

The Mechanism of the Reverse Recovery Step, Phosphate Release, and Actin Activation of *Dictyostelium* Myosin II^{*[5]}

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The rate-limiting step of the myosin basal ATPase (*i.e.* in absence of actin) is assumed to be a post-hydrolysis swinging of the lever arm (reverse recovery step), that limits the subsequent rapid product release steps. However, direct experimental evidence for this assignment is lacking. To investigate the binding and the release of ADP and phosphate independently from the lever arm motion, two single tryptophan-containing motor domains of *Dictyostelium* myosin II were used. The single tryptophans of the W129+ and W501+ constructs are located at the entrance of the nucleotide binding pocket and near the lever arm, respectively. Kinetic experiments show that the rate-limiting step in the basal ATPase cycle is indeed the reverse recovery step, which is a slow equilibrium step ($k_{\text{forward}} = 0.05 \text{ s}^{-1}$, $k_{\text{reverse}} = 0.15 \text{ s}^{-1}$) that precedes the phosphate release step. Actin directly activates the reverse recovery step, which becomes practically irreversible in the actin-bound form, triggering the power stroke. Even at low actin concentrations the power stroke occurs in the actin-attached states despite the low actin affinity of myosin in the pre-power stroke conformation.

The mechanism of the phosphate (P_i) release of the myosin ATPase is of particular interest, because it is the key step that is presumed to be activated by actin and is coupled to the working stroke of the cross-bridge cycle (1, 2). The observed rate of phosphate release in the absence of actin is of the order of 0.05 s^{-1} for rabbit skeletal muscle myosin II at 20°C and provides the major contribution to the rate of the overall basal ATPase activity. Phosphate binding is, however, weak ($K_d > 1 \text{ mM}$), which led Trentham *et al.* (3, 4) to propose that phosphate release was controlled by a slow isomerization followed by a rapid phosphate release step (*steps 4 and 5* in Scheme 1, respec-

tively). This suggestion was made on the grounds that the second-order rate constant for phosphate binding would be abnormally low if steps 4 and 5 were combined into a single step.

This idea was supported by the experiments of Mannherz *et al.* (5) who demonstrated that a small amount of labeled $M^*\text{ATP}^3$ could be synthesized by the addition of millimolar concentrations of ^{32}P -labeled phosphate to myosin subfragment 1 (S1) in the presence of saturating ADP. From the kinetics and extent of incorporation, they estimated that the $M^{**}\text{ADP}\cdot P_i$ to $M^*\text{ADP}\cdot P_i$ isomerization was slow but freely reversible ($K_4 = 15.6$ at pH 8 and 0.9 at pH 6) while phosphate binding to $M^*\text{ADP}$ was weak ($K_5 = 7.3 \text{ mM}$ at pH 8 and 55 mM at pH 6). Subsequently, these authors modified their view (6) and considered that the apparent saturation they observed in ^{32}P incorporation was a consequence of the increase in ionic strength with increasing phosphate concentrations. Their important conclusion about the very high affinity of ATP for myosin in the $M^*\text{ATP}$ complex was unaffected by this problem, but the values of K_4 and K_5 could no longer be reliably disentangled. Thus, direct evidence for two steps remains elusive.

It was demonstrated in both solution and fiber experiments (3, 7–9) that phosphate release precedes the release of ADP. The ADP-bound myosin state possesses strong actin binding properties, whereas both ADP and phosphate binding to the active site of myosin results in the weakening of the actin-myosin interaction. It was shown in solution experiments that more than half of the free energy obtained from ATP hydrolysis is liberated during phosphate release (8), therefore this step was suggested to be coupled to the power-generating lever arm movement. However, Goldman & Brenner pointed out that solution kinetics might differ from fiber kinetics (10). They and others demonstrated that in single fiber experiments the power stroke precedes the fast phosphate release step and that the starting state of the power stroke in this case is a weak actin binding $A\cdot M\cdot\text{ADP}\cdot P_i$ state (11–14).

It is well known that addition of phosphate analogs, such as BeF_x and AlF_4^- , to skeletal muscle myosin-ADP complexes, generates an enhancement in tryptophan fluorescence, indicative

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³ The abbreviations used are: M^* , enhanced fluorescent state of myosin S1 (~5%); M^{**} , enhanced fluorescent state of myosin S1 (~15%); S1, subfragment 1; dmATP, 3'-(*N*-methylanthraniloyl)-2'-deoxy-ATP; dmADP, 3'-(*N*-methylanthraniloyl)-2'-deoxy-ADP; *Dd*, *D. discoideum*; mant, *N*-methylanthraniloyl.

TABLE 1

Nomenclature of the enzymatic steps

It is essential to distinguish the post-recovery step conformational state ($M^* \cdot \text{ATP}$ and $M^* \cdot \text{ADP} \cdot \text{P}_i$ states) from the pre-power-stroke state ($A \cdot M^* \cdot \text{ADP} \cdot \text{P}_i$). The latter is in the actin-bound conformational state, which is different from the post-recovery step state from both energetic and structural aspects.

Enzymatic steps		Rate constant		Initial and end states of the forward steps (lever arm position/nucleotide form)	
K_1	ATP binding collision step	k_1	Forward	Apo	ATP-binding
K_2	ATP binding-induced conformational change	k_{-1}	Reverse	Collision complex with ATP	
		k_2	Forward	Collision complex with ATP	
K_{3a}	Recovery step ^b	k_{-2}	Reverse	Down ^c ATP state	Effective hydrolysis step
		k_{3a}	Forward	Down ATP state	
K_{3b}	Hydrolysis step	k_{-3a}	Reverse	Up ^c ATP state	
		k_{3b}	Forward	Up ATP state	
K_4	Reverse recovery step ^{d,e}	k_{-3b}	Reverse	Up ADP·P _i state	
		k_4	Forward	Up ADP·P _i state	
K_4'	Power stroke	k_{-4}	Reverse	Down ADP·P _i state	
		k_4'	Forward	Actin-bound up ADP·P _i state	
		k_{-4}'	Reverse	Actin-bound down state ^e	

^a The down lever arm state is also called the post-rigor and post-power stroke states, although it also exists in the actin-detached state.

^b We recommend the nomenclature of the recovery/reverse recovery step instead of the recovery/reverse recovery stroke, because a stroke cannot be explained by an equilibrium step.

^c The up lever arm state is generally called pre-power stroke state, although it also exists in actin-detached state.

^d Reverse recovery step is also called the power stroke, even in the absence of actin. We suggest the distinction between the actin-induced power stroke from the reverse recovery step, which occurs in the absence of actin or in the actin-detached form of myosin.

^e Because the sub-steps of the power stroke have not been resolved in detail, the final state is undefined, as yet.

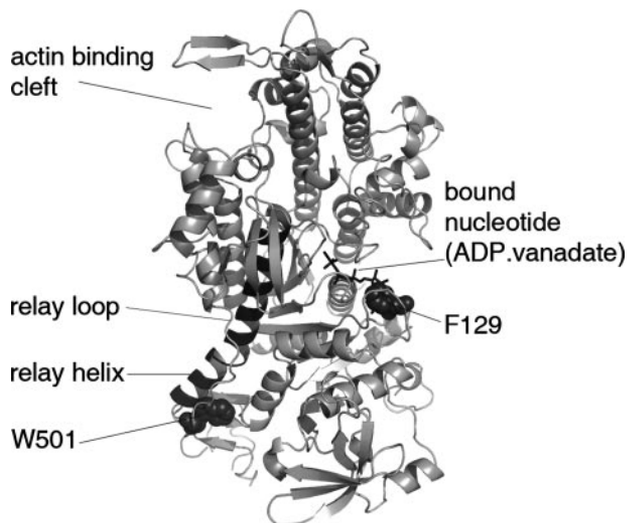


FIGURE 1. X-ray crystal structure of *Dd* myosin II MD complexed with ADP-vanadate (1VOM). Trp-501 and Phe-129 residues are rendered in space-filling appearance. W501+ and W129+ motor domain constructs contain single tryptophans at the positions of 501 and 129, respectively. Phe-129 is in close proximity to bound nucleotide (here Mg·ADP·VO₄ is shown as a black stick), whereas Trp-501 is located in the relay region.

obtained from Roche Applied Science and Jena Bioscience GmbH, respectively.

Phosphate Buffers—Care was taken to maintain the constant ionic strength in the experiments using different phosphate concentrations or pH. Ionic strength values were calculated according to (27) using the Buffers for pH Control website, and ionic strength was kept constant with additional NaCl to the phosphate buffers.

Reduction of Pyrophosphate Contamination—To reduce the pyrophosphate contamination in the phosphate buffers, 0.5 M Na₂HPO₄ buffer (pH ≤ 12.6) was autoclaved at 121 °C for 30 min immediately before the experiments.

Expression and Purification of Proteins—In this study we used *Dd* myosin II motor domain mutants, which are based on the M761 construct (28). The W501+ construct possesses W36F, W432F, and W584F mutations (18), the

W129+ construct contains W36F, W432F, W501F, W584F, and F129W mutations (17), and both were cloned into the pDXA-3H vector (29). Both W501+ and W129+ were expressed in *Dd* cells cultured in HL-5 (Formedium) medium at 21 °C and purified as in a previous study (18). After purification proteins were dialyzed against an assay buffer (20 mM HEPES, pH 7.2, 40 mM NaCl, 1 mM MgCl₂, 3 mM β-mercaptoethanol, 5 mM benzamidine), and after dialysis, proteins were frozen to liquid nitrogen in 50-μl droplets. Actin was purified as described (30). Protein concentrations were determined with the Bradford method.

Steady-state Kinetic Measurements—Steady-state spectra and time courses of the purified proteins were measured on a FluoroMax SPEX-320 spectrofluorometer. All measurements were carried out at 20 °C. Trp fluorescence was excited at 296 nm with 2 nm bandwidth for both excitation and emission sides, and emission was detected from 305 to 420 nm. Time courses were followed by tryptophan fluorescence excited at 296 nm and detected at 340 nm.

Transient Kinetic Measurements—Stopped-flow measurements were carried out with a KinTek-2004 equipped with a Super Quiet Mercury-Xenon Lamp (Hamamatsu) or with a BioLogic SFM-300/400 stopped-flow fluorometer equipped with a mercury-xenon lamp (Hamamatsu). Trp fluorescence was excited at 297 nm where the mercury-xenon lamp has a discrete emission peak. Slits were adjusted to 2 nm in the case of the KinTek-2004 stopped-flow and a 340 nm interference filter (Corion CFS-001999 9L134) was used on the emission side in both instruments. *N*-Methylantraniloyl (mant) derivatives were excited either directly at 365 nm or via energy transfer (Forster resonance energy transfer) at 280 nm. In both cases, a WG420 cut-off filter (Comar Instruments) was used to selectively measure mant fluorescence. The cuvette block and the syringe house were thermostatted at 20 °C both in the KinTek 2004 and the BioLogic SFM-300/400 stopped-flow solutions. The dead-time parameter of the KinTek 2004 apparatus at 18 ml/s flow-rate was determined to be 1 ms (31). The BioLogic SFM-300/400 was equipped with an 8-μl cuvette (μFC-08) or a

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15- μl cuvette (FC-15), which had <0.2 ms or 2 ms dead time at 18 ml/s, respectively, which increases if slower flow-rate is applied. Unless otherwise noted post-mix concentrations of the reactants are presented.

Actin-activated ATPase Measurement—Actin-activated ATPase activity of W501+ was measured using a pyruvate kinase/lactate dehydrogenase-coupled assay at 20 °C as follows: 2% pyruvate kinase/lactate dehydrogenase enzyme mixture, 1 mM ATP, 1 mM phosphoenolpyruvate, and 200 μM NADH was used in the 5 mM HEPES, 1 mM MgCl_2 , and 1 mM KCl buffer at pH 7.2, with 0.5 μM W501+ and an actin concentration of 0 to 94 μM .

NMR Spectroscopy— ^{31}P NMR spectra were collected at 101.25 MHz on a Bruker Avance 250 spectrometer using a 5-mm inverse $^1\text{H}/^{13}\text{C}/^{31}\text{P}/^{19}\text{F}$ probe head at room temperature. The inverse-gated proton-decoupling sequence was applied with 4- μs time, 30° pulse, 1-s FID acquisition time, and 1.2-s acquisition delay, and 21,000 scans collected. Chemical shifts were referenced to an external 85% H_3PO_4 standard at 0.00 ppm. All NMR samples were prepared by mixing 0.6 μl of 0.5 M Na_2HPO_4 solution (pH 7.5) and 0.06 μl of D_2O .

RESULTS

Detection and Elimination of Pyrophosphate Traces from Phosphate Buffer—We determined that untreated phosphate buffers contained $\sim 0.01\%$ pyrophosphate contamination (Fig. 2). Because pyrophosphate has high affinity for myosin ($K_{d,\text{PP}_i} = 200$ nM), even this small concentration of pyrophosphate can block the binding of the substrates with lower affinity such as ADP ($K_{d,\text{ADP}} \sim 1$ μM) or phosphate ($K_{d,\text{P}_i} \sim 10$ mM). However, according to the NMR spectroscopy results, heat treatment at alkaline pH (see “Experimental Procedures”) can reduce the pyrophosphate level by 5- to 10-fold and enabled us to investigate reactions of myosin and substrates at high phosphate concentrations. A detailed description of the experiments with the related figures (supplemental Fig. S1, a–c) can be found in the supplemental materials.

Phosphate Binding of W501+•ADP Complex—To study ligand binding and the possible reverse reaction of the recovery step ($\text{M}^+\text{•ADP•P}_i \leftrightarrow \text{M}^*\text{•ADP•P}_i$), W501+ was preincubated with ADP and then rapidly mixed with different concentrations of a phosphate solution in the stopped-flow apparatus. The ionic strength of the solutions was kept constant in all experiments by adding an appropriate amount of NaCl. Also the ionic strength of the solutions was identical in both syringes of the stopped-flow apparatus. The reactions were studied at $I = 244$ mM at pH 7.2 and $I = 118$ mM at pH 7.2 and 6.7. At all phosphate concentrations studied, large and slow fluorescence increases (35% enhancement with $k_{\text{obs}} = 0.14$ s^{-1} at 25 mM phosphate concentration, pH 7.2, $I = 118$ mM) were detected, which indicates the formation of $\text{M}^*\text{•ADP•P}_i$. Because the rate constant of ADP release is 5 s^{-1} and accompanied with a 10–15% fluorescence increase, the larger than 15% amplitude increase and the slower than 5 s^{-1} observed rate constants rule out this fluorescence enhancement being simply due to dequenching on displacement of ADP from $\text{M}^+\text{•ADP}$. Upon phosphate binding to W501+, in the absence of ADP, a small ($<3\%$) quench was detected. At each applied ionic strength and pH, stopped-flow

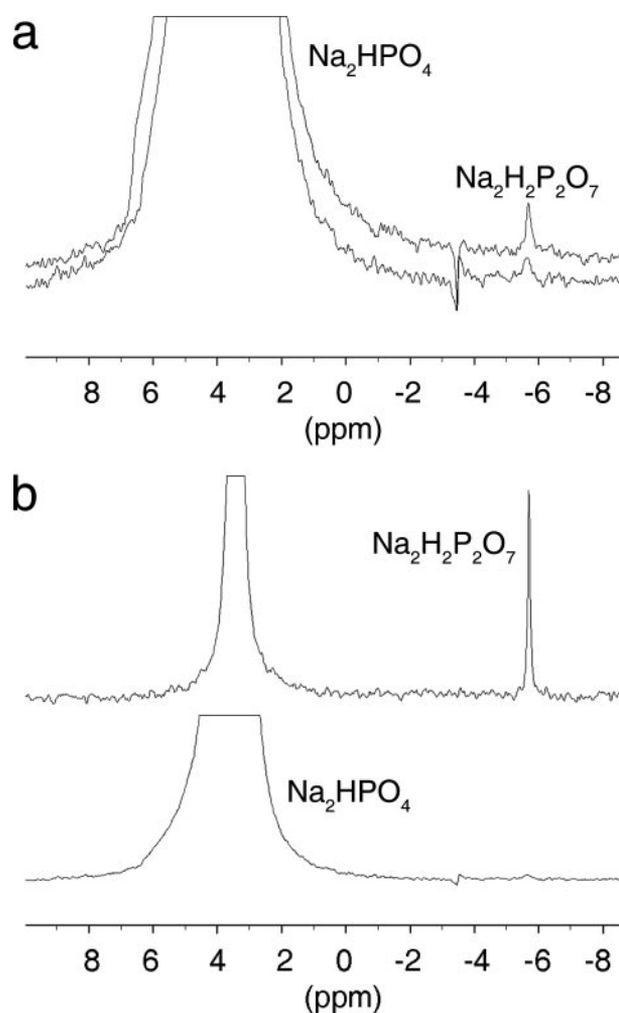


FIGURE 2. Determination of the contaminating pyrophosphate traces in phosphate buffer by NMR spectroscopy. a, $^{31}\text{P}\{^1\text{H}\}$ NMR spectra of samples containing 0.6 μl of 0.5 M Na_2HPO_4 (pH 7.5) and 0.06 μl of D_2O . The bottom spectrum belongs to the boiled and the top spectrum to the non-treated solution demonstrating that boiling the buffer for 30 min at 121 °C in an autoclave can reduce the pyrophosphate level 5- to 10-fold. b, $^{31}\text{P}\{^1\text{H}\}$ NMR spectra of samples containing 0.6 μl of boiled 0.5 M Na_2HPO_4 solution (pH 7.5) and 0.06 μl of D_2O , recorded with 21,000 scans (bottom), and after addition to this solution of 30 μl of 0.1 M $\text{Na}_2\text{H}_4\text{P}_2\text{O}_7$ (pH 7.5), recorded with 54 scans (top). This experiment demonstrates that the contaminating compound in phosphate buffer, which can be reduced by heat treatment, is pyrophosphate.

records could be fitted with single exponentials (Fig. 3a). In addition, a very slow (0.005 s^{-1}) fluorescence quench also was observed that is the consequence of binding of residual pyrophosphate. This reaction was faster and became dominant when untreated phosphate samples were used (see supplemental Fig. S1c). The amplitudes and the observed rate constants of the fluorescence enhancement increased with phosphate concentrations. The slope of the initial linear fit of the plot of rate constants versus phosphate concentration ($k_{\text{Pi,on,app},118\text{mM},\text{pH}7.2}^{\text{W501+•ADP}} = 0.0036$ $\text{mM}^{-1}\text{s}^{-1}$) depends on both the ionic strength and pH (Fig. 3b), whereas the extrapolated intercepts ($k_{\text{Pi,on,app},118\text{mM},\text{pH}7.2}^{\text{W501+•ADP}} = 0.052$ s^{-1}) were practically unchanged. From these values the dissociation constant of phosphate for W501+•ADP complex is $K_{d,\text{app},\text{P}_i} = 14.2$ mM at 118 mM ionic strength. The intercept value is the same as the basal ATPase rate constant. This indicates that k_{+4} determines the ATPase

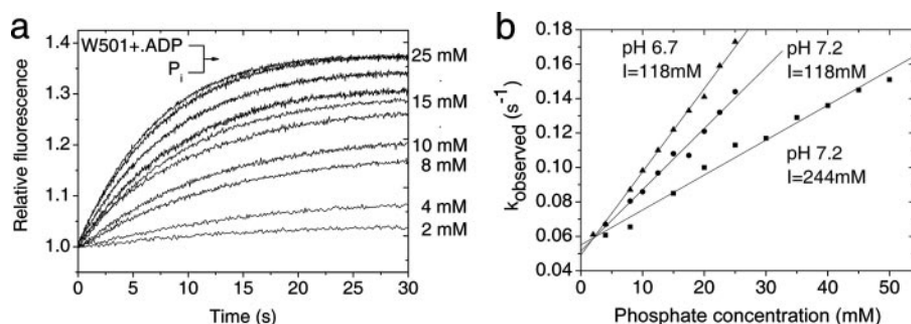


FIGURE 3. **Phosphate binding of W501 + ADP.** *a*, stopped-flow records of 3 μM W501 + ADP complex rapidly mixed with different amount of phosphate (2–25 mM), while the ionic strength ($I = 118$ mM) and pH (pH 7.2) were kept constant. The large enhancement in fluorescence shows that the $\text{M}^*\text{-ADP}\cdot\text{P}_i$ complex develops upon phosphate binding. *b*, plot of the observed rate constants of the reaction (described in *a*) against phosphate concentration at different ionic strengths ($I = 118$ mM or 244 mM) or pH (pH 6.7 or 7.2). From this plot the on- and off-rate constants of phosphate binding sensed by Trp-501 can be calculated: $k_{\text{on,app}}^{\text{W501+ADP}} = 0.0036 \text{ mM}^{-1}\text{s}^{-1}$ and $k_{\text{off,app}}^{\text{W501+ADP}} = 0.052 \text{ s}^{-1}$ (other parameters are shown in Table 2).

TABLE 2

Summary of the rate and equilibrium constants measured in this study

Each section shows the experiment from which the values are originated. (Arrangement of stopped-flow reactions are simplified as: component(s) in one syringe \times component in the other syringe.)

	Measured constants
W501 + ADP \times P_i	
P_i half sat ($I = 118$ mM, pH 6.7)	12 mM
P_i half sat ($I = 118$ mM, pH 7.2)	14.2 mM
P_i half sat ($I = 244$ mM, pH 7.2)	23 mM
$k_{\text{on,Pi}}$ ($I = 118$ mM, pH 6.7)	$0.0048 \text{ mM}^{-1}\text{s}^{-1}$
$k_{\text{on,Pi}}$ ($I = 118$ mM, pH 7.2)	$0.0036 \text{ mM}^{-1}\text{s}^{-1}$
$k_{\text{on,Pi}}$ ($I = 244$ mM, pH 7.2)	$0.0020 \text{ mM}^{-1}\text{s}^{-1}$
$k_{\text{off,Pi}}$ ($I = 118$ mM, pH 6.7)	0.049 s^{-1}
$k_{\text{off,Pi}}$ ($I = 118$ mM, pH 7.2)	0.052 s^{-1}
$k_{\text{off,Pi}}$ ($I = 244$ mM, pH 7.2)	0.055 s^{-1}
$k_{\text{max,Pi}}$ ($I = 118$ mM, pH 7.2)	$\sim 0.2 \text{ s}^{-1}$
$K_{\text{app,Pi}}$ ($I = 118$ mM, pH 7.2)	14.2 mM
W501 + $\text{P}_i \times$ ADP	
$K_{\text{app,ADP}}$	45 μM
$K_{\text{on,ADP}}$	$0.011 \mu\text{M}^{-1}\text{s}^{-1}$
$k_{\text{max,ADP}}$	0.2 s^{-1}
W129 + $\text{P}_i \times$ ADP	
ADP half sat	15 μM
$k_{\text{PD,on}}$	$1.6 \mu\text{M}^{-1}\text{s}^{-1}$
$k_{\text{PD,off}}$	19 s^{-1}
$k_{\text{max,ADP}}$	370 s^{-1}
W129 + ADP \times P_i	
k_{obs}	$7.5 \pm 1 \text{ s}^{-1}$
W129 + \times P_i	
$k_{\text{fast,on,Pi}}^a$	$50.3 \text{ mM}^{-1}\text{s}^{-1}$
$k_{\text{fast,off,Pi}}^a$	35.0 s^{-1}
$k_{\text{fast,max,Pi}}^a$	300 s^{-1}
k_{slow}^a	5 s^{-1}
W129 + \times ATP	
$k_{\text{ATP,fast}}$	1140 s^{-1}
$k_{\text{ATP,slow}}$	350 s^{-1}
W129 + $\text{P}_i \times$ ATP	
$k_{\text{P,off}}$	260 s^{-1}
W501 + dmADP \times ATP	
k_{obs}	2.5 s^{-1}
W501 + dmADP \cdot $\text{P}_i \times$ ATP	
k_{fast}	5.8 s^{-1}
k_{slow}	0.04 s^{-1}
$A_{\text{fast}}/A_{\text{slow}}$	0.50
W501 + \times ADP	
$k_{\text{D,on}}$	$1.2 \mu\text{M}^{-1}$
$k_{\text{D,off}}$	7 s^{-1}
$K_{\text{D}} (K_{\text{AD}})$	$0.17 \mu\text{M}^{-1}$ (5.8 μM)
W501 + ADP \cdot AIF ₄ \times PyA	
$K_{\text{D}} \text{ A}\cdot\text{M}^*\text{-ADP}\cdot\text{P}_i$	57 μM

^a Results can be found in the supplemental material.

rate constant of the cycle. It also supports the conclusion that the $\text{M}^*\text{-ADP}\cdot\text{P}_i$ state is the one generated by addition of ADP and phosphate to W501 +. The parameters under other condi-

tions are presented in Table 2. Because the increase of phosphate concentration was limited by the constant ionic strength of the solution, it could not be increased above 25 mM at pH 7.2, therefore the saturation of values of rate constants could not be determined. Furthermore, it is difficult to characterize these values, because rate constants are strongly dependent on ionic strength and it is hard to maintain a constant value with different counter ions. Even a little overcompensation of the ionic strength level with sodium chloride could cause the apparent saturation of the observed

rate constant. In summary, there is some indications that the rate constant might saturate at around $k_{\text{Pi,max,app}}^{\text{W501+ADP}} \approx 0.2 \text{ s}^{-1}$. To overcome this uncertainty we carried out the inverse experiment in which W501 + was premixed with phosphate and then rapidly mixed with ADP.

ADP Binding of W501 + P_i Complex—W501 + was preincubated with 25 mM phosphate at pH 7.2 and rapidly mixed in the stopped-flow apparatus with different concentration of ADP. Both stopped-flow syringes contained 25 mM phosphate to keep phosphate concentration constant at pH 7.2 and $I = 118$ mM. Figs. 4*a* and 6 show a stopped-flow record of the reaction of W501 + P_i complex with 350 μM ADP. A small (5%) and fast (400 s^{-1}) fluorescence quench was followed by a 25% fluorescence increase. The apparent rate constant of the fluorescence increase depended on the ADP concentration (Fig. 4*b*), the initial slope of the plot is $k_{\text{ADP,on,app}}^{\text{W501+Pi}} = 0.011 \mu\text{M}^{-1}\text{s}^{-1}$. The observed rate constant of the reaction saturated at $k_{\text{ADP,max,app}}^{\text{W501+Pi}} = 0.20 \text{ s}^{-1}$, which is a similar value to the estimated maximum rate constant of phosphate binding of W501 + ADP complex ($k_{\text{Pi,max,app}}^{\text{W501+ADP}} \approx 0.2 \text{ s}^{-1}$).

ADP Binding of W129 + P_i Complex—In contrast to Trp-501, which senses the lever arm swing, Trp-129 is sensitive to the binding events of ligands (17, 18). The fluorescence of Trp-129 is quenched by all adenosine nucleotides (ADP, ATP, and ADP \cdot P_i) and its phosphate analog (ADP \cdot AIF₄) by 55%, whereas phosphate binding slightly (10%) increases its fluorescence. Using W501 + and W129 +, we can potentially distinguish the lever arm swing from product binding events if the kinetics is favorable. When 4 μM W129 + was mixed with different concentrations of ADP (pH 7.2, $I = 118$ mM) in the presence of a saturating amount of phosphate, a large fluorescence quench was detected (Figs. 5 and 6). Single exponentials were fitted to all records. Increasing the ADP concentration increased both the amplitudes and the observed rate constants of the fluorescence changes. At higher ADP concentrations the rate constant was so high that significant amplitude loss occurred during the dead time of the stopped-flow apparatus. A hyperbola was fitted to the plot of rate constants versus ADP concentration, which yielded an initial slope, $k_{\text{ADP,on,app}}^{\text{W129+Pi}} = 1.6 \mu\text{M}^{-1}\text{s}^{-1}$ and an intercept (extrapolated observed rate constant at zero ADP concentration), $k_{\text{ADP,off,app}}^{\text{W129+Pi}} = 19 \text{ s}^{-1}$. The hyperbola saturates

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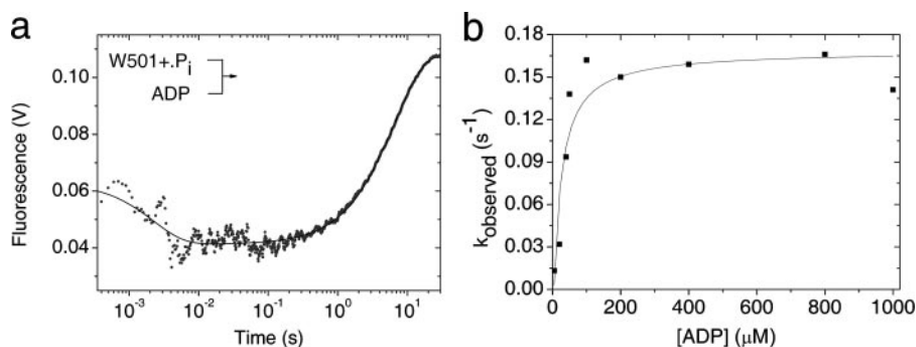


FIGURE 4. **ADP binding of W501+ in the presence of phosphate.** *a*, stopped-flow record of 3 μM W501+ mixed with 350 μM ADP in the presence of 25 mM phosphate on both sides. A fast (400 s^{-1}) quenching phase with small amplitude change (5%) is followed by an [ADP]-dependent slow (0.2 s^{-1}) increasing phase. This experiment also confirms that, in the presence of ADP and phosphate, the closed (Trp-501 high fluorescence) state can be developed. *b*, plot of the observed rate constants of the ADP binding to W501+ \cdot P_i complex. The fitted hyperbola saturated at $k_{\text{ADP,max,app}}^{\text{W501+}\cdot\text{P}_i} = 0.20\text{ s}^{-1}$.

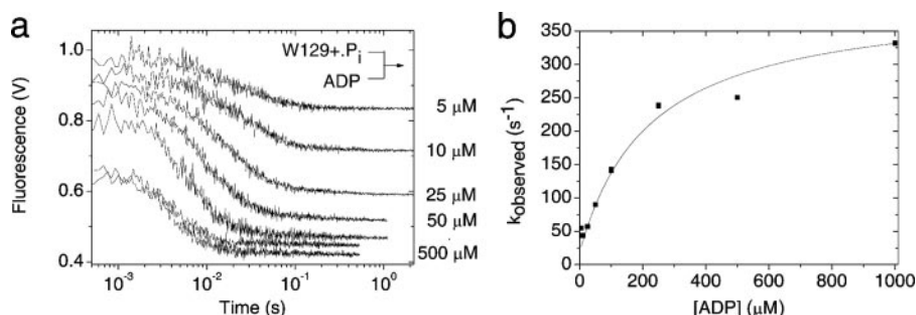


FIGURE 5. **ADP binding of W129+ in the presence of phosphate.** *a*, stopped-flow records of the ADP binding reaction of 4 μM W129+ in the presence of 25 mM phosphate. The fluorescence quench sensed by Trp-129 responds to the nucleotide binding. Note that ADP binding to M \cdot P_i complex results in a fast fluorescent decrease in the case of W129+ but causes a slow fluorescence enhancement in the W501+. At high ADP concentrations significant amplitude is lost in the dead time of the stopped-flow. *b*, single exponentials were fitted to the traces and the observed rate constants were plotted against [ADP], which showed the maximal rate constant for the reaction is $k_{\text{ADP,max,app}}^{\text{W129+}\cdot\text{P}_i} = 370\text{ s}^{-1}$. The on- and off-rate constants of ADP binding to M \cdot P_i was calculated from the initial slope and the intercept of the fitted hyperbola, respectively: $k_{\text{ADP,on,app}}^{\text{W129+}\cdot\text{P}_i} = 1.6\text{ }\mu\text{M}^{-1}\text{ s}^{-1}$ and $k_{\text{ADP,off,app}}^{\text{W129+}\cdot\text{P}_i} = 19\text{ s}^{-1}$.

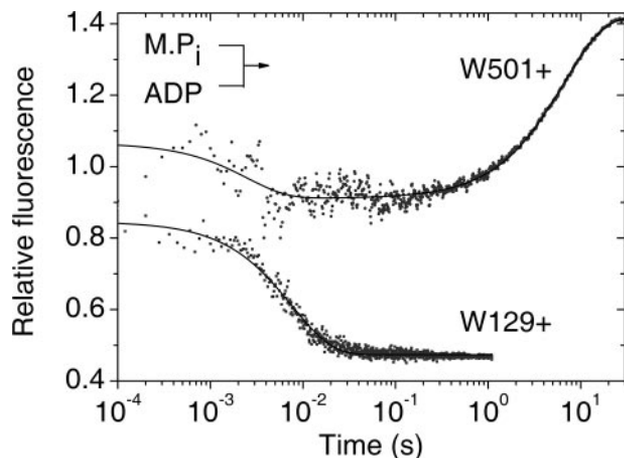


FIGURE 6. **Stopped-flow records of ADP binding to the M \cdot P_i complex for W501+ (upper trace) and W129+ (lower trace).** Please note that the data used here are from Figs. 3 and 4. They are normalized to a starting fluorescence of 1 to make the amplitudes comparable. Trp-129 responds to ADP binding to W129+ \cdot P_i with a fast fluorescence quench ($\sim 150\text{ s}^{-1}$ at 100 μM ADP), whereas the same reaction produces a small fast quench followed by a dominant slow ($\sim 0.15\text{ s}^{-1}$ at 350 μM ADP) fluorescence enhancement of Trp-501. This implicates that W129+ is sensitive for the nucleotide binding itself, whereas W501+ senses the large isomerization step coupled with the power stroke of myosin. Using these two mutants allowed us to study the properties of the nucleotide binding pocket separately from the isomerization step.

at the rate constant $k_{\text{ADP,max,app}}^{\text{W129+}\cdot\text{P}_i} = 370\text{ s}^{-1}$, however, it is probably underestimated because of the amplitude loss in the dead time. The half-maximum of the amplitude of the fluorescence change was at $[\text{ADP}]_{\text{half-sat}}^{\text{W129+}\cdot\text{P}_i} = 15\text{ }\mu\text{M}$.

Phosphate Binding of W129+ \cdot ADP Complex—When W129+ \cdot ADP complex was rapidly mixed with phosphate in a stopped-flow apparatus, a small fluorescence change was detected. Because the fluorescence level of W129+ \cdot ADP and W129+ \cdot ADP \cdot P_i are identical, the small signal change was the consequence of the re-equilibration of the high (apo and W129+ \cdot P_i) and the low (W129+ \cdot ADP and W129+ \cdot ADP \cdot P_i) fluorescence states when the enzyme was not completely saturated with ligands. Supplemental Fig. S2 shows the stopped-flow records of the reaction when 4 μM W129+ was mixed with 50 mM phosphate at pH 7.2, I = 118 mM, while the same ADP concentration was maintained in both syringes. The records show the reactions at four different ADP concentrations (50 μM , 150 μM , 450 μM , and 1 mM). At 1 mM ADP concentration, a 3.5% fluorescence increase was detected whose amplitude increased to 17% at 50 μM ADP concentrations. Single exponentials were fitted to the records. In the range of ADP concentrations studied, the observed rate constants were similar ($7.5\text{ s}^{-1} \pm 1\text{ s}^{-1}$) (for more details see supplemental Fig. S2, *a–d*).

Phosphate Binding of W129+—In contrast to W501+, W129+ responds with a small (10–15%) fluorescence increase upon phosphate binding. The kinetics of the signal change when 5 μM W129+ was mixed with different concentration of phosphate were characterized at pH 7.2, I = 118 mM. The records were fitted with double exponentials (supplemental Fig. S3*a*), in contrast to the records of W129+ \cdot ADP mixed with phosphate (see above). The observed rate constant of the slow phase was 5 s^{-1} and independent of the phosphate concentrations studied (supplemental Fig. S3*c*). The rate of the fast phase was dependent on and plotted against phosphate concentration and the plot between 0.5 and 5 mM phosphate concentrations was fitted with linear (supplemental Fig. S3*b*). The slope was $k_{\text{Pi,on,app}}^{\text{W129+}} = 50.3\text{ mM}^{-1}\text{ s}^{-1}$ and its intercept $k_{\text{Pi,off,app}}^{\text{W129+}} = 35.0\text{ s}^{-1}$, and the rate constant of the fast phase was maximum at $k_{\text{Pi,max,app}}^{\text{W129+}} = 300\text{ s}^{-1}$ at 5 mM phosphate concentration and did not exceed 300 s^{-1} even at 25 mM phosphate concentration. We were interested whether this phosphate binding site was part of the functional ATP binding site or an alternative

phosphate binding region sensed by W129+. To test these possibilities, phosphate bound to W129+ was chased by large excess of ATP, and the record was compared with the reaction record when W129+ was mixed with ATP in the absence of phosphate. In the latter reaction the fluorescence change has two phases, as described in an earlier study (17). The observed rate constants of the slow and fast phases were $k_{\text{ATP,slow}}^{\text{W129+}} = 350 \text{ s}^{-1}$ and $k_{\text{ATP,fast}}^{\text{W129+}} \geq 1000 \text{ s}^{-1}$. When motor domain was complexed with phosphate and chased with ATP, the fluorescence change became a single phase and its observed rate constant slowed to $k_{\text{Pi,off,ATPchase}}^{\text{W129+}} = 260 \text{ s}^{-1}$ (supplemental Fig. S4). When W129+ was not saturated with phosphate, a slow and a fast phase could be distinguished. The ratios of the amplitudes of these phases were difficult to determine because of the significant loss of the fast phase in the dead time of the stopped-flow apparatus.

Because the ATP chasing experiment showed a much higher off-rate constant for phosphate than the off-rate constant determined by the intercept in the phosphate binding experiment (260 s^{-1} and 35 s^{-1} , respectively), and also the binding constants differed significantly in these experiments, we suggest that, in the direct P_i binding experiment, an alternative phosphate binding site is involved. This is presumed to be close to the substrate binding pocket, because Trp-129 senses its binding, but it does not overlap with the functional ATP binding site, because ATP does not displace it (*i.e.* there was no slow phase of ATP binding limited by the off-rate constant of 35 s^{-1}).

Dissociation of ADP from $M \cdot \text{ADP} \cdot \text{P}_i$ by Chasing dmADP —To study the mechanism of the dissociation of ADP and phosphate directly, $8 \mu\text{M}$ W501+ motor domain was preincubated with $100 \mu\text{M}$ dmADP (premix concentrations) in the presence and absence of 42 mM phosphate, and the ligands were chased with 2 mM ATP (dmADP was prepared by incubation of dmATP with W501+ for 15 min at 20°C). In both experiments, the mant fluorescence signal decrease was followed, which is limited by the dissociation of bound dmADP . In the absence of phosphate, a single exponential fitted to the fluorescence record to yield a rate constant, $k_{\text{off,mantADP}} = 2.5 \text{ s}^{-1}$. The off-rate constant of mantADP is slightly smaller than that of ADP (7 s^{-1}) (17). This effect is caused by the mant group, which interacts with the protein (32), and results in higher affinity of dmADP to myosin (33–35). By contrast, in the presence of phosphate, two phases of displacement were detected. The rate constants of the fast and the slow phases were $k = 5.8 \text{ s}^{-1}$ and $k = 0.04 \text{ s}^{-1}$, whereas the ratio of their amplitudes was $A_{\text{fast}}/A_{\text{slow}} = 0.50$ (Fig. 7). The slower rate constant indicates that some $M^* \cdot \text{dmADP} \cdot \text{P}_i$ formed during the preincubation reaction.

Actin Binding of $W501+ \cdot \text{ADP} \cdot \text{AlF}_4$ Complex—We investigated actin binding of the $W501+ \cdot \text{ADP} \cdot \text{AlF}_4$ complex by following the light scattering signal. $109 \mu\text{M}$ W501+ was mixed with 1 mM ADP, 1 mM AlCl_3 , and 5 mM NaF on ice for 3 h, and the formation of the complex was followed by the fluorescent steady-state emission intensity enhancement of Trp-501. To investigate the actin binding properties of the complex, a varying amount of $W501+ \cdot \text{ADP} \cdot \text{AlF}_4$ and $1 \mu\text{M}$ F-actin was mixed in a stopped-flow apparatus and light scattering signal was followed. Very fast reactions ($>1000 \text{ s}^{-1}$) were detected even at

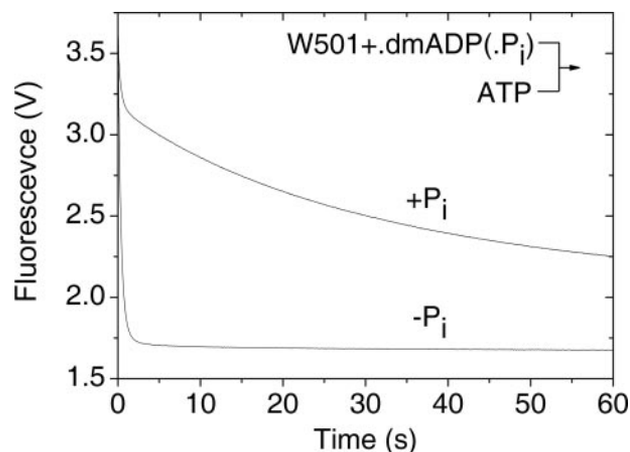


FIGURE 7. Stopped-flow records of $8 \mu\text{M}$ W501+ complexed with $100 \mu\text{M}$ dmADP and mixed with 2 mM ATP in the absence (lower trace) or in the presence (upper trace) of 42 mM phosphate (premix concentrations). The fast (2.5 s^{-1}) ADP release step sensed by mant fluorescence enhancement shows single exponential decay in the absence of phosphate, however, in the presence of phosphate the fast phase (5.8 s^{-1}) is followed by a slow (0.04 s^{-1}) phase. The ratio of the amplitudes in the latter case is $A_{\text{fast}}/A_{\text{slow}} = 0.50$.

low $W501+ \cdot \text{ADP} \cdot \text{AlF}_4$ concentrations, and most of the signal change was lost in the dead time of the stopped-flow apparatus. The signal in the stopped-flow apparatus was so stable that almost all of the single traces overlapped with each other. This enabled us to determine the starting and the endpoints of the reactions at specific $W501+ \cdot \text{ADP} \cdot \text{AlF}_4$ concentrations where the signal difference represented the total amplitude of the reaction. The amplitudes of the reactions were plotted against W501+ concentration, and the fitted hyperbola resulted in the dissociation constant $K_d \cdot \text{AlF}_4 \cdot \text{actin} = 57 \mu\text{M}$ (supplemental Fig. S5).

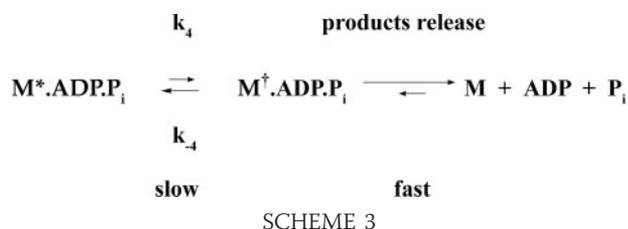
Actin-activated ATPase Experiments with W501+—Actin-activated steady-state ATPase activity was measured on W501+ by a pyruvate kinase/lactate dehydrogenase-coupled assay measurement. Activities were determined in the presence of different amount of F-actin and were plotted against actin concentration. The maximum of actin-activated ATPase activity was $v_{\text{max}} = 3.8 \text{ s}^{-1}$, and half-saturation was determined at $K_m = 67 \mu\text{M}$.

DISCUSSION

In the Bagshaw-Trentham scheme, the relationship between the reverse recovery step (K_4) and the product release steps (K_5 , K_6 , and K_7) had not been fully resolved. Because Trp-129 and Trp-501 are very sensitive signals for ligand binding and lever arm motion, respectively, the kinetics of the two processes can be investigated separately. To exploit these probes to measure phosphate binding, we found it necessary to minimize pyrophosphate contamination by boiling the stock phosphate solution. It is a well known requirement that vanadate solutions need to be boiled to remove polymeric species before using orthovanadate as phosphate analog for myosin product complex formation (36, 37). However, it is less appreciated that phosphate stock solutions need similar treatment.

With this experimental approach, the mechanism of the basal ATPase activity can be elucidated, and we can compare the processes to the “actin-induced” ATPase mechanism. In the

Mechanism of Phosphate Release of Myosin II



following sections we describe step by step how the relevant kinetic parameters were determined.

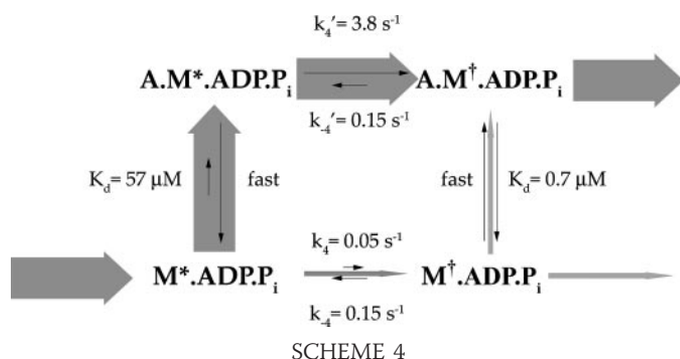
Separation of the Lever Arm Motion from the Ligand Binding/Release Processes—Previous studies have verified that the measured rate constants of W129+, W501+, and the wild-type motor domain are similar, and thus the introduced mutations do not change the kinetic mechanism significantly (17–19). Because Trp-129 and Trp-501 are located at the nucleotide binding pocket and in the relay helix, respectively, upon ADP and phosphate binding, different local molecular events are detected by following tryptophan fluorescence changes (Fig. 6). To follow ligand binding events, W129+ was mixed with ADP in the presence of 25 mM phosphate, and a large, relatively fast ($k_{\text{ADP,max,app}}^{\text{W129+P}_i} = 370 \text{ s}^{-1}$) fluorescence decrease was detected. This event is slightly slower than ADP binding in the absence of phosphate. In an earlier study we showed that Trp-129 senses only ligand binding events, and it is insensitive to the conformational changes that occur in the switch 2 and the relay/converter regions. Under the same conditions, when W501+·P_i was mixed with ADP a much slower fluorescence increase ($k_{\text{ADP,max,app}}^{\text{W501+P}_i} = 0.20 \text{ s}^{-1}$) was detected than in the case of W129+·P_i (Fig. 6). The Trp-501 fluorescence change responds to the large isomerization coupled to the lever arm swing (18). Because the fluorescence change of Trp-129, located at the binding pocket, is much faster than that of Trp-501, located in the relay region, it indicates that ADP and phosphate binding events occur separately from the lever arm swing. Consequently, in the basal ATPase cycle, the rate-limiting step is the slow lever arm swing ($\text{M}^* \cdot \text{ADP} \cdot \text{P}_i \leftrightarrow \text{M}^\dagger \cdot \text{ADP} \cdot \text{P}_i$) and not the phosphate release step. Thus the supposition in the original Bagshaw-Trentham scheme has been validated (3, 4). In Scheme 3 below, k_{-4} is the rate of the recovery step with bound ADP·P_i (it is important to note that this is different from the k_{+3a} of the recovery step with bound ATP). k_4 is the rate of the reverse recovery step with bound ADP·P_i.

If the slow fluorescence increase of W501+ ($k_{\text{ADP,max,app}}^{\text{W501+P}_i}$) is assigned to the lever arm swing, the relaxation of this equilibrium is $k_{\text{ADP,max,app}}^{\text{W501+P}_i} = k_4 + k_{-4}$. In the absence of phosphate, the basal ATPase rate constant ($k_{\text{basal}} = 0.05 \text{ s}^{-1}$) is determined by k_4 , therefore $k_{-4} = k_{\text{ADP,max,app}}^{\text{W501+P}_i} - k_4 = 0.15 \text{ s}^{-1}$, and the equilibrium constant is $K_4 = k_4/k_{-4} = 0.33$. The mechanism is confirmed by the fact that a small, fast fluorescence quench was detected in the reaction when W501+·P_i was mixed with ADP (Figs. 4 and 6). The quench represents the initial M[†]·ADP·P_i state, which goes through the slow isomerization reaction resulting M*·ADP·P_i state. The rate constant of the fast quench is as fast as the rate constants of ADP binding of W129+·P_i, which also indicates that binding events are being monitored in both cases.

This two-step mechanism is also confirmed by the result of the experiment when dmADP was chased by ATP in the presence of saturating concentration of phosphate (Fig. 7). The observed two phases correspond to the two populations of the M·dmADP·P_i complex. The fast phase is the release of the ligands from M[†], and the slow phase represents the reverse recovery step, because shifting this pre-equilibrium (which is established in the preincubation of myosin with ADP and phosphate) becomes the rate-limiting step of the ligand release after the quick release of ADP and phosphate from M[†]. The ratio of the amplitudes of the two phases ($A_{\text{fast}}/A_{\text{slow}} = 0.50$) gives the equilibrium constant of the reverse recovery step. Taking the values calculated for the reverse recovery step ($K_4 = 0.33$) and for phosphate affinity for M·ADP complex ($K_{\text{DP}} = 42 \text{ mM}$), the expected ratio of the amplitudes of the two phases should be $A_{\text{fast}}/A_{\text{slow}} = 0.78$. The difference from the expectation could be the effect of the mant group of dmADP, which has been demonstrated to affect the affinity of ADP (33–35), and probably influences the binding properties of phosphate as well. If we suppose that the equilibrium between the M*·ADP·P_i and M[†]·ADP·P_i is not affected by the mant group, then a 2.7-fold stronger binding of phosphate to M·ADP ($K_{\text{DP}} = 9.6 \text{ mM}$ instead of 42 mM) is indicated. On the other hand, if phosphate affinity was unaltered, the equilibrium of the reverse recovery step would change a little: $K_4 = 0.12$ for dmADP instead of 0.33 for ADP.

How Does Actin Activation Occur?—As we showed above, the rate-limiting step of the basal ATPase activity is the reverse recovery step, whereas dissociation of ADP and phosphate is comparatively fast. Previous studies showed that actin causes relatively small changes (2–5 times) in the rate constants of ligand binding/dissociation and of the recovery step (38). Because actin activation causes a 76 times overall increase of the ATPase ($V_{\text{max}} = 3.8 \text{ s}^{-1}$ from 0.05 s^{-1}), the largest effect of actin must be on the rate-limiting step of basal ATPase. Thus, the approximate rate of the actin-induced power stroke is then $k_4' = 76 * k_4 = 3.8 \text{ s}^{-1}$ for the $\text{A} \cdot \text{M}^* \cdot \text{ADP} \cdot \text{P}_i \rightarrow \text{A} \cdot \text{M}^\dagger \cdot \text{ADP} \cdot \text{P}_i$ reaction. Note that this reaction starts from the weak actin binding conformation of myosin, in which switch 2 is still closed, however actin binding might induce the opening of the switch 1 loop. Very recently this state was kinetically characterized by Jahn (39) with an ATP analogue (γ -amido-ATP), which favors the closed switch 2 state. He observed actin binding of myosin II with closed switch 2 pre-power stroke conformation. However, k_{-4}' still remains unresolved by a direct experimental measurement, but our simulation results support the assumption that it is very similar to $k_4 = 0.15 \text{ s}^{-1}$. Otherwise the flux through the actin-detached M[†]·ADP·P_i state (see Scheme 4) would dominate the flux via the actin-attached A·M[†]·ADP·P_i state and lead to wasteful ATP hydrolysis. If we assume that the rate constant of the reverse step of the power stroke (k_{-4}') is approximately the same as that of the reverse step of the reverse recovery step (k_{-4}), i.e. $k_{-4}' = k_{-4} = 0.15 \text{ s}^{-1}$, then the equilibrium constant for the lever arm movement increases from $K_4 = 0.33$ in the absence of actin to $K_4' = 25$ upon actin-activation. This equilibrium constant represents a $\Delta\Delta G_0 = 10 \text{ kJ/mol}$ difference in free energy change. In other words, actin pushes the equilibrium toward the lever arm-down, post-power

stroke state. This energy change accompanying the swing of the lever arm ($M^* \rightarrow M^\dagger$) is paid for the improved binding to actin, as the K_d goes from $57 \mu\text{M}$ for $A \cdot M^* \cdot \text{ADP} \cdot \text{P}_i$ to $0.7 \mu\text{M}$ for $A \cdot M^\dagger \cdot \text{ADP} \cdot \text{P}_i$, *i.e.* changing by the factor of 81. The strength of actin binding further increases in the following steps of product release: the whole sequence of actin dissociation constants between the pre-power stroke and rigor states decreases as follows: $K_{d,A \cdot M^* \cdot \text{ADP} \cdot \text{P}_i} = 57 \mu\text{M}$, $K_{d,A \cdot M^\dagger \cdot \text{ADP} \cdot \text{P}_i} = 0.7 \mu\text{M}$, $K_{d,A \cdot M^* \cdot \text{ADP}} = 0.12 \mu\text{M}$, and $K_{d,A \cdot M} = 0.03 \mu\text{M}$ (40). This stronger actin binding is the driving force that pulls the reaction through the weakly bound $A \cdot M^* \cdot \text{ADP} \cdot \text{P}_i$ state (see Scheme 4). As a consequence, despite the fact that $M^* \cdot \text{ADP} \cdot \text{P}_i$ binds actin very weakly ($K_d = 57 \mu\text{M}$), the main flux of the reaction occurs via the actin bound form, even if the actin concentration is low (Fig. 8a). Our kinetic models show that at $0.5 \mu\text{M}$ actin concentration, the flux of the actin-bound and -unbound pathways are equal. By increasing actin concentration the ratio of the fluxes



dramatically increases: *e.g.* at $5 \mu\text{M}$ actomyosin concentration the ratio is $\text{Flux}_{\text{actin-bound}}/\text{Flux}_{\text{actin-unbound}} = 14$ (Fig. 8b).

In earlier studies, phosphate release was proposed to precede the working-stroke step in the actin-attached states (21, 38, 41) based on the assumption that the $A \cdot M^* \cdot \text{ADP} \cdot \text{P}_i \rightarrow A \cdot M^\dagger \cdot \text{ADP} \cdot \text{P}_i$ transition is not poised strongly to the right, *i.e.* $k_{-4}' \gg k_{-4} = 0.15 \text{ s}^{-1}$. If this assumption was valid, this would imply that phosphate release provides the required free energy change for the working stroke, which therefore should precede the lever arm swing avoiding significant futile cycle. This is inconsistent with physiological studies, which have demonstrated significant tension generation before phosphate release, thus demanding significant free energy change from the $A \cdot M^* \cdot \text{ADP} \cdot \text{P}_i$ to $A \cdot M^\dagger \cdot \text{ADP} \cdot \text{P}_i$ transition itself (2, 11–14, 42, 43). Takagi *et al.* (14) also demonstrated in single fiber experiments that the binding of the $M \cdot \text{ADP} \cdot \text{P}_i$ complex to actin is a fast and weak binding process. They proposed a scheme, which was supported by other studies (11–13) in which the weak actin binding $A \cdot M^* \cdot \text{ADP} \cdot \text{P}_i$ state undergoes an isomerization to a strongly bound tension generating state. This is then followed by phosphate release, which might strengthen further the actin-myosin interaction. The model based on solution kinetic measurements presented here is consistent with this mechanism, because they show that the order of events is: weak actin binding occurs first, then the actin activated initiation of the power stroke, and finally the product release steps. However, in contrast to previous studies (14, 44–46), which suggested that the weak-to-strong transition of actin binding should precede the power stroke, our results suggest that the weak-to-strong transition does not necessarily precede the lever arm swing of the power stroke and they might be concurrent processes.

This mechanism also answers another interesting question: if the recovery step that occurs in the $M \cdot \text{ATP}$ form is a simple reversal of the power stroke that occurs in $M \cdot \text{ADP} \cdot \text{P}_i$, and both nucleotide states have weak actin affinities, why does the first occur in actin detached form, while the latter reaction must happen predominantly in actin-bound form (otherwise a futile cycle would occur)? The answer is that the recovery step is rapid in the absence of actin and is not activated by actin (38), whereas the power stroke step (and not phosphate release) is accelerated by actin, which pulls the reaction flux toward the actin-bound reaction pathway (Scheme 4). In other words, actin activation diverts the reaction flow to the actin-bound pathway inducing the actin-bound power stroke (Fig. 8). Consequently, there is a kinetic reason why the

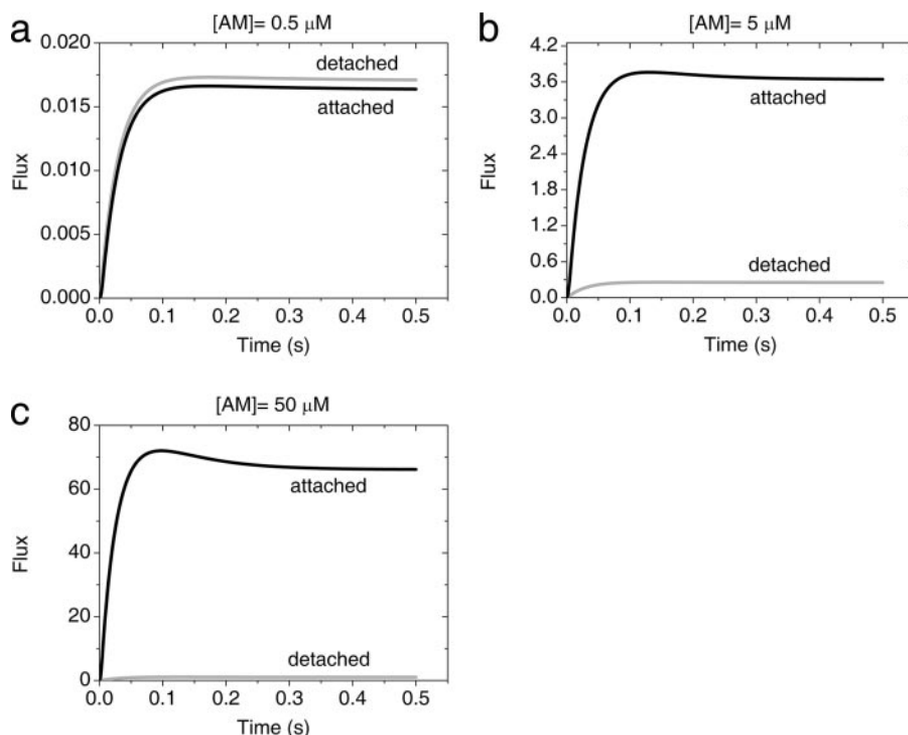


FIGURE 8. Simulation of the power stroke step at different actin concentrations. *a*, at low actin concentrations ($0.5 \mu\text{M}$) the isomerization reaction in the actin attached (*black*) and detached (*gray*) forms are practically equal. *b*, at moderately increased ($5 \mu\text{M}$) actin concentration $>90\%$ of the reaction flux occurs in the actin-bound state; this distribution is further pushed toward the actin attached form at higher ($50 \mu\text{M}$) actin concentration (*c*). Note the different scaling of the flux on the different panels.

TABLE 3
Kinetic parameters of the reaction sub-steps measured in this study

	Rate constants	Equilibrium/dissociation constants
$K_{\text{ATP-binding}}$ (55)		$120 \mu\text{M}^{-1}$ (8 nM)
k_{3a} (19)	350 s^{-1}	
k_{-3a} (19)	870 s^{-1}	
K_{3a} (19)		0.40 (2.5)
$K_{\text{hydrolysis}}$ (19)		82 (0.012)
$K_{\text{DP}} (K_{\text{DP}}\text{DP})$		0.024 mM^{-1} (42 μM)
$k_{\text{PD,on}}$	$1.6 \mu\text{M}^{-1}\text{s}^{-1}$	
$k_{\text{PD,off}}$	19 s^{-1}	
$K_{\text{PD}} (K_{\text{DP}}\text{PD})$		$0.084 \mu\text{M}^{-1}$ (12 μM)
$k_{\text{P,off}}$	260 s^{-1}	
$k_{\text{D,on}}$ (17)	$1.2 \mu\text{M}^{-1}\text{s}^{-1}$	
$k_{\text{D,off}}$ (17)	7 s^{-1}	
$K_{\text{D}} (K_{\text{D}}\text{D})$ (17)		$0.17 \mu\text{M}^{-1}$ (5.8 μM)
k_4	0.05 s^{-1}	
k_{-4}	0.15 s^{-1}	
K_4		0.33
$K_{\text{DP}} \text{A}\cdot\text{M}^*\text{ADP}\cdot\text{P}_i^a$		57
$K_{\text{DP}} \text{A}\cdot\text{M}^+\text{ADP}\cdot\text{P}_i^b$		0.7
$K_{\text{DP}} \text{A}\cdot\text{M}^+\text{ADP}$ (40)		0.12
$K_{\text{DP}} \text{A}\cdot\text{M}$ (40)		0.03

^a Based on W501+·ADP·AlF₄ pyrene-labeled actin binding experiments.

^b Calculated value from the thermodynamic box of Scheme 4.

reaction path is diverted to the actin-detached states during the recovery step, while the main flux flows through the actin-bound states in the case of the power stroke. Moreover, the weakly actin-bound $\text{A}\cdot\text{M}^*\text{ADP}\cdot\text{P}_i$ and $\text{A}\cdot\text{M}^+\text{ADP}\cdot\text{P}_i$ states represent higher free energy states compared with those in the detached states ($\text{M}^*\text{ADP}\cdot\text{P}_i$ and $\text{M}^+\text{ADP}\cdot\text{P}_i$, respectively), whose energy can also be utilized upon the power stroke of myosin.

Structural Aspects and Questions—Several questions arise from the present findings. First, is the myosin structure in the $\text{M}^*\text{ADP}\cdot\text{P}_i$ state the same as in the M^*ATP state? Some structural change must account for the fact that the up-down motion of the lever arm is four orders of magnitude slower for $\text{M}^*\text{ADP}\cdot\text{P}_i$ (0.05 s^{-1}) than for M^*ATP ($\sim 1000 \text{ s}^{-1}$) (Table 3). By *in silico* experiments the initial events of that structural change have been described recently. Combined quantum/classical molecular mechanics calculations of the ATP hydrolysis reaction path in myosin have shown that the nucleotide reorganizes first, in particular via a motion of the magnesium, which moves between the β and γ phosphate groups, thereby breaking its interaction with the switch 1 loop. This destabilizes the closed switch 1 conformation and prepares for its subsequent opening upon actin binding (47). Molecular dynamics simulations have shown that this is then followed by changes in the hydrogen bond network between the switch 2 loop, the relay helix, and the wedge loop (His-572 through Ala-574), leading to a locking of the wedge loop in the pre-power stroke configuration (48). Because the wedge loop motion is coupled to the lever arm swing (24), this locking can contribute to the much reduced rate of the reverse recovery step from the $\text{M}^*\text{ADP}\cdot\text{P}_i$ state.

Scheme 2 of the basal ATPase raises an intriguing question: Why is the reverse recovery step (k_4 , which is the rate-limiting step of the basal ATPase) needed before the products can be released (K_5 , K_6 , and K_7)? In other words, because switch 1 opening is likely to be needed to allow product release, why can't switch 1 open in the $\text{M}^*\text{ADP}\cdot\text{P}_i$ state of myosin? The reason is that when switch 1 opens, it is coupled to the closing of

the actin-binding cleft between the upper and lower 50-kDa domains (49). Although cleft closing can occur freely when the lever arm is "down" (*i.e.* in the pre-recovery step orientation, M^+), the same closing is not allowed when the lever arm is "up" in M^* . In other words, the actin-binding cleft and the lever arm are coupled (note that it is precisely this coupling that drives the power stroke when the cleft closing is induced by actin binding). In the absence of actin, this coupling prevents the closure of the actin-binding cleft, thus preventing the coupled opening of switch 1. Therefore, the pathway of basal ATPase involves first reversal of the recovery step ($\text{M}^*\text{ADP}\cdot\text{P}_i \rightarrow \text{M}^+\text{ADP}\cdot\text{P}_i$) to allow the combined switch 1 opening/cleft closing, which in turn allows the subsequent release of the products. In addition to switch 1 opening, the release of the γ -phosphate is likely to require also the opening of the switch 2 loop. Because that loop is strongly correlated to the lever arm orientation, another reason for the need of the reverse recovery step is to open switch 2 and allow γ -phosphate release.

When actin binds to $\text{M}^*\text{ADP}\cdot\text{P}_i$, actin induces closure of the actin-binding cleft and opening of the switch 1 loop (23, 50–54). Takagi *et al.* (14), suggested that the actin-induced opening of switch 1 weakens the interaction between switch 1 and 2, thus letting switch 2 open. This in turn would weaken the binding of the γ -phosphate and allow its release, leading to the increased ATPase turnover rate. But this explanation assumes that switch 2 can open independently from the motion of the lever arm. This is not likely, because, as long as the lever arm is in the pre-power stroke orientation, the wedge loop blocks the opening of switch 2, and the wedge loop is coupled to the lever arm orientation. Therefore, a more complete explanation for the actin-induced acceleration of the ATPase cycle is that actin binding also accelerates the lever arm reorientation (*i.e.* the power stroke), which in turn is coupled to switch 2 opening, thus facilitating the subsequent release of the products.

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