the closed state stabilized by the salt bridge R4-D100 and the open states are present in equilibrium. The presence of a $closed_{(noSB)}$ population in unphosphorylated RLC, and its absence in phosphorylated RLC, indicates that the $closed_{(noSB)}$ conformation is probably responsible for inhibition of ATPase activity. We found that the state $closed_{(SB)}$ is present in both phosphorylated and unphosphorylated RLC; the proximity of the C-lobe of the RLC to the IQ and ELC (**Fig. 1A**) suggests that the salt bridge R4-D100 is necessary for the correct placement of the highly mobile PD onto the C-terminal lobe of the RLC during the open-to-closed transition. Because salt bridge R4-D100 is still partially present in phosphorylated RLC, it is likely that the alignment of PD and C-terminal lobe facilitated by this salt bridge is required for signal transmission from the RLC toward the catalytic domain via the ELC. It is possible that these differences are linked to the unique physicochemical properties of phosphoserine. For instance, it has been shown that salt bridge interactions are stronger for phosphoserine than for acidic residues^{24, 25}. In addition, phosphoserine has a significantly higher probability than glutamate to form hydrogen bonds with arginein and lysine ²⁵. Stronger electrostatic interactions in the presence of phosphoserine might also impose additional structural restraints necessary for the correct placement of the PD domain onto the C-lobe (i.e., via pS19-R16 salt bridges ⁹). We propose that the absence of the salt bridge R4-D100 in mutant T18D/T19D results in an impaired signal transmission from the RLC to the catalytic domain of SMM through the ELC ⁵, which explains the low ATPase activity of these mutants ¹². Conversely, partial activation of ATPase activity of T18E/S19E probably results from a shift toward the open state, the presence of a small fraction of the closed_(*SB*) state, and the presence of inhibitory closed_(*noSB*) state (**Fig. 5**).

In conclusion, we found that simple charge replacement at position S19 is not sufficient to induce the structural changes necessary for activation of both ATPase activity and motility of SMM. We also found that pseudophosphorylation mutants that induce a structural shift toward the open state can also partially or fully mimic the motility observed with phosphorylated RLC. This finding suggests that an increase in the mol fraction of the open state and motility in SMM are coupled. Structural analyses of pseudophosphorylation mutants T18D/S19D and T18E/S19E suggest that alterations of an interdomain salt bridge between residues R4 and D100 result in impaired signal transmission from the RLC to the catalytic domain of SMM, which explains the low of ATPase activity of these mutants. Our results demonstrate that phosphorylation produces

a unique conformational balance in the RLC, which plays a key role in smooth muscle activation. Our findings regarding the structural landscape of the RLC that is vital for activation reach beyond smooth muscle to different muscle types and species, including the residues we have investigated, where the RLC sequence remains quite conserved. RLCs are very powerful governors of motile effects, underscored by the striking capability of smooth muscle RLC to regulate and modulate force generation characteristic of smooth muscle, even when it is bound to the IQ motif of striated myosin ¹². Taken together, these findings are not only relevant to smooth muscle, but also to other muscle types. For instance, mono- and biphosphorylation of RLC in tarantula thick filaments from striated muscle are important for activation (i.e., destabilization of the super-relaxed state ²⁶) and force potentiation, respectively . These observations have important implications for our understanding of the structural aspects of activation and force potentiation in smooth and striated muscle.

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