Binding modes of decavanadate to myosin and inhibition of the actomyosin ATPase activity

Teresa Tiago a,b,d,⁎, Paulo Martel a,c,1, Carlos Gutiérrez-Merino d, Manuel Aureliano a,b

a Departamento de Química e Bioquímica, FCT, Universidade do Algarve, Faro, Portugal
b CCMar, Universidade do Algarve, Faro, Portugal
CBME, Universidade do Algarve, Faro, Portugal
d Departamento de Bioquímica y Biología Molecular, Facultad de Ciencias, Universidad de Extremadura, Badajoz, Spain

Received 9 November 2006; received in revised form 15 January 2007; accepted 6 February 2007
Available online 20 February 2007

Abstract

Decavanadate, a vanadate oligomer, is known to interact with myosin and to inhibit the ATPase activity, but the putative binding sites and the mechanism of inhibition are still to be clarified. We have previously proposed that the decavanadate (V_{10}O_{28}^{6−}) inhibition of the actin-stimulated myosin ATPase activity is non-competitive towards both actin and ATP. A likely explanation for these results is that V_{10} binds to the so-called back-door at the end of the Pi-tube opposite to the nucleotide-binding site. In order to further investigate this possibility, we have carried out molecular docking simulations of the V_{10} oligomer on three different structures of the myosin motor domain of Dictyostelium discoideum, representing distinct states of the ATPase cycle. The results indicate a clear preference of V_{10} to bind at the back-door, but only on the "open" structures where there is access to the phosphate binding-loop. It is suggested that V_{10} acts as a "back-door stop" blocking the closure of the 50-kDa cleft necessary to carry out ATP-γ-phosphate hydrolysis. This provides a simple explanation to the non-competitive behavior of V_{10} and spurs the use of the oligomer as a tool to elucidate myosin back-door conformational changes in the process of muscle contraction.

© 2007 Elsevier B.V. All rights reserved.

Keywords: Back-door; Decavanadate; Myosin; Molecular Docking

1. Introduction

Myosin is a highly specialized protein involved in the process of muscle contraction, which along with actin converts the chemical energy of ATP hydrolysis into mechanical work. The motor function of myosin is located on its globular head, the subfragment-1 (S1), which contains both the nucleotide and actin binding sites [1]. The triphosphate portion of ATP is tightly bound in a tube-like structure, termed the Pi-tube, located at the bottom of the active site [2,3]. On the basis of high-resolution X-ray structures of the catalytic domain of Dictyostelium discoideum myosin (S1dC) complexed with several nucleotide analogues [4–9], it has been proposed the existence of a secondary entrance/exit to the active site at the rear opening of the Pi-tube opposite to the "front door". This "back-door" appears to be both a likely approach route for a water molecule to attack the γ-phosphate during ATP hydrolysis, and a likely exit route of the cleaved γ-phosphate after hydrolysis [10,11]. The γ-phosphate of ATP is partially visible in the structure of S1dC complexed with MgADP·BeFx or MgATP, in which there is a clear opening of the Pi-tube into the large cleft that divides the 50 kDa fragment into upper and lower domains. In contrast, in the S1dC structures complexed with MgADP·VO_{4} or MgADP·AlF_{4} the opening of the Pi-tube is blocked off by a closure of the 50 kDa cleft and the γ-phosphate is no longer visible (Fig. 1). The latter conformational change has been suggested to be necessary for the hydrolysis of ATP [12–14] meaning that the phosphate group will be trapped in the Pi-tube until a second conformational rearrangement occurs, which was proposed to occur upon myosin rebinding to actin [4]. Besides elucidating the kinetics of ATP hydrolysis, this "back door" mechanism has been used to explain a number of experimental observations, including the
formation of stable myosin-MgADP-vanadate complexes in muscle fibres [10].

The mechanism of inhibition of the myosin ATPase activity by vanadate is relatively well characterized. Monomeric vanadate ($VO_4^{3−}$) mimics the transition state for the γ-phosphate hydrolysis freezing myosin in a pre-power stroke state [15]. However, there is a considerably difference in the ability of vanadate to inhibit the myosin and the actomyosin ATPase activity. While in the absence of actin vanadate inhibits myosin ATPase activity in the submicromolar concentration range, in the presence of actin much higher concentrations (>900 μM) are needed [16,17]. Clearly, the need for such high concentrations strongly suggested that this effect could be due to oligomeric vanadate species that are favored at higher vanadate concentrations [18,19] and likely to occur through a different mechanism. In fact it was recently shown that the ATPase activity of the actomyosin complex is inhibited ($Ki=0.27±0.05$ μM) by decameric but not by the monomeric form of vanadate [20,21]. The results were consistent with binding of decavanadate (V$_{10}$) to the conserved regions of the phosphate binding-loop (P-loop) which shape part of the Pi-tube [22]. This $V_{10}$ high-affinity binding site produces non-competitive inhibition of the actin-stimulated S1-ATPase activity, without causing dissociation of the ATP-free rigor acto-S1 complex. Moreover, the affinity of S1 for $V_{10}$ is modulated by the conformational changes that takes place in S1 during the catalytic cycle, as indicated by the two- to three-fold increase of the dissociation constant produced in the presence of ADP·VO$_4$ and ADP·AlF$_4$ [21]. Two questions arose from this work: (1) how does $V_{10}$ access the P-loop? (2) How can the $V_{10}$ binding mode account for non-competitive inhibition of the ATPase activity of the actomyosin complex towards the nucleotide and actin? In order to address the previous questions, we performed a theoretical study in which decavanadate was docked in three different myosin motor domain conformations representing different intermediate states of the contractile cycle.

The source for these conformations were three X-ray structures of the $D. discoideum$ myosin S1 fragment, namely the apo (ligand-free) form, and the S1dC-MgATP and S1dC-MgADP·VO$_4$ complexes [5,8]. The first two conformations considered (free and ATP-bound) are quite similar and they both show an open back-door conformation characteristic of the pre-hydrolysis state. By binding at this site $V_{10}$ could block the Pi-tube and prevent ATP hydrolysis without competing with ATP or actin for association with myosin. The third conformation is an analogue of the transition state for ATP hydrolysis, and is expected to bind $V_{10}$ with lower affinity since the back-door is in a closed conformation, rendering the P-loop inaccessible to this ligand.

2. Methods

2.1. Selection and preparation of structures

Three $D. discoideum$ myosin S1 structures were retrieved from the Brookhaven Protein Databank: ligand-free (PDB code: 1FMV), in complex with MgATP (PDB code: 1FWM) and in complex with MgADP·VO$_4$ (PDB code: 1VOM) [5,8]. The program Needle (EMBL-EBI site) was used to calculate the percentage identity between the amino acid sequence of myosin II from $D. discoideum$ and from rabbit skeletal muscle using a global alignment procedure. Several short loops, missing from the x-ray structures, were built and optimized with the MODELLER software, following standard procedures [23]. All models were then validated with the PROCHECK software [24]. All crystallographic water molecules were removed prior to the docking procedure.

2.2. Decavanadate and protein parameters

Coordinates for the decavanadate ion ($V_{10}$) were retrieved from the Cambridge Structural Database. There are three non-equivalent positions for vanadium atoms in decavanadate: two VA atoms lie at the center of the equatorial plane, four VB atoms are in the equatorial plane and four VC atoms lie in an axial position. Partial charges for all atoms were assigned after SCF calculations of Kempf et al. [25]. These authors found that the inner vanadate oxygens should be close to ionic state, with a charge of $−2$, while all other oxygens would have charges close to $−1$. The remaining charge should be evenly distributed among the 10 vanadium atoms. For the purpose of this work, a charge of $−2$ was assigned to the OA and OB oxygen, a charge of $−1$ to all remaining oxygen atoms, and a charge of +2.4 to the 10 vanadium atoms. The non-bond parameters for vanadium were copied from the corresponding iron values in the AutoDock library, a reasonable assumption given the similarity between these two elements. Partial charges for the protein, MgATP and MgADP atoms were taken from the Amber 95 forcefield library [26]. For the VO$_4$ ion, the total charge of $−3$ was evenly distributed among the four oxygen atoms.
2.3. Molecular docking

All docking simulations were produced with the software AutoDock 3.0 [27], using Simulated Annealing algorithm (SA) and a total of 128 runs per docked complex. Since no torsional degrees of freedom were assigned to decavanadate, the SA docking algorithm was chosen because it performs better in rigid-body docking situations. For the protein, the lack of conformational flexibility is implied in the algorithm implemented in this version of the AutoDock program.

The termination criterion for docking runs was either a maximum number of 25,000 energy evaluations or a maximum of 27,000 generations. The docking trajectories were calculated within a cubic grid of size 36 Å and spacing 0.3 Å centered on the active site and the final solutions were clustered with a tolerance of 1.0 Å RMSD (root mean square deviation). Within each cluster, solutions are ranked according to docking energies from lowest to highest.

3. Results

The docking of decavanadate was performed on the structure of truncated D. discoideum myosin motor domain (S1dC) because the crystallographic structure of rabbit skeletal myosin has not been experimentally solved. Moreover, S1dC offers a greater number of nucleotide-bound complexes with high resolution compared to other myosins. At 43% sequence identity, the two molecules have a sufficient degree of structural similarity [28] to allow interpretation of the results here presented on the basis of our previously published experimental data on rabbit myosin.

The three myosin D. discoideum structures used in the present work were chosen as they represent three distinct states of the actomyosin ATP hydrolysis cycle relevant to the interpretation of our previously published data on the interaction between rabbit skeletal myosin S1 and decavanadate [21,22]. The docking region is limited to a cube with side 36 Å centered on the active site, since V10 is experimentally predicted to bind in the vicinity of the catalytic center through the phosphate-loop [22]. Docking calculations with the AutoDock software were run on the three crystal structures, and the results were evaluated in terms of the grouping (clustering) of solutions found in each case. The number of solutions found in each location (back- or front-door) and the lowest docking energy are indicative of the stability of the complex and are displayed in Table 1.

Table 1

<table>
<thead>
<tr>
<th></th>
<th>No. of solutions at “Back-door”</th>
<th>No. of solutions at “Front-door”</th>
<th>No. of solutions in top cluster*</th>
<th>Lowest energy (Kcal mol⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Apo (1FMV)</td>
<td>89</td>
<td>2</td>
<td>48</td>
<td>−3.20</td>
</tr>
<tr>
<td>MgATP (1FMW)</td>
<td>95</td>
<td>3</td>
<td>38</td>
<td>−2.02</td>
</tr>
<tr>
<td>MgADP·V10 (1VOM)</td>
<td>0</td>
<td>79</td>
<td>14</td>
<td>−1.95</td>
</tr>
</tbody>
</table>

Protein coordinates were taken from the Brookhaven Protein Databank; accession numbers are given in parenthesis. *Cluster with lower docking energies.

3.1. The apo-myosin structure

The crystal structure of S1dC myosin in its ligand-free (apo) form is believed to be representative of the pre-hydrolysis state after dissociation from actin [8]. The back-door is open in this structural state (Fig. 1a), allowing V10 to easily access the P-loop from this side. Of the 128 docking runs calculated for this structure, 89 ended with V10 docked on the back-door (Table 1 and Fig. 2a) and 79 of these conformations were the lowest
energy conformations, when compared to the remaining 49 runs. The lowest docking energy (in the back-door) was $-3.20$ kcal mol$^{-1}$ (this energy value is only indicative, due to the highly idiosyncratic nature of V$_{10}$ when compared to the test set used to parameterize Autodock). The preference thus shown by V$_{10}$ to bind at the back-door region leads support to our previous finding that V$_{10}$ does not compete with ATP for binding to myosin [21,22]. Also, this location is consistent with the non-competitive behavior of V$_{10}$ towards actin binding to myosin as it does not interfere with the actin binding interface.

A careful screening of the back-door solutions reveals another feature correlated to our previous experimental findings: back-door binding of V$_{10}$ brings it very close to Ser-181 (Fig. 3a), a residue located on the 178–185 sequence which forms the phosphate binding-loop (P-loop) motive, previously shown to be cleaved, upon irradiation, in the presence of V$_{10}$ [22]. In the “front-door”, besides being farther from Ser-181 (11.84 Å) than in the back-door (6.27 Å for the lowest docked energy conformation), the docked location of decavanadate is partially shielded by side chains of other residues which would prevent the access to Ser-181. One must note however that the use of a rigid protein model in the docking procedure makes these distances merely indicative.

3.2. The MgATP–myosin complex

The crystal structure of S1dC myosin complexed with ATP is also an open back-door structure, very similar to the apo-S1dC structure already discussed above. As pointed out by Bauer et al. [8], this could explain the high affinity of myosin for ATP, and is in agreement with this structure belonging to a pre-hydrolysis state, and thus unable to hydrolyze ATP. Given the extent of structural similarity, it is no surprise that the docking results are very similar to those obtained with the apo-S1dC structure. Of the 128 docking runs, 95 ended with V$_{10}$ docked in the back-door region, and 62 of these conformations were the solutions with lower energy (Table 1 and Fig. 2b). The lowest energy for a docked conformation was slightly higher than in the apo-S1dC docking ($-2.02$ kcal/mol), as expected because of the repulsion between the negative charge of V$_{10}$ and ATP (however, see the previous remark on the significance of Autodock energies). The preference for V$_{10}$ binding at the back-door is again clear, in spite of the electrostatic effect of ATP at the active site. This result agrees well with the experimentally observed non-competitive effect of V$_{10}$ towards myosin ATP hydrolysis [21]. The most stable conformation of V$_{10}$ is again close to Ser-181, but not as close as in the apo-S1dC dockings (Fig. 3b). This increased distance (approximately 2 Å for the lowest energy conformations) could explain why only partial V$_{10}$ photo-cleavage of myosin is attained in the presence of nucleotide [22].

3.3. The MgADP·VO$_4$–myosin complex

The S1dC–MgADP–VO$_4$ crystal structure is believed to be representative of the post-hydrolysis transient state before the power-stroke [5,29,30]. In this state, the 50-kDa cleft has closed (Fig. 1b), blocking the Pi-tube and preventing both the release of Pi and the access of V$_{10}$ to the back-door and consequently the P-loop. Accordingly, out of the 128 simulation runs there were 79 final V$_{10}$ conformations docked at the entrance of the ATP-binding site (“front-door”), and these were the lowest energy solutions (Table 1 and Fig. 2c). With the access to the P-loop barred, binding at the back-door region no longer represents an energetically favorable solution, and the front door remains as the only favorable solution for docking V$_{10}$ on the myosin surface. The smaller stabilization offered by binding at the nucleotide site could explain the experimentally observed two- to three-fold increase for the dissociation constant of the V$_{10}$–myosin complex in the presence of ADP·VO$_4$ or ADP·AlF$_4$ (another analogue of the transition.

Fig. 3. Lowest energy docked conformations of V$_{10}$ on: (a) the back-door (left) and nucleotide binding pocket (right) regions of the ligand-free structure of the S1 domain of Dictyostelium discoideum; (b) the S1dC structure in complex with Mg·ATP. Serine 181 is colored green in both structures; ATP is yellow and Mg$^{2+}$ violet. The P-loop, switch I and switch II segments forming the phosphate-tube are represented in orange, cyan and pink, respectively. V$_{10}$, Mg·ATP and Ser181 are represented as spheres.
state for ATP hydrolysis; S1dC-ADPAIF$_4$ structure is very similar to the ADP·VO$_4$ structure) [21]. Therefore, if different conformational states of myosin display different affinities for V$_{10}$, it is expected that an altered distribution of conformational states during the steady state catalytic activity takes place in the presence of V$_{10}$.

4. Discussion

The computational molecular docking calculations presented here, together with previous experimental data, lend strong support for the binding of V$_{10}$ at the back-door of the S1dC myosin fragment in the free or pre-hydrolysis nucleotide-bound state. This binding location both explains how the 10 Å decavanadate molecule gains access to the P-loop without having to squeeze through the narrow entrance of the phosphate-tube at the nucleotide binding site, and provides a simple mechanism for the experimentally observed non-competitive pattern of inhibition of V$_{10}$ towards binding of both ATP and actin to myosin. By binding at the back-door, at the end of the Pi-tube, V$_{10}$ will not interfere with the nucleotide binding site or the actin binding surface. Photo-cleavage of myosin by V$_{10}$ at the Ser-181 position can also be accounted for, since the lowest energy docked solutions of V$_{10}$ at the back-door place it in very close contact with this residue. Binding of V$_{10}$ to the back-door is also supported by the experimentally observed strong quenching of IAEDANS fluorescence by V$_{10}$ [21], when the probe is bond to a cysteine residue close to the back-door (Cys697). Also consistent with this picture is the previously observed competition of V$_{10}$ binding with AP$_5$A [21], a γ-phosphate modified nucleotide analogue that has been shown to bind myosin in a bimodal way that extends from the catalytic crevice through the Pi-tube and to the 50 kDa cleft leading to the back-door [31]. However, the question still remains: at which step in the actomyosin ATPase cycle does V$_{10}$ bind to myosin?

Since the closed states of myosin display low affinity for V$_{10}$ binding must take place when the back-door is open, either at the pre-hydrolysis or the “power-stroke” states of the actomyosin ATP hydrolysis cycle. In the latter state myosin is strongly bound to actin, so V$_{10}$ would have to interfere with formation of the transient acto-myosin complex. However, we have previously shown by light-scattering studies that V$_{10}$ binding does not dissociate the ATP-free rigor acto-myosin complex [21]. On the other hand, these experimental results indicated a positive correlation between the time after addition of ATP needed for the re-association of actin to myosin in the presence of V$_{10}$ and the extent of V$_{10}$ inhibition of the acto-myosin ATPase activity. Besides, decavanadate addition with MgADP is without effect on the actomyosin complex. Together, these results show that while V$_{10}$ does not promote dissociation of the acto-myosin complex, it does prevent its formation until ATP level in the solution is sufficiently low for the re-association process to occur. Consequently, V$_{10}$ must bind to myosin at its dissociated state, most likely the pre-hydrolysis state after actin release and before closure of the 50 kDa cleft and concomitantly

Fig. 4. Relevant steps of the actomyosin ATP hydrolysis cycle schematically represented as perturbed by V$_{10}$ binding: ATP binding to myosin in the rigor state (a) promotes opening of the 50-kDa cleft leading to dissociation of the actomyosin complex (b). In the absence of V$_{10}$, closure of the active site leads to ATP hydrolysis and a conformational change of the S1 domain (c). At this point, the back-door is closed and the hydrolyzed Pi group is trapped inside the active site until myosin rebinds actin, leading to the opening of the back-door and Pi release (d). When present, V$_{10}$ acts as “back-door stop”, blocking closure of the 50-kDa cleft and concomitant closure of the back-door thus preventing ATP hydrolysis.
the back-door. Decavanadate binding at this step would act as a “back-door stop” blocking the conformational change necessary to carry out the ATP-γ-phosphate hydrolysis. This proposal is schematically represented in Fig. 4. When attempting to extrapolate the myosin behavior in muscle fibers from the above proposal, one must keep in mind that all structures used in this work are static intermediates, frozen snapshots taken out of the conformational complexity of myosin in solution. As such, there may be other transient states, as of yet unknown, playing a key role in the myosin cycle in vivo.

Very often studies describing interactions of vanadate with proteins consider only monomeric vanadate as the active species, disregarding the possible contribution of other vanadate oligomers which are favored at higher vanadate concentrations [18,19]. Unlike the simple and labile vanadate oligomers including monomeric (V1), dimeric (V2), tetrameric (V4) and pentameric (V5) species, which interchange with each other on the millisecond to second time scale in neutral and alkaline aqueous solutions, decameric species (V10) interconversion is limited time periods because hydrolysis into other vanadate oligomers is very slow which makes it an excellent molecular probe to investigate the nature of polyoxoanions interactions with proteins. Among the free or complexed forms of vanadate, decameric species is one of the most potent inhibitors. The enzymes inhibited by decavanadate appear to be all prearranged to bind phosphoryl groups, but V10 is structurally different from V1 and cannot mimic orthophosphate in its interaction with enzymes. Moreover, decavanadate inhibition is by no means a general property of all the enzymes that bind phosphoryl groups and therefore the structural and conformational characteristics of the enzymes must be important. For example, Pezza et al. [33] have shown that the Walker A motive of ABC ATPases is a highly adapted anion-binding domain that can bind decavanadate with high affinity. Unlike V1, decavanadate interacts outside the Walker A loop (corresponding to the P-loop in myosin) and is stabilized by certain residues nearby. Therefore, according to these authors, if the presence of these residues is essential for the binding stabilization, only some members of the ABC superfamily are able to interact with decavanadate.

In what concerns myosin, vanadate is able to populate different conformational states of the myosin ATPase cycle depending on its oligomerization state. While monomeric vanadate (VO$_4^{3-}$) mimics the transition state for the γ-phosphate hydrolysis blocking myosin in a pre-power-stroke state, decameric vanadate induces the formation of the intermediate myosin-MgATP-V$_{10}$ complex blocking the actomyosin cycle presumably in the pre-hydrolysis state. Therefore, the mode of action of vanadate on the inhibition of the actin-stimulated myosin ATPase activity depends, at least in part, on the charge and size of the vanadate species, being favored for those with higher oligomerization state, such as the decameric species. The specific phosphate-binding domains in the vicinity of the back-door provide electrostatic interactions which favor an approach of these polyanionic species to the rear opening of the Pi-tube. In addition, the large size of decavanadate is likely to interfere with movements associated with closure of the 50–kDa cleft and concomitant closure of the back-door. This allows rationalizing on simple grounds (1) the non-competitive inhibition of V$_{10}$ towards both actin and ATP of the actin-stimulated myosin-ATPase activity; (2) the delayed re-association of the actomyosin complex but without a dissociation of the ATP-free rigor complex in the presence of decavanadate; (3) the preference of V$_{10}$ to bind at the back-door entrance of the Pi-tube, but only on the “open” structures were there is access to the phosphate binding-loop. Finally, this study lends further support to the myosin back-door mechanism hypothesis and spurs the use of decavanadate as a biochemical tool to gain a deeper knowledge of the myosin back-door modulation and the relevance of its conformational changes in the process of muscle contraction.

Acknowledgments

This work has been supported by Joint Spanish-Portuguese Grant HP2004-0080 (to C.G.-M. and M.A.), by POCTI program financed through FEDER for the research project 38191/QUI/2001 (to M.A.), and by Grant 3PR05A078 of the Junta de Extremadura (to C.G.-M.). Dr. T. Tiago is the recipient of a post-doctoral fellowship (SFRH/BPD/20777/2004) from the Portuguese Foundation for Science and Technology (FCT).

References


