

Asahikawa Medical College Repository http://amcor.asahikawa-med.ac.jp/

Journal of Biological Chemistry. (2008) 283(43): 29144-29155.

Roles of Tyr^<122>-hydrophobic Cluster and K⁺ Binding in Ca^{<2+>-releasing Process of ADP-insensitive Phosphoenzyme of Sarcoplasmic Reticulum Ca^{<2+>-ATPase.}}

Yamasaki K, Wang G, Daiho T, Danko S, Suzuki H.

Roles of Tyr¹²²-Hydrophobic Cluster and K⁺ Binding in Ca²⁺-releasing Process of ADP-insensitive Phosphoenzyme of Sarcoplasmic Reticulum Ca²⁺-ATPase^{*}

Kazuo Yamasaki, Guoli Wang, Takashi Daiho, Stefania Danko, and Hiroshi Suzuki

Department of Biochemistry, Asahikawa Medical College, Asahikawa 078-8510, Japan

Running Title: Tyr¹²²-Hydrophobic Cluster of SERCA1a for Ca²⁺ Release

Correspondence address to: Kazuo Yamasaki, Dept. of Biochemistry, Asahikawa Medical College,

Midorigaoka-Higashi, Asahikawa, 078-8510, Japan, Tel.: +81-166-68-2353; Fax: +81-166-68-2359;

E-mail: kyamasak@asahikawa-med.ac.jp.

Tyr¹²²-Hydrophobic cluster (Y122-HC) is an interaction network formed by the top part of second transmembrane helix and the cytoplasmic Actuator and Phosphorylation domains of sarcoplasmic reticulum Ca²⁺-ATPase. We have previously found that Y122-HC plays critical roles in the processing of **ADP-insensitive** phosphoenzyme (E2P) after its formation by the isomerization from ADP-sensitive phosphoenzyme (E1PCa₂) (Wang *et al.* (2005) J. Biol. Chem. 280, 26508-26516). Here, we further explored kinetic properties of the alanine-substitution mutants of Y122-HC to examine roles of Y122-HC for Ca²⁺release process in E2P. In the steady state, the amount of E2P decreased thereby that of $E1PCa_2$ increased with increasing lumenal Ca^{2+} concentration in the mutants with $K_{0.5}$ 110 ~ 320 μ M at pH 7.3. These lumenal Ca²⁺ affinities in E2P agreed with those estimated from the forward and lumenal Ca²⁺-induced reverse kinetics of the E1PCa₂-E2P isomerization. $K_{0.5}$ of the wild type was estimated to be 1.5 mM in this kinetics. Thus, E2P of the mutants possess significantly higher affinities for lumenal Ca²⁺ than that of the wild type. The kinetics further indicated that the rates of lumenal Ca^{2+} access and binding to the transport sites of E2P were substantially slowed by the mutations. Therefore the proper formation of Y122-HC and resulting compactly organized structure is critical for both decreasing Ca^{2+} affinity and opening the lumenal gate, thus for Ca²⁺ release from *E*2PCa₂. Interestingly, when K^+ was omitted from the medium of the wild type, the properties of the wild type became similar to those of Y122-HC mutants. K⁺ binding likely functions *via* producing the compactly organized structure, in this sense, similarly to Y122-HC.

Ca²⁺-ATPase Sarcoplasmic reticulum (SERCA1a¹) of the P-type ion transporting ATPase family catalyzes Ca^{2+} transport coupled with ATP hydrolysis from the cytoplasm to lumen against ~10000-fold concentration gradient (Refs. 1-8). In the initial steps (steps 1 and 2 in Scheme 1), the enzyme is activated by binding of two cytoplasmic Ca^{2+} ions at the transport sites with a submicromolar high affinity (E2 to $E1Ca_2$). The activated enzyme is then auto-phosphorylated at Asp³⁵¹ by ATP and forms a phosphoenzyme intermediate (EP) (step 3), thereby the bound Ca²⁺ ions are occluded in the transport sites. This EP is rapidly dephosphorylated by ADP in the reverse reaction reproducing ATP, therefore "ADPsensitive $E\hat{P}$ " (E1P). In the next step (step 4), E1PCa₂ is isomerized to the ADP-insensitive form, E2PCa₂. Upon this change at the catalytic site, the Ca^{2+} sites are deoccluded and opened to the lumenal side and the Ca^{2+} affinity is largely reduced, releasing the bound Ca^{2+} ions into the lumen (step 5). The Ca^{2+} release process is thought to be very rapid with wild type Ca^{2+} -ATPase, and the the accumulation of E2PCa2 intermediate had actually never been found until we recently identified and trapped successfully this intermediate by a mutation study (9). In the final step, the Asp^{351} -acylphosphate of E2P is hydrolyzed to reproduce the dephosphorylated and inactive E2 form (step 6). The transport cycle is totally reversible, e.g. E2P can be formed from E2 by P_i in the absence of Ca^{2+} , and the subsequent lumenal Ca²⁺ binding to E2P produces E1PCa₂.

Three-dimensional structures in several intermediate states and their analogs have been solved (10-18). The Ca²⁺-ATPase has three cytoplasmic domains; P (phosphorylation), N (nucleotide binding), and A (actuator or

anchor), and ten transmembrane helices (M1-M10). The two Ca^{2+} binding sites consist of residues on the M4, M5, M6, and M8 (10). The P domain possesses the phosphorylation site (Asp³⁵¹) and is directly linked to the long helices M4 and M5. The ATP binding site is on the N domain connected to the P domain. The A domain is linked to M1, M2, and M3 via the A/M1-, A/M2- and A/M3-linkers. The cytoplasmic three domains largely move and change their organization states during the Ca^{2+} -transport cycle (19-21), and these changes are linked with the rearrangements in the transmembrane helices for the Ca²⁺ transport. As a most remarkable change, in the EP isomerization (loss of ADP-sensitivity) and Ca²⁺ release, the A domain largely rotates and the P domain largely inclines toward the A domain and these domains produce their tight association (see Fig. 1 for the change $E1Ca_2 \cdot AlF_4 \cdot ADP \rightarrow E2 \cdot MgF_4^{2-}$ as the model for the overall process $E1 \sim PCa_2 \cdot ADP \rightarrow E2 \cdot P_i$ including the EP isomerization and Ca^{2+} release). These structural changes therefore involve distinct events in distinct regions, yet they are coordinated; namely 1) the loss of ADP-sensitivity at the cytoplasmic region, 2) the decrease in the Ca^{2+} -affinity at the transmembrane region, 3) the opening of the Ca^{2+} -releasing pathway (lumenal gating).

Recently, we found that mutations in a specific hydrophobic interaction network, "Tyr¹²²-hydrophobic cluster" (Y122-HC) at the A-P domain interface disrupt markedly the processing of ADP-insensitive EP formed from ATP with Ca^{2+} and also the hydrolysis of *E*2P formed from P_i without Ca^{2+} , thus causing nearly complete inhibition of the Ca²⁺-ATPase activity (22, 23). In these Y122-HC mutants, the high affinity binding of cytoplasmic Ca²⁺ the resulting E1PCa₂ formation, and the loss of the ADP-sensitivity were all found to occur normally as in the wild type (22, 23). Y122-HC is formed by gathering of the seven residues of the three regions upon their motions; *i.e.* the largely rotated A domain (Ile¹⁷⁹, Leu¹⁸⁰, and Ile²³²), the inclined P domain (Val⁷⁰⁵ and Val⁷²⁶), and the top part of the largely inclined M2 (or the A/M2-linker) (Leu¹¹⁹ and Tyr¹²²). Thus Y122-HC produces the compactly organized structure of E2P. Our previous analyses indicated that in the Y122-HC mutants, there is a kinetic limit after the loss of ADP-sensitivity and before the hydrolysis of the Ca^{2+} -free E2P, therefore the Ca^{2+} release from *E*2PCa₂ is likely retarded (22, 23). Almost the same kinetic results were found with the mutations in another A-P

domain interaction network at the Val²⁰⁰ loop of the A domain (24). Notably, $E2PCa_2$, the ADP-insensitive EP with two Ca²⁺ ions occluded at the transport sites was recently identified and trapped successfully by the elongation of the A/M1-linker with two or more amino-acid insertions (9). In the elongation mutants, Y122-HC is not formed properly yet in $E2PCa_2$ trapped, but it is properly formed in the Ca²⁺-released form of E2P produced by P_i without Ca²⁺. Thus the observation is consistent with the involvement of Y122-HC in the Ca²⁺-release process from $E2PCa_2$.

In this study, to further clarify roles of in the Ca^{2+} -deocclusion/release Y122-HC and thus in the long-range processes communication between the cytoplasmic and transmembrane regions, we explored kinetic features of the alanine-substitution mutants of Y122-HC. The results revealed that the mutations cause a marked increase in the apparent affinity of E2P for lumenal Ca^{2+} and also a substantial retardation of the lumenal Ca^{2+} access to E2P. Therefore, the formation of Y122-HC is critical for decreasing the affinity for Ca²⁺, for lumenal gating (opening of the release pathway), and thus for Ca^{2+} release into lumen. Importantly, the assembling manner of the seven residues in Y122-HC in the very recently revealed crystal structure $E2 \cdot BeF_3^{-1}$ (17, 18) somewhat differs from that in $E2 \cdot AlF_4$ and $E2 \cdot MgF_4^{2-}$. Therefore we discussed the significance of this difference in terms of the possible sequential gathering of the seven residues into Y122-HC on the basis of the observed difference in the extents of the mutational effects of the seven residues in Y122-HC. In addition, we found with the wild type that its kinetic behavior became similar to that of Y122-HC mutants when K^+ was omitted from the medium of the wild type. Results revealed for the first time the critical role of K^+ binding in the wild type for Ca^{2+} deocclusion/release from E2PCa₂.

EXPERIMENTAL PROCEDURES

Mutagenesis and Expression — Mutations were created by the QuikChangeTM sitedirected mutagenesis kit (Stratagene) and plasmid pGEM7-Zf(+) or pGEM3-Zf(+) (Promega) containing ApaI-KpnI or KpnI-SalI fragments of rabbit SERCA1a cDNA as a template. The ApaI-KpnI or KpnI-SalI fragments were then excised from the products and used to replace the corresponding region in the full-length SERCA1a cDNA in the pMT2 expression vector (25). The pMT2 DNA was transfected into COS-1 cells by the liposomemediated transfection method. Microsomes were prepared from the cells as described previously (26). The "control microsomes" were prepared from COS-1 cells transfected with the pMT2 vector containing no SERCA1a cDNA.

ATPase Activity — The rate of ATP hydrolysis was determined at 25 °C in a mixture containing 20 μ g/ml microsomal protein, 0.1 mM [γ -³²P]ATP, 3 μ M A23187, 0.1 M KCl, 7 mM MgCl₂, various concentrations of CaCl₂ up to 3 mM, 0.01 mM EGTA, and 50 mM MOPS/Tris (pH 7.3).

Formation and Hydrolysis of EP ----Phosphorylation of SERCA1a in microsomes with $[\gamma^{-32}P]ATP$ or ${}^{32}P_i$, and dephosphorylation of ³²P-labeled SERCA1a was performed under conditions described in the figure legends. The reactions were quenched with ice-cold trichloroacetic acid containing P_i. Rapid kinetics measurements of phosphorylation and dephosphorylation were performed with a handmade rapid mixing apparatus (27), otherwise the method was as above. The precipitated proteins were separated at pH 6.0 by 5% SDS-polyacrylamide gel electrophoresis, according to Weber and Osborn (28). The radioactivity associated with the separated Ca²⁺-ATPase was quantitated by described digital autoradiography as previously (29). The amount of EP formed with the expressed SERCA1a was obtained by subtracting the background radioactivity with the control microsomes. This background was less than 1% of the radioactivity of EP formed with the expressed wild-type SERCA1a.

Miscellaneous — Protein concentrations were determined by the method of Lowry *et al.* (30) with bovine serum albumin as the standard. Free Ca^{2+} concentrations were calculated by the Calcon program. Data were analyzed by nonlinear regression using the program Origin (Microcal Software, Inc., Northampton, MA). Three-dimensional models of the enzyme were reproduced by the program VMD (31).

RESULTS

 Ca^{2+} -induced Change in Accumulation of ADP-insensitive EP in the Presence of 0.1 M K^+ at Steady State — We first determined the steady state Ca^{2+} -ATPase activity in the presence of increasing Ca^{2+} and ionophore

A23187 with the alanine-substitution mutants of the seven residues of Y122-HC and the wild type. The Ca²⁺-ATPase activity was nearly completely inhibited in all the mutants in agreement with our previous observation (22, 23), and the complete inhibition was found at all the Ca^{2+} concentrations examined (see Supplemental Figure 1 for the representative mutant Y122A). Thus the possible lumenal Ca^{2+} effect was not revealed by this type of measurements. Therefore in Fig. 2, to assess the affinity of the lumenally oriented Ca²⁻ transport site of E2P (known as the low affinity sites with the mM \sim over 10 mM $K_{\rm d}$ value), the amounts of ADP-insensitive EP were determined with the representative mutant Y122A at steady state at various Ca² concentrations and pHs in the presence of A23187 and KCl. The total amounts of EP (ADP-sensitive *EP* plus ADP-insensitive *EP*) were nearly the same under all the sets of conditions.

In the mutant Y122A, the fraction of the ADP-insensitive EP was very high at the low Ca^{2+} concentrations at all pHs (Fig. 2). This agrees with the property of this mutant (22, 23) that the hydrolysis of *E*2P is nearly completely inhibited thus causing its accumulation. The fraction of ADP-insensitive EP in the mutant markedly decreased and it was converted to the ADP-sensitive EP with increasing Ca^{2+} concentration at several tens μM to sub-mM range. The apparent Ca²⁺ affinity in this Ca²⁺induced change increased with increasing pH and the Hill coefficients were found to be 2 in all pHs (see the legend to Fig. 2). In the wild type, the fraction of ADP-insensitive EP was low at pH 7.3 and 7.8 being approximately 10% or less, and was a significant level, 35% at pH 6.8 (Supplemental Figure 2). These levels were not changed at 1 μ M to 3 mM Ca²⁺. Consistently, the lumenal Ca^{2+} affinity of E2P of the wild type is known to be $mM \sim 10 mM$ range (see Ref. 32-34). The results suggested that the lumenal Ca²⁺ affinity of transport sites of E2P in the mutant may be significantly higher than that in the wild type.

Time Courses of Forward and Ca^{2+} -induced Reverse Conversions between ElPCa₂ and E2P — In Fig. 3 with Y122A, the ADPinsensitive EP and the ADP-sensitive EP was first accumulated at steady state at 10 μ M Ca²⁺ and 1 mM Ca²⁺, respectively at pH 7.3. Then the Ca²⁺ concentration jump was made from 10 μ M to 1 mM or from 1 mM to 80 nM, and the change in the fraction of the ADP-insensitive EP was followed. Because the hydrolysis of E2P was nearly completely blocked in Y122A (with the rate $\ll 0.01 \text{s}^{-1}$) (22, 23), the time courses represent the forward and reverse isomerization between $E1PCa_2$ and E2P. When Ca^{2+} was increased from 10 μ M to 1 mM, the fraction of ADP-insensitive EP rapidly decreased from 80% to 10% (*i.e.* it was converted to the ADP-sensitive EP) with a rate 0.4 s⁻¹. On the other hand, when the Ca²⁺ concentration was decreased from 1 mM to 80 nM, *i.e.* virtually Ca²⁺ was removed, the ADPsensitive EP was converted to the ADPinsensitive EP with a rate 0.022 s⁻¹.

Effect of Ca^{2+} Ionophore A23187 in Accumulation of ADP-insensitive EP — To ascertain that the Ca^{2+} dependent changes in the fraction of ADP-insensitive EP (Figs. 2 and 3) are caused by lumenal Ca^{2+} , we examined also in the absence of A23187 the Ca^{2+} dependence of the steady state fraction of ADP-insensitive EP (Fig. 4). The Ca^{2+} dependent change was rather small in the absence of A23187, in contrast to the very large change in its presence. Therefore, the observed Ca^{2+} -induced conversion from the ADP-insensitive EP to ADP-sensitive one in Y122A is due to the Ca^{2+} binding to the lumenally oriented transport sites.

 Ca^{2+} -induced Change in Accumulation of ADP-insensitive EP of Seven Y122-HC Mutants in the Presence of 0.1 $M K^+$ — In Fig. 5, each of the other six residues involved in the (Leu¹¹⁹/Ile¹⁷⁹/Leu¹⁸⁰/Ile²³²/Val⁷⁰⁵/ Y122-HC Val⁷²⁶) was substituted with alanine, and its effect on the ADP-insensitive EP level was examined as in Fig. 2 in the presence of A23187 and K⁺. All the Y122-HC mutants exhibited the marked Ca²⁺-dependent change in the ADP-insensitive EP fraction². The apparent affinities for lumenal Ca²⁺ in the Y122-HC mutants were found to be between 110 and 320 μ M with the Hill coefficients approximately 2 (Table 1). The results showed that all these Y122-HC mutants possess the lumenally oriented transport sites with the affinities as high as that of Y122A.

 Ca^{2+} -induced Change in Accumulation of ADP-insensitive EP of Mutants and Wild Type in the Absence of K^+ — In Fig. 6, the same sets of steady-state analysis as in Fig. 2 were done with the wild type and Y122A but here in the absence of K^+ . It is well known (35, 36) that in the absence of K^+ , the E2P hydrolysis of the wild type is markedly slowed and therefore the ADP-insensitive EP significantly accumulates. The fraction of ADP-insensitive EP in the wild type in the absence of K^+ decreased with increasing Ca^{2+} concentration as in Y122A with the Hill coefficient approximately 2. The

apparent affinity for lumenal Ca²⁺ increased with increasing pH in the wild type as in Y122A. The pH dependent changes are consistent with the fact that the residues for Ca^{2+} ligation at the transport sites are also involved in the proton binding (and its counter transport); the observed Ca^{2+} -induced changes reflect the Ca^{2+} binding to the lumenally oriented transport sites of E2P. At each pH, the affinity of the wild type was similar to or slightly lower than that of Y122A. Thus in the absence of 0.1 M K^+ , the property of the wild type became similar to that of Y122A. In Y122A, elimination of K^+ exhibited no significant effect on the apparent affinity for lumenal Ca^{2+} (*cf.* Fig. 2).

The observed effect of K^+ on the wild type is probably due to its binding at cytoplasmic region. In the crystallographic as well as mutational studies (12, 37), the K^+ binding site of the Ca²⁺-ATPase was identified to be in the cytoplasmic region but not in the lumenal or transmembrane regions (see Fig. 11). Actually, we found experimentally that when K^+ at 0.1 M was added without any K⁺-ionophore to the Ca²⁺-ATPase in SR vesicles phosphorylated in the absence of K^+ , the Ca²⁺-dependence of the ADP-insensitive EP fraction observed as in Fig. 6A became immediately (within 10 sec after the K^+ addition) that in the presence of 0.1 M K^+ as in Supplemental Figure 2 (data not shown).

Kinetics of Lumenal Ca²⁺-induced E2P to EIPCa₂ Reverse Transition Followed by Its ADP-induced Rapid Decay to $E1Ca_2$ in the Presence of 0.1 M K^+ — Then with the representative mutant Y122A, we explored kinetically the lumenal Ca²⁺ accessibility to the lumenally oriented transport sites of E2P formed from P_i without Ca^{2+} and the resulting lumenal Ca^{2+} -induced E2P to E1PCa₂ reverse transition. In Fig. 7, we included ADP and thereby followed the Ca²⁺- and ADP-induced decay of E2P to E1Ca₂ via E1PCa₂ in the reverse reaction. The E2P hydrolysis in the absence of Ca^{2+} was extremely slow (as previously demonstrated with the Y122-HC mutants (22, 23)), and the E2P decay was dramatically accelerated by the addition of Ca^{2+} and ADP (Fig. 7A). For example, the rate in the presence of 1 mM Ca^{2+} was 200-times faster than that of the forward E2P hydrolysis in the absence of Ca^{2+} . ADP alone without Ca^{2+} or Ca^{2+} alone without ADP did not accelerate the EP decay (data not shown). As shown in Fig. 3, the increase of Ca^{2+} to 1 mM converted the ADP-insensitive EP (E2P) to the ADP-sensitive one ($E1PCa_2$), and $E1PCa_2$ thus

formed was not decomposed in the absence of ADP. Therefore the Ca^{2+} and ADP-induced decay of E2P in Fig. 7A obviously occurred in the reverse reaction by the lumenal Ca² binding; $E2P + 2Ca^{2+} \rightarrow E1PCa_2$, then $E1PCa_2$ $+ ADP \rightarrow E1Ca_2 + ATP$ (Scheme 1). This view agrees with the previous demonstration with SR Ca²⁺-ATPase (34). The rate of the EP decay in the presence of Ca^{2+} and ADP increased almost linearly with increasing Ca²⁺ concentrations and was not saturated even at 3 mM (Fig. 7B)³. Here, note that the steady state level of ADP-sensitive EP ($E1PCa_2$) in the conversion from the ADP-insensitive EP (E2P) upon the lumenal Ca²⁺ binding in Y122A was almost fully saturated at 1 mM Ca²⁺ (see Fig. 2 at pH 7.3). The results indicate that the lumenal Ca^{2+} binding to *E*2P and formation of $E2PCa_2 (E2P + 2Ca^{2+} \rightarrow E2PCa_2)$ is likely the rate-limiting for the overall Ca²⁺- and ADPinduced reverse decay via the E2P to E1PCa₂ conversion with subsequent extremely rapid ADP-induced $E1PCa_2$ decay to $E1Ca_2$. This means that the slope in Fig. 7B in the Ca^{2+} dependence probably reflects the rate constant for the lumenal Ca²⁺ access and binding to the lumenally oriented transport sites of E2P and resulting E2PCa₂ formation.

The Ca²⁺- and ADP-dependent acceleration of the reverse *E*2P decay was assayed also with all the other Y122-HC mutants (Supplemental Figure 3). The rates of the reverse *E*2P decay increased almost linearly with increasing Ca²⁺ concentrations even at 3 mM, except those of I232A and V705A⁴ over ~1 mM Ca²⁺. Nevertheless, the slope of the Ca²⁺ dependence below 1 mM Ca²⁺ was estimated to be approximately 0.2 s⁻¹mM⁻¹ in all the mutants as in Y122A. Therefore, the rate of the lumenal Ca²⁺ access and binding to the transport sites is similar in all the mutants of Y122-HC.

Kinetics of Lumenal Ca^{2+} Access to E2P of Wild Type and Y122A with and without K^+ — Then in Fig. 8, with the wild type and the representative mutant Y122A in the presence and absence of 0.1 M K⁺, we analyzed the Ca^{2+} and ADP-dependent acceleration of the reverse decay of E2P formed from P_i without Ca^{2+} . As the well characterized property of the wild type, the forward hydrolysis of E2P without bound Ca^{2+} is very slow in the absence of K⁺, but markedly accelerated and thus very rapid in the presence of 0.1 M K⁺ (35, 36) (see the rates without Ca^{2+} in Fig. 8A). Nevertheless, even with the wild type in the presence of K⁺, we observed an apparently single exponential decay of E2P after the addition of Ca^{2+} and ADP at all the Ca^{2+} concentrations examined (time courses are not shown for simplicity). This is consistent with the kinetics described in the textbook by Fersht (39) that in the parallel reactions in which a compound undergoes two or more single-step reactions simultaneously, its disappearance rate is described by a single exponential decay. In our case, the two reactions are the forward *E*2P hydrolysis and the Ca^{2+} -/ADP-induced reverse *E*2P decay. The single decay rates thus obtained are plotted in Fig. 8*A*.

In the wild type in the presence of 0.1 M K^+ , the Ca^{2+} dependence of the EP decay rate was complicated because of the rapid E2P hydrolysis without Ca²⁺ (~0.4 s⁻¹), no change in the rate at $0\sim0.6$ mM Ca^{2+} , and the gradual increase above 0.6 mM. On the other hand, in the wild type in the absence of K^+ in which the E2P hydrolysis without bound Ca^{2+} is markedly slowed, the nearly linear increase in the rate of Ca^{2+} -/ADP-induced reverse E2P decay was observed at least up to $\sim 3 \text{ mM Ca}^{2+}$ as in the Y122-HC mutants in the presence of K^+ . The slope of the wild type without K^+ was actually close to that of Y122A with K^+ . Therefore, the rate of lumenal Ca²⁺ access and binding to the transport sites of E2P of the wild type in the absence of K^+ is similar to that of Y122A. With the wild type in the presence of K^+ , evaluation of the lumenal Ca^{2+1} access rate by this approach was not possible because of the complicated Ca²⁺-dependence curve. In Y122A, little effect was seen when K^+ was omitted at $0\sim1$ mM Ca²⁺, although the slope became gradually less steep at the higher Ca²⁺ concentration in the absence of K^+ .

In Fig. 8B, by using the rates of the E2Pdecay in the presence of added Ca²⁺ and ADP (determined in Fig. 8A) and the rates of the forward $E1PCa_2$ to E2Ptransition (as determined in Fig. 3), we simulated the fraction of the steady-state level of the ADPinsensitive EP(E2P) in the total amount of EPat each Ca²⁺ concentration. Note that this simulation was made possible by the fact that nearly all the phosphorylation sites are phosphorylated at steady state (in either E1P or E2P form) under the conditions used for the steady-state and kinetic analyses at all the Ca²⁺ concentrations in this study. Namely, the E2 to $E1Ca_2$ transition and the $E1PCa_2$ formation from $E1Ca_2$ with ATP are rapid enough to be ignored from the simulation. Therefore, the fraction of ADP-insensitive EP (E2P) in the steady state will be determined by the rate of its formation in the forward E1PCa₂ to E2P transition, v_1 , and by the rate of its decay, v_2 that includes both the forward hydrolysis of Ca^{2+} -unbound E2P to E2 and the Ca^{2+} -induced reverse transition to E1PCa₂ with the subsequent ADP-induced decay. This means that the simulation can be made even with the wild type in the presence of K^+ (as v_2 can include the forward E2P hydrolysis). In the steady-state conditions, the decay rate (v_2) and formation rate (v_1) should be equal, therefore the fraction of ADP-insensitive $EP(F_{E2P})$ in the total amount of EP will be estimated by an equation: $F_{E2P} = v_1/(v_1 + v_2)$. Here, the E2P decay rate (v_2) was obtained in Fig. 8A at each Ca^{2+} concentration. The $E1PCa_2$ to E2Ptransition rate (v_1) was estimated from the Ca²⁺ jump experiments from high (1 mM) to low (80 nM) for Y122A with and without 0.1 M K^+ and the wild type without K^+ , as described in Fig. 3. With the wild type in the presence of 0.1 M K^+ , the forward decay rate of E1PCa₂ formed by ATP was used as v_1 , because the *E*1PCa₂ to E2P transition (the loss of ADP-sensitivity) is rate-limiting for the E1PCa2 decay via E2P and its hydrolysis.

The Ca^{2+} -dependent curves thus obtained by the simulation for the steady-state level of ADP-insensitive EP for Y122A with and without K^+ and the wild type without K^+ agreed very well with the respective ones determined at the steady state (cf. Figs. 2 (with K^+) and 6 (without K^+) at pH 7.3). The affinities for lumenal Ca^{2+} estimated from the simulated curves are in fact almost the same as those actually determined at steady state (Table 2). The agreements assure the validity of the simulation and further allow us to estimate the lumenal Ca^{2+} affinity of E2P of the wild type in the presence of K^+ . In the simulation for the wild type in the presence of K^+ (open circles and inset in Fig. 8B), the fraction of ADP-insensitive EP was very low and the extent of its change was extremely small as expected from the steady-state measurements (cf. Supplemental Figure 2 (pH 7.3)). The apparent affinity of wild type for lumenal Ca^{2+} in the presence of K^+ was thus estimated by the small change to be 1.5 mM (see Table 2). This affinity was approximately 3.5-times lower than that of wild type without K^+ and 10-times lower than that of Y122A with and without K^+ . Thus by omitting K^+ , the lumenal Ca^{2+} affinity of *E*2P in the wild type became higher and similar to that in Y122A.

Kinetics of Lumenal Ca^{2+} -induced E2P to E1PCa₂ Reverse Transition of E2P of Wild Type in the Presence of 0.1 M K⁺ Was Revealed by the Absence of ADP — Unfortunately in the above experimental design and approach of Fig. 8A, we were not able to estimate the lumenal Ca²⁺ access rate in E2P of the wild type in the presence of K^+ because of the observed complexity of the Ca²⁺-dependent curve. In Fig. 9, we therefore employed a modified and thus different approach to examine the lumenal Ca²⁺-induced reverse conversion from E2P to E1PCa₂. Namely, E2P was formed with P_i and then a medium containing various concentrations of Ca^{2+} but without ADP (in contrast to its presence in Fig. 8) was added to E2P, and the subsequent EP decay was followed (Fig. 9A). In the absence of Ca^{2+} , E2P was all hydrolyzed rapidly to E2 in a single exponential function. The E2P hydrolysis was inhibited gradually with increasing Ca^{2+} concentrations (over 0.1 mM) and the decay time course became biphasic as typically seen with 1 mM Ca^{2+} . With increasing the Ca^{2+} concentration, the fraction of the first and rapid phase decreased, that of the second phase increased, and the rate of the second phase became further slower. The observation agrees with the previous kinetic analysis (9, 40). The first phase corresponds to the rapid and forward hydrolysis of the Ca^{2+} -unbound E2P to E2. The EP species in the second phase was all ADPsensitive (data not shown), therefore *E*1PCa₂ formed from *E*2P by the lumenal Ca^{2+} binding. E1PCa₂ decayed very slowly in the absence of ADP because the $E1PCa_2$ to E2P transition is much slower than the E2P hydrolysis and this transition is retarded by the Ca²⁺ replacement of Mg^{2+} at the catalytic site of E1PCa₂ at the ~mM high Ca^{2+} concentrations (41, 42).

The fraction of the second and slow phase of the EP decay was obtained by extrapolating to the zero time, and plotted versus the Ca²⁺ concentration (Fig. 9*B*). The plot showed saturation at $5\sim10$ mM Ca²⁺. Here it is critical to note that, as previously discussed in detail (9), the fraction of EP of the second phase (the fraction remaining after the first phase) is dependent on the ratio between the rates of the forward E2P hydrolysis and of the reverse E2P to $E1PCa_2$ conversion upon the lumenal Ca^{2+} binding to E2P. Namely, the plot in Fig. 9B reflects the relation between these forward and reverse rates of E2P rather than the lumenal Ca^{2+} affinity of E2P. For example, at the 50% saturation of the curve, the rate of the Ca²⁺induced $E1PCa_2$ formation from E2P is equal to that of the E2P hydrolysis to E2. Then in Fig. 9C for the wild type in the presence of K^+ , the rate of the E1PCa₂ formation from E2P by the lumenal Ca^{2+} binding to E2P was calculated $(k_{rev}, open circles)$ at each Ca^{2+} concentration

by using the fraction of the slow and second phase (F_s) and the *E*2P hydrolysis rate (k_h) with the equation; $k_{rev} = k_h F_s/(1 - F_s)$. The rate increased largely with increasing Ca²⁺ concentration.

In this kinetics, we eliminated the contribution of forward E2P hydrolysis on the overall E2P decay kinetics, and thereby revealed the rate of reverse E2P transition to $E1PCa_2$ induced by the lumenal Ca^{2+} binding of the wild type in the presence of K^+ . For comparison in Fig. 9C, the rates of the lumenal Ca^{2+} -induced reverse E2P decay estimated for the wild type without K^+ and Y122A with K^+ in Fig. 8A were replotted. Note again that in these cases, the hydrolysis of Ca²⁺-unbound E2P was very slow and retarded; therefore the observed Ca2+-/ADP-induced decay rates in their linear regions up to 3 mM Ca^{2+} reflect mostly the rates of the lumenal Ca²⁺ access and binding to E2P in the reverse E2P decay. Note also that the experimental design in Fig. 9A employed for the wild type with K⁺ was not applicable to the wild type without K^+ and Y122A, because the E2P hydrolysis is very slow and almost completely retarded in these cases and therefore the E2P decay upon the Ca^{2+} addition can not be described as the biphasic decay. Conversely, the experimental design employed in Fig. 8A to estimate the rates of the lumenal Ca²⁺ access was not applicable to the wild type in the presence of K^+ because of the complexity of the Ca²⁺dependent curve as described above in Fig. 8A.

Thus in Fig. 9C, employing the inevitably different but most suitable experimental designs depending on the different kinetic properties, we were able to compare the rates of the E2P to $E1PCa_2$ reverse transition induced by the lumenal Ca^{2+} binding to the transport sites of E2P at the limited Ca²⁺ concentration range up to 3 mM. In the wild type in the presence of K^+ , the rate was Ca^{24} dependent and not saturated even at 3 mM, thus reflecting at least the Ca²⁺-dependent and ratelimiting process; *i.e.* the lumenal Ca^{2+} -induced change from E2P to E2PCa₂. This reverse transition rate in the wild type in the presence of K⁺ was significantly faster than those in the wild type in the absence of K^+ and in Y122A (as well as in the other Y122-HC mutants (Supplemental Figure 3) especially at the high Ca^{2+} concentration over 1 mM.

Here it is also interesting to note that the affinity of *E*2P for the lumenal Ca^{2+} in the wild type without K⁺ and the Y122-HC mutants is significantly higher than in the wild type with K⁺ (see Fig. 8*B*). If the rate of lumenal Ca^{2+}

access and binding to *E*2P is solely slowed in the wild type without K^+ and Y122-HC mutants, a decrease in the affinity is rather the consequence, which is in contrast to the observed increase. Therefore the rates of the Ca^{2+} release from *E*2PCa₂ in the wild type without K^+ and in the Y122-HC mutants are also presumably retarded significantly as compared with that in the wild type in the presence of K^+ . Namely, the mutations of Y122-HC and the lack of K^+ binding affect the energy levels of Ca^{2+} -free and -bound *E*2P states and also that of the transition state for lumenal gating (opening), and favor the Ca^{2+} bound state *E*2PCa₂ and the closed lumenal gate.

DISCUSSION

Roles of Y122-HC in Ca^{2+} -release from $E2PCa_2$ and in E2P Hydrolvsis — In this study, we found that the mutations of any of the seven residues in Y122-HC increase the lumenal Ca²⁺ affinity and retard the lumenal Ca^{2+} access to the transport sites in *E*2P. These mutations also retard markedly the hydrolysis of the Ca^{2+} -released form of E2P (22, 23). Thus, the proper formation of Y122-HC from the seven residues is critical for both Ca²⁺ release into lumen from E2PCa₂ (reducing the Ca²⁺ affinity and opening the lumenal gate), and formation of the E2P catalytic site for the subsequent Asp³⁵¹-acylphosphate hydrolysis. The formation of Y122-HC therefore functions critically for realizing and stabilizing the compactly organized and thus distorted structure of the Ca^{2+} -released form of *E*2P. The stabilization of this state is certainly important for making the time period long enough for Ca²⁺ release into lumen and likely for proton bindings to the empty Ca²⁺ sites, and for the fine rearrangement of the catalytic site for the subsequent Asp³⁵¹-acylphosphate hydrolysis.

As shown in Fig. 10 and Supplemental Figure 4, the extents of the mutational effects on the lumenal Ca^{2+} affinities and on the *E*2P hydrolysis rates varied significantly among the seven Y122-HC mutants, and depended on their positions. The residues of which mutation exhibited strongest effects on increasing the lumenal Ca^{2+} affinity were Leu¹¹⁹, Tyr¹²², and Leu¹⁸⁰. This agrees with the critical role of M2 for rearrangement of the transmembrane helices for the Ca^{2+} -release; *i.e.* the tight association of the top part of M2 with the largely rotating A domain in Y122-HC functions for the lever-like inclination of M2 to push the lumenal part of M4 to open the

lumenal gate (14). In fact, in $E2PCa_2$ trapped by the elongation of the A/M1-linker, the Leu¹¹⁹/Tyr¹²² region on the top part of M2 is not involved fully in Y122-HC (9).

The residues of which mutations exhibited the strongest retardation of the E2P hydrolysis were Ile²³² at the top part of the A/M3-linker and, again, Leu¹¹⁹ and Tyr¹²² on the A/M2linker (top part of M2). Thus these residues on the linkers seem to contribute most critically to produce the proper configuration of the catalytic site. Consistently, the proteolytic cleavage at Leu¹¹⁹ on the A/M2-linker causes a marked inhibition of the E2P hydrolysis (43). The structural changes producing the Ca² release may be transmitted to the catalytic site via these residues of Y122-HC on the linkers, thereby ensuring the E2P hydrolysis to occur after the Ca²⁺ release. In any case, the different degree of the contributions of the seven residues of Y122-HC to the Ca^{2+} release and subsequent formation of the E2P catalytic site may suggest a possible sequential gathering of the seven residues. This possibility will be discussed more in the last section of "DISCUSSION" in relation to the crystal structure $E2 \cdot \text{BeF}_3$ (17, 18).

Structural Mechanism Involving Y122-HC and Other Critical Elements — In E1Ca₂·AIF₄ $\cdot ADP \rightarrow E2 \cdot MgF_4^{2-}$ as an overall structural change including the EP isomerization and Ca^{2+} release (Supplemental Figure 5A), the A domain largely rotates and M2 largely inclines. Also the P domain markedly inclines toward the underneath of the A domain, and rotates by approximately 20° around the phosphorylation site (Asp³⁵¹) parallel to the membrane and in the opposite direction of the A-domain rotation. These motions involve (can be dissected into) the horizontal and vertical factors, parallel and perpendicular to the membrane plane. As a consequence of the motions, the A and P domains and M2 will come to their appropriate positions producing their tight association at Y122-HC. At the A-P domain interface in the E2P analog structures, there is another interaction network between these domains at the Val²⁰⁰ loop, Asp¹⁹⁶-Asp²⁰³ of the A domain (Fig. 1, and see Supplemental Figure III in Ref. 23 for the details of the interactions and central role of Val²⁰⁰). Our previous mutations of Val²⁰⁰ showed (24) that this A-P domain interaction is critical for Ca2+ release from E2PCa₂ and for formation of the E2P catalytic site, thus very similarly to Y122-HC. Then note that in the E2P analog structures (see Supplemental Figure 5A for $E2 \cdot MgF_4^{-2}$), the two networks at Y122-HC and at Val²⁰⁰ are

located at each side of the A-P domain interface on its Top view and at the bottom and upper parts of the interface, respectively on its Side view. Thus the two are situated horizontally and vertically with the specific relative positioning. It is very likely that this positioning of the two is most efficiently functioning to realize and stabilize the compactly organized and distorted structure of the Ca²⁺-released E2P: *i.e.* the interactions at the two positions are most appropriate to produce the horizontal and vertical motions of the P and A domains and M2 required for Ca^{2} release from $E2PCa_2$, and to stabilize the Ca^{2+} released E2P state. Certainly these motions rearrangements cause the in the transmembrane helices for Ca^{2+} release: *e.g.* the P-domain inclination with slight rotation is directly associated with the bending and slight rotation of connected M4/M5 and downward movement of M4, thus their twisting-like motion. The largely inclining M2 pushes the lumenal part of M4 (Supplemental Figure 5B). Hence the Ca²⁺ sites are destroyed and lumenal gate is opened.

It should be noted that, for the loss of the ADP-sensitivity $E1PCa_2 \rightarrow E2PCa_2$, the large rotation of the A domain and its docking onto the P domain should occur so as to bring the T¹⁸¹GES loop above Asp³⁵¹-acylphosphate to block the ADP access from the N domain. As the motive force of this large A-domain rotation approximately parallel to membrane plane, the strain imposed on the A/M3-linker in $E1PCa_2$ was predicted to be critical (13, 14, 20, 44). Also, the sufficiently long length of the A/M1-linker was revealed to be critical for this EP isomerization, in this case, probably for realizing the E2PCa₂ structure, in which the A domain is positioned above the P domain (9, 45). For the subsequent Ca^{2+} release in *E2PCa*₂ \rightarrow E2P + 2Ca²⁺, the A/M1-linker with its appropriately short length (therefore its strain) is critical (9): Actually, the elongation of this blocks completely Ca^{2+} linker deocclusion/release from E2PCa₂, thus traps this E2PCa₂ state, in which Y122-HC is not properly formed yet in contrast to its proper formation in the Ca^{2+} -released form of E2Pwith the lumenally opened normal Ca^{2+} release pathway (9). The results clearly demonstrated that the native and appropriately short length of A/M1-linker functions critically in inducing the motions from the $E2PCa_2$ state, especially inclination of the A and P domains and M2, to accomplish the Y122-HC formation and the Ca²⁺-deocclusion/release from *E*2PCa₂. During the Y122-HC formation, the interaction-force being produced in Y122-HC will likely function to induce final process of the vertical and horizontal motions of the P and A domain and M2 to realize and stabilize the Ca^{2+} released E2P structure (Supplemental Figure 6). Importantly also, the E2P catalytic site is produced by these rearrangements. In this mechanism, a possible hydrolysis of Asp³⁵¹acylphosphate without releasing Ca^{2+} will be avoided; thereby the ordered reaction sequence of the Ca^{2+} release from $E2PCa_2$ and the subsequent E2Phydrolysis will be accomplished for the energy coupling.

Possible Structural Role of K^+ for Reducing Ca^{2+} Affinity and Lumenal Gating — K⁺ is known to markedly accelerate the E2P hydrolysis (35, 36), and also to modulate the E2 to $E1Ca_2$ transition in the nonphosphorylated Ca²⁺-ATPase (46, 47). In the present study, we further found that the K^+ binding is important for reducing the affinity for Ca^{2+} and lumenal gating thus for Ca^{2-} release from E2PCa₂. In the crystal structure $E1Ca_2 \cdot AIF_4 \cdot ADP$, K⁺ is situated at the bottom part of the P domain and coordinated by the backbone carbonyls of the loop Leu⁷¹¹-Glu⁷¹⁵ and by the Glu^{732} side chain (Fig. 11). The K⁺ binding at this site was indeed previously found by the mutations to be critical for the stimulation of the E2P hydrolysis (37). In the structures E2P analogs and E2(TG), this K⁺ site of the P domain comes very close to the A/M3-linker, and actually K^+ at this site is further coordinated by the Gln^{244} side chain on the A/M3-linker (see $E2 \cdot AlF_4$ in Fig. 11). Since the alanine-substitution of Gln²⁴⁴ and those of Glu-Gln-Asp²⁴⁵ gave virtually no effect on Ca^{2+} transport activity (48), K⁺ at this region may be coordinated by their neighboring residues or backbone carbonyls on the A/M3-linker and thereby play structural function. In the present study, we found that the lack of K^+ binding have the consequences very similar to those of the mutations at Y122-HC. It is therefore possible that the K^+ binding functions with similar structural effects as Y122-HC to produce the proper structure of the Ca^{2+} -released form of *E2P*.

Then note that the K⁺ site of the P domain in $E1Ca_2 \cdot AIF_4 \cdot ADP$ is situated at much higher position from the membrane plane than the Gln^{244} region on the A/M3-linker (Fig. 11) and that, in the change $E1Ca_2 \cdot AIF_4 \cdot ADP \rightarrow E2 \cdot AIF_4$ (or $E2 \cdot BeF_3$ and $E2 \cdot MgF_4^{2-}$), the P domain with connected M4/M5 largely inclines toward the A domain, hence the K⁺ site with bound K⁺ on the P domain moves down to the Gln^{244} region on the A/M3-linker

to contact with. The interactions between the bottom part of the P domain and the A/M3-linker *via* bound K^+ thus produced would likely cross-link them and hence contribute to formation and stabilization of this compactly organized Ca²⁺-released structure of *E*2P with the reduced Ca²⁺ affinity and lumenally opened gate. Alternatively, it is also possible that the appropriate P-domain structure produced by K⁺ binding on this domain solely contributes to the formation of the Ca²⁺-released *E*2P structure.

Y122-HC in Crystal Structure of $E2 \cdot BeF_3^-$ — The crystal structures of $E2 \cdot BeF_3$, the analog for the E2P ground state (21), were solved at atomic level very recently with and without bound thapsigargin, TG (E2BeF₃⁻ and $E2BeF_{3}(TG)$ (17, 18)). Surprisingly, in this crystallized $E2 \cdot BeF_3$, the side chains of Ile¹¹⁹ and Tyr¹²² are somewhat pointing away from the clustered other five residues on the A and P domains (Ile¹⁷⁹/Leu¹⁸⁰/Ile²³² and Val⁷⁰⁵/Val⁷²⁶), although all these seven residues are closely located in the $E2 \cdot BeF_3$ structures of both 2ZBE (17) and 3B9B (18). On the other hand, Y122-HC is formed fully from all these seven residues in $E2BeF_3$ (TG) as well as in the other E2P analogous structures, $E2 \cdot AlF_4$ and $E2 \cdot MgF_4^{2-}$. Thus, the assembling manner of the seven residues in the crystal structure $E2 \cdot BeF_3$ seemingly conflicts somehow with our results that the gathering of all the seven residues including Tyr^{122}/Ile^{119} in Y122-HC is required for producing the Ca²⁺-released *E2P*. Furthermore, Tyr^{122} and Ile^{119} on the top part of M2 (A/M2-linker) are likely most critical in Y122-HC and play central roles (Fig. 10). Our previous biochemical structural analysis of SR Ca²⁺-ATPase in solution by the proteolysis and the lumenal Ca^{2+} accessibility demonstrated (21) that in $E2 \cdot BeF_3^-$ without TG, Leu^{119}/Tyr^{122} are surely gathered and involved in Y122-HC, and thereby the lumenal gate is opened and the lumenal Ca^{2+} is accessible to the transport sites. Thus the crystal structure $E2 \cdot BeF_3^-$ seems to conflict also with these biochemical results obtained in solution.

Nevertheless, as a comprehensive idea, the crystal structure of $E2 \cdot BeF_3^-$ may be consistent with (or indicative of) the view that the gathering of the seven residues to form Y122-HC upon motions of the A and P domains and M2 (A/M2-linker) occurs in some ordered sequence but not necessarily at once (see Fig. 10 and its related "DISCUSSION"). Most peculiar to us is that Tyr¹²² and Leu¹¹⁹, of which mutations exhibited the most inhibitory effects, are not involved yet in the hydrophobic

cluster in the crystal structures $E2 \cdot BeF_3$. Then here note that in the structures $E2 \cdot BeF_3$, a Mg^{2+} ion is bound near the Ca^{2+} binding sites in the transmembrane domain as in extremely high Mg^{2+} concentration employed for crystallization (18) or protonation on the residues of transmembrane helices including Ca^{2+} ligands must have been occurred as in low pH for crystallization (17). Thus, these ligations are probably involved critically in the stabilization of the transmembrane helices for the crystallization. This might mean that the transmembrane structure thus stabilized differs from that without any ligations, *i.e.* the state immediate after the Ca^{2+} release (the empty Ca^{2+} sites) that is realized by the contribution of Y122-HC. Therefore, Y122-HC is, in return, disrupted or not properly produced yet in the crystal structure $E2 \cdot BeF_3^-$ as if it occurs with the lumenal Ca^{2+} binding in the E2PCa₂ state as postulated in this study. Therefore the following sequential gathering of the seven residues to produce Y122-HC can be speculated: The five hydrophobic residues on the A domain (Ile¹⁷⁹/Leu¹⁸⁰/Ile²³²) and P domain (Val⁷⁰⁵/Val⁷²⁶) are first gathered upon the motions of the A and P domains and top part of M2 (A/M2-linker), and subsequently, the top part of M2 including Ile¹¹⁹ and Tyr¹²² takes further motions during the final process

of the M2-inclination to join them and produce the fully assembled Y122-HC, thereby to realize and stabilize fully the gathered state of the A and P domains and top part of M2 (A/M2-linker) as in the Ca²⁺-released form of E2P. Being in agreement with this view, in $E2PCa_2$ trapped by the elongation of the A/M1-linker, Leu¹¹⁹/Tyr¹²² on the top part of M2 is not fully involved yet in Y122-HC (9). The Ca²⁺-released and empty Ca²⁺ sites (without any protonation and stabilization immediately after the Ca²⁺ release) will be subsequently protonated producing the E2P ground state for its hydrolysis.

Alternatively, if a possible contribution of such ligation in the transmembrane domain $(Mg^{2+} \text{ or protonation})$ should not be concerned in the crystallization of $E2 \cdot BeF_3$, the followings might be possible: First of all, the arrangements of helices of the Ca²⁺-released empty transport sites must be unstable, for example, due to possible repulsions between the negative charges of the Ca^{2+} ligands. Then to relieve the instability, the most effective gathering of Tyr^{122}/Leu^{119} in Y122-HC, which produces and stabilizes the Ca²⁺-released empty state, might possibly be disrupted, thereby the helices may be rearranged so as to form the more stabilized arrangements that can be crystallized.

Acknowledgments — We would like to thank Dr. David H. MacLennan, University of Toronto, for his generous gift of SERCA1a cDNA and Dr. Randal J. Kaufman, Genetics Institute, Cambridge, MA, for his generous gift of the expression vector pMT2. We are also grateful to Dr. Chikashi Toyoshima, University of Tokyo, for helpful discussions.

REFERENCES

- 1. Hasselbach, W., and Makinose, M. (1961) Biochem. Z. 333, 518-528
- 2. Ebashi, S., and Lipmann, F. (1962) J. Cell Biol. 14, 389-400
- 3. Inesi, G., Sumbilla, C., and Kirtley, M. E. (1990) Physiol. Rev. 70, 749-776
- 4. Møller, J. V., Juul, B., and le Maire, M. (1996) Biochim. Biophys. Acta 1286, 1-51
- 5. MacLennan, D. H., Rice, W. J., and Green, N. M. (1997) J. Biol. Chem. 272, 28815-28818
- 6. McIntosh, D. B. (1998) Adv. Mol. Cell Biol. 23A, 33-99
- 7. Toyoshima, C., and Inesi, G. (2004) Annu. Rev. Biochem. 73, 268-292
- 8. Toyoshima, C. (2008) Arch. Biochem. Biophys. doi 10.1016/j.abb.2008.04.017
- 9. Daiho, T., Yamasaki, K., Danko, S., and Suzuki, H. (2007) J. Biol. Chem. 282, 34429-34447
- 10. Toyoshima, C., Nakasako, M., Nomura, H., and Ogawa, H. (2000) Nature 405, 647-655
- 11. Toyoshima, C., and Nomura, H. (2002) Nature 418, 605-611
- 12. Sørensen, T. L-M., Møller, J. V., and Nissen, P. (2004) Science 304, 1672-1675
- 13. Toyoshima, C., and Mizutani, T. (2004) Nature 430, 529-535
- 14. Toyoshima, C., Nomura, H., and Tsuda, T. (2004) Nature 432, 361-368
- 15. Olesen, C., Sørensen, T. L.-M., Nielsen, R. C., Møller, J. V., and Nissen, P. (2004) Science 306, 2251-2255
- Takahashi, M., Kondou, Y., and Toyoshima, C. (2007) Proc. Natl. Acad. Sci. U. S. A. 104, 5800-5805

- 17. Toyoshima, C., Norimatsu, Y., Iwasawa, S., Tsuda, T., and Ogawa, H. (2007) Proc. Natl. Acad. Sci. U. S. A. 104, 19831-19836
- Olesen, C., Picard, M., Winther, A.-M. L., Gyrup, C., Morth, J. P., Oxvig, C., Møller, J. V., Nissen, P. (2007) *Nature* 450, 1036-1042.
- 19. Danko, S., Daiho, T., Yamasaki, K., Kamidochi, M., Suzuki, H., and Toyoshima, C. (2001) FEBS Lett. 489, 277–282
- 20. Danko, S., Yamasaki, K., Daiho, T., Suzuki, H., and Toyoshima, C. (2001) *FEBS Lett.* **505**, 129-135
- 21. Danko, S., Yamasaki, K., Daiho, T., and Suzuki, H. (2004) J. Biol. Chem. 279, 14991-14998
- 22. Yamasaki, K., Daiho, T., Danko, S., and Suzuki, H. (2004) J. Biol. Chem. 279, 2202-2210
- 23. Wang, G., Yamasaki, K., Daiho, T., and Suzuki, H. (2005) J. Biol. Chem. 280, 26508-26516
- 24. Kato, S., Kamidochi, M., Daiho, T., Yamasaki, K., Wang, G., and Suzuki, H. (2003) J. Biol. Chem. 278, 9624-9629
- 25. Kaufman, R. J., Davies, M. V., Pathak, V. K., and Hershey, J. W. B. (1989) Mol. Cell. Biol. 9, 946-958
- 26. Maruyama, K., and MacLennan, D. H. (1988) Proc. Natl. Acad. Sci. U. S. A. 85, 3314-3318
- 27. Kanazawa, T., Saito, M., and Tonomura, Y. (1970) J. Biochem. (Tokyo) 67, 693-711
- 28. Weber, K., and Osborn, M. (1969) J. Biol. Chem. 244, 4406-4412
- 29. Daiho, T., Suzuki, H., Yamasaki, K., Saino, T., and Kanazawa, T. (1999) FEBS Lett. 444, 54-58
- 30. Lowry, O. H., Rosebrough, N. J., Farr, A. L., and Randall, R. J. (1951) J. Biol. Chem. 193, 265-275
- 31. Humphrey, W., Dalke, A., and Schulten, K. (1996) J. Mol. Graphics 14, 33-38
- 32. Coan, C., Verjovski-Almeida, S., and Inesi, G. (1979) J. Biol. Chem. 254, 2968-2974
- 33. Prager, R., Punzengruber, C., Kolassa, N., Winkler, F., and Suko, J. (1979) Eur. J. Biochem. 97, 239-250
- 34. de Meis, L., and Inesi, G. (1982) J. Biol. Chem. 257, 1289-1294
- 35. Shigekawa, M., and Pearl, L. J. (1976) J. Biol. Chem. 251, 6947-6952
- 36. Shigekawa, M., and Dougherty, J. P. (1978) J. Biol. Chem. 253, 1451-1457
- 37. Sørensen, T. L-M., Clausen, J. D., Jensen, A.-M. L., Vilsen, V., Møller, J. V., Andersen, J. P., and Nissen, P. (2004) *J. Biol. Chem.* **279**, 46355-46358
- 38. de Meis, L., Martins, O. B., and Alves, E. W. (1980) Biochemistry 19, 4252-4261
- 39. Fersht, A. R. (1999) Structure and Mechanism in Protein Science: A Guide to Enzyme Catalysis and Protein Folding, pp. 132-168, W. H. Freeman and Co., New York
- 40. Sato, K., Yamasaki, K., Daiho, T., Miyauchi, Y., Takahashi, H., Ishida-Yamamoto, A., Nakamura, S., Iizuka, H., and Suzuki, H. (2004) *J. Biol. Chem.* **279**, 35595-35603
- 41. Shigekawa, M., Wakabayashi, S., and Nakamura, H. (1983) J. Biol. Chem. 258, 8698-8707
- 42. Wakabayashi, S., and Shigekawa, M. (1987) J. Biol. Chem. 262, 11524-11531
- 43. Lenoir, G., Picard, M., Gauron, C., Montigny, C., Le Maréchal, P., Falson, P., le Maire, M., Møller, J.V., and Champeil, P. (2004) *J. Biol. Chem.* **279**, 9156-9166
- 44. Møller, J.V., Lenoir, G., Marchand, C., Montigny, C., le Maire, M., Toyoshima, C., Juul, B. S., and Champeil, P. (2002) *J. Biol. Chem.* **277**, 38647-38659
- 45. Daiho, T., Yamasaki, K., Wang, G., Danko, S., and Suzuki, H. (2003) J. Biol. Chem. 278, 39197-39204
- 46. Lee, A. G., Baker, K., Khan, Y. M., and East, J. M. (1995) Biochem. J. 305, 225-231
- 47. Champeil, P., Henao, F., and de Foresta, B. (1997) Biochemistry 36, 12383-12393
- 48. Clarke, D. M., Maruyama, K., Loo, T. W., Leberer, E., Inesi, G., and MacLennan, D. H. (1989) *J. Biol. Chem.* **264**, 11246-11251

FOOTNOTES

*This work was supported by a Grant-in-Aid for Scientific Research (C) (to K.Y.) and (B) (to H.S.) from the Ministry of Education, Culture, Sports, Science and Technology of Japan.

¹The abbreviations used are: SERCA1a, adult fast-twitch skeletal muscle sarcoplasmic reticulum Ca^{2+} -ATPase; *E*P, phosphoenzyme; *E*1P, ADP-sensitive phosphoenzyme; *E*2P, ADP-insensitive phosphoenzyme; MOPS, 3-(*N*-morpholino) propanesulfonic acid; TG, thapsigargin; Y122-HC, Tyr¹²²-hydrophobic cluster.

²Maximal level of the ADP-insensitive *E*P of L180A at the low Ca^{2+} concentrations (44% of total amount of *E*P) was significantly lower than those of the other mutants. It may be due to the significantly faster *E*2P hydrolysis rate in L180A as compared with the rates in the others, as shown in Supplemental Figure 4.

³Its slope was approximately 0.2 s⁻¹mM⁻¹ at 0~2 mM Ca²⁺. In Fig. 3, the rate of the forward *E*1PCa₂ to *E*2P conversion was estimated to be 0.02 s⁻¹. Therefore the calculation with these values, 0.02 s⁻¹ divided by 0.2 s⁻¹mM⁻¹, gave the apparent affinity for lumenal Ca²⁺ in *E*2P of Y122A as 100 μ M. This value agreed very well with that (160 μ M) obtained in Fig. 2 under the same conditions (pH 7.3) at the steady state.

⁴As a possible reason for the less steep Ca²⁺-dependent curve observed with I232A and V705A over ~ 1 mM, it might be possible that the rate in the transition from *E*2PCa₂ to *E*1PCa₂ is slower in these mutants than in the other mutants, and this step became a rate-limiting at the high Ca²⁺ concentrations where the lumenal Ca²⁺-induced change to *E*2PCa₂ became fast.

FIGURE LEGENDS

FIGURE 1. Structure of SERCA1a and formation of Tyr¹²²-hydrophobic cluster. The coordinates for the structures $E1Ca_2 \cdot AlF_4 \cdot ADP$, (the analog for the transition state of the phosphoryl transfer $E1 \sim PCa_2 \cdot ADP$, *left panel*) and $E2 \cdot MgF_4^{2-}$ ($E2 \cdot P_i$ analog (21), *right panel*) of Ca^{2+} -ATPase were obtained from the Protein Data Bank (PDB accession code 1T5T and 1WPG, respectively (12, 14)). The arrows indicate approximate movements of the A and P domain and the top part of M2 (Leu¹¹⁹/Tyr¹²²) in the change from $E1 \cdot AlF_4 \cdot ADP$ to $E2 \cdot MgF_4^{2-}$. The seven hydrophobic residues, Ile¹⁷⁹/Leu¹⁸⁰/Ile²³² on the A domain, Leu¹¹⁹/Tyr¹²² on the A/M2-linker, Val⁷⁰⁵/Val⁷²⁶ on the P domain are depicted as van der Waals spheres. They gather to form a hydrophobic cluster around Tyr¹²² in the change $E1Ca_2 \cdot AlF_4 \cdot ADP \rightarrow E2 \cdot MgF_4^{2-}$ (Y122-HC, surrounded by *red dotted circle*). The top part of M2 including Leu¹¹⁹/Tyr¹²² is unwound in $E2 \cdot MgF_4^{2-}$, $E2 \cdot AlF_4$, and $E2 \cdot BeF_3$ with bound thapsigargin (TG), thus becomes the A/M2-linker loop (see the region of M2 colored by *pink*).

FIGURE 2. Ca^{2+} dependence of accumulation of ADP-insensitive *EP* in the steady state in mutant Y122A. Microsomes expressing the mutant Y122A were phosphorylated with $[\gamma^{-32}P]ATP$ at various Ca^{2+} concentrations and pHs as indicated at 0 °C for 5 min in a mixture containing 20 µg/ml microsomal protein, 50 mM MOPS/Tris, 0.1 M KCl, 7 mM MgCl₂, 0.01 mM EGTA, 3 µM A23187, 10 µM $[\gamma^{-32}P]ATP$, and various concentrations of CaCl₂. For the determination of the accumulated ADP-insensitive *EP*, an equal volume (50 µl) of a mixture containing 10 mM ADP, 7 mM MgCl₂, 10 mM EGTA, 50 mM MOPS/Tris (pH 6.8, 7.3, or 7.8 as indicated), and 0.1 M KCl was added to the above phosphorylation mixture. At 1 s after this addition, the reaction was quenched with trichloroacetic acid. ADP-sensitive *EP* disappeared entirely within 1 s after the addition of ADP. The total amounts of *EP* were nearly the same under all the conditions (data not shown). The amount of ADP-insensitive *EP* is shown as a percentage of the total amount of *EP*. *Solid lines* show the least squares fit to the Hill equation. Apparent Ca²⁺ affinities and Hill coefficients thus obtained were 360 µM and 2.1 (pH 6.8), 160 µM and 1.9 (pH 7.3), and 76 µM and 2.0 (pH 7.8).

FIGURE 3. Time course of the change in the fraction of ADP-insensitive *EP* upon Ca²⁺concentration jump. Microsomes expressing the mutant Y122A (20 μ g/ml) were phosphorylated with [γ -³²P]ATP at pH 7.3 for 5 min in the presence of 10 μ M (\circ) or 1 mM CaCl₂ (\bullet) without EGTA in the phosphorylation solution otherwise as described in Fig. 2. Then an equal volume of the solution containing 2 mM CaCl₂ or 2 mM EGTA (otherwise as in the phosphorylation solution) was added to give final free Ca²⁺ concentrations 1 mM and 80 nM, respectively. At the indicated times after this Ca²⁺ jump, the total amount of *EP* and the amount of ADP-insensitive *EP* was determined as in Fig. 2. *Solid lines* show the least squares fit to a single exponential. The total amount of *EP* was not changed during the period of observation in both cases (data not shown). The amount of ADP-insensitive *EP* is shown as a percentage of the total amount of *EP*. FIGURE 4. Effect of calcium ionophore on the Ca²⁺ dependence of ADP-insensitive *EP* accumulation in the steady state. Microsomes expressing the mutant Y122A (20 μ g/ml) were phosphorylated at 0 °C for 5 min in the medium containing 50 mM MOPS/Tris (pH 7.3), 0.1 M KCl, 7 mM MgCl₂, 10 μ M CaCl₂, 10 μ M [γ -³²P]ATP, and 50 μ M Ruthenium Red (to block Ca²⁺ channels as much as possible) in the presence (\circ) or absence (\bullet) of 3 μ M A23187. Then the free Ca²⁺ concentration was changed as indicated on abscissa by mixing with an equal volume of a solution containing 50 mM MOPS/Tris (pH 7.3), 0.1 M KCl, 7 mM MgCl₂, 2 mM EGTA, and various concentrations of CaCl₂. After 10 s of this Ca²⁺ jump, the total amount of *EP* and the fraction of the ADP-insensitive *EP* were determined as described in Fig. 2. The inclusion of Ruthenium Red in the presence of A23187 caused a slight decrease in the ADP-insensitive *EP* fraction at the Ca²⁺ concentrations below ~100 μ M (*cf.* Fig. 2) for unknown reasons.

FIGURE 5. Ca^{2+} dependence of accumulation of ADP-insensitive *EP* in mutants for Tyr¹²hydrophobic cluster. Microsomes expressing each of the seven Y122-HC mutants (indicated in the figure) were phosphorylated with [γ -³²P]ATP at various concentrations of Ca²⁺ and pH 7.3, otherwise as described in Fig. 2. *Solid lines* show the least squares fit to the Hill equation. The fitting parameters including the apparent Ca²⁺ affinity and the Hill coefficient thus obtained were listed in Table 1.

FIGURE 6. Ca^{2+} dependence of accumulation of ADP-insensitive *EP* in the absence of K⁺. Microsomes expressing the wild type (*A*) or Y122A (*B*) SERCA1a were phosphorylated with [γ -³²P]ATP at various Ca²⁺ concentrations and pH (6.8 (\circ), 7.3 (\bullet), and 7.8 (Δ)) in the presence of 0.1 M LiCl in place of KCl otherwise under the exactly same conditions as those described in Fig. 2. The amount of ADP-insensitive *EP* was determined by the addition of the ADP solution that contains 0.1 M LiCl in place of KCl otherwise as in Fig. 2. The total amount of *EP* was nearly constant under all the conditions (data not shown). The amount of ADP-insensitive *EP* is shown as a percentage of the total amount of *EP*. *Solid lines* show the least squares fit to the Hill equation. Apparent Ca²⁺ affinities and Hill coefficients thus obtained with the wild type (*panel A*) were 930 μ M and 1.8 (pH 6.8), 400 μ M and 1.5 (pH 7.3), and 310 μ M and 1.9 (pH 7.8), and those obtained with Y122A (*panel B*) were 1000 μ M and 1.9 (pH 6.8), 220 μ M and 2.2 (pH 7.3), and 130 μ M and 1.7 (pH 7.8).

FIGURE 7. Lumenal Ca²⁺ and ADP-induced reverse *E*2P decay of Y122A in the presence of K⁺. *A*, microsomes (100 μ g/ml) expressing the mutant Y122A were phosphorylated with ³²P_i for 10 min at room temperature in the absence of Ca²⁺ in a medium containing 50 mM MOPS/Tris (pH 7.3), 7 mM MgCl₂, 1 mM EGTA, 15 μ M A23187, 0.1 mM ³²P_i, and 20% Me₂SO (that favors extremely the *E*2P formation (38)), and then the reaction mixture was chilled on ice. Subsequently, the phosphorylated sample was diluted at 0 °C with a 20-fold volume of a chase solution containing 50 mM MOPS/Tris (pH 7.3), 105 mM KCl, 7 mM MgCl₂, 1 mM EGTA, 0.1mM non-radioactive P_i and 0.105 mM ADP without or with various concentrations of CaCl₂ to give the final free Ca²⁺ concentrations as indicated. At the indicated periods, the chase reaction was terminated by trichloroacetic acid and the amount of *E*P was determined. *Solid lines* show the least squares fit to a single exponential decay. The decay rates thus obtained were plotted *versus* the Ca²⁺ concentration in *panel B*.

FIGURE 8. Lumenal Ca²⁺ and ADP-induced reverse *E*2P decay of wild type and Y122A in the absence and presence of K⁺. *A*, microsomes (100 μ g/ml) expressing wild type (WT) or Y122A were phosphorylated with ³²P_i in the presence of A23187 and absence of Ca²⁺, and then chilled on ice, as described in Fig. 7. Subsequently, the phosphorylated sample was diluted at 0 °C with a 20-fold volume of a chase solution containing non-radioactive P_i, various concentrations of CaCl₂, and ADP in the presence of 105 mM KCl (*open symbols*) or 105 mM LiCl in place of KCl (*closed symbols*), otherwise as described in Fig. 7. The time courses of *E*P decay were fitted to single exponential (data not shown, see Fig. 7*A* as an example). The rates thus obtained were plotted *versus* the Ca²⁺ concentrations. *B*, the fraction ADP-insensitive *E*P (F_{*E*2P}) in the total amount of *E*P at steady state was simulated by using the rate of *E*2P decay (v₂) and the rate of the *E*1PCa₂ to *E*2P isomerization (loss of ADP-sensitivity, v₁) with an equation $F_{E2P} = v_1/(v_1 + v_2)$. Here, v₂ is the *E*2P decay rate obtained above in *panel A* at each Ca²⁺ concentrations. The v₁ was obtained as described in Fig. 3 by the Ca²⁺ (without K⁺) and for the wild type with 0.1 M Li⁺ (without K⁺). For the wild type with 0.1 M K⁺, the forward decay rate of *E*1PCa₂ formed from ATP was used as the v₁ value, because the *E*1PCa₂ to *E*2P transition (the loss of ADP-sensitivity) is rate-limiting for the *E*1PCa₂ decay *via E*2P and its

subsequent rapid hydrolysis. The v₁ values actually used for the calculation were 0.049 s⁻¹ (wild type with K⁺), 0.071 s⁻¹ (wild type with Li⁺), 0.021 s⁻¹ (Y122A with K⁺), and 0.034s⁻¹ (Y122A with Li⁺). The fraction of ADP-insensitive *E*P thus calculated was plotted *versus* the Ca²⁺ concentrations. The solid lines show the least squares fit to the Hill equation. In the *inset*, the ordinate is in a magnified scale for wild type with K⁺. In Table 2, the affinities and the Hill coefficients of the wild type and the Y122-HC mutants thus "*Estimated by kinetic analyses*" in the absence and presence of K⁺ at pH 7.3 are summarized together with those actually "*Determined by the steady-state analyses*" of the lumenal Ca²⁺-induced change of the fraction of ADP-insensitive *E*P under otherwise the same conditions in Figs. 2 and 6.

FIGURE 9. Kinetics of lumenal Ca^{2+} -induced change of E2P to E1PCa₂ in the wild type in the presence of K^+ without ADP. A, the microsomes expressing the wild type were phosphorylated with ${}^{32}P_i$ in the presence of A23187 and absence of Ca^{2+} and chilled on ice, as described in Fig. 7. Subsequently, the phosphorylated sample was mixed at 0 °C with a 20-fold volume of chase solution containing 105 mM KCl and various concentrations of CaCl₂ without ADP, otherwise as described in Fig. 7. The final free Ca^{2+} concentrations were indicated in the figure. At the indicated time periods after this addition, the chase reaction was terminated by trichloroacetic acid and the amount of EP was determined. Solid lines show the least squares fit to a double exponential decay. The EP remaining in the second and slow phase was all in the ADP-sensitive form (thus $E1PCa_2$, data not shown), and the first and rapid phase is the forward hydrolysis of E2P without bound Ca^{2+} . B, the fraction of EP in the second phase in the total amount of EP was obtained by extrapolating to the zero time in the double exponential decay fitting, and plotted versus the Ca²⁺ concentration. The data were fitted well with the Hill equation (solid line), and the Ca²⁺ concentration giving the 50% saturation and the Hill coefficient were found to be 750 μ M and 1.5. Here note that the EP amount in the second phase is dependent on the ratio between the rate of the forward E2P hydrolysis and that of the reverse E2P to E1PCa₂ conversion upon the lumenal Ca^{2+} binding to E2P: The plot reflects the relative values between these forward and reverse rates of E2P rather than the lumenal Ca^{2+} affinity of E2P. C, by using the data obtained in A and B with the wild type in the presence of K⁺, the rate of the lumenal Ca²⁺-induced E1PCa₂ formation from E2P (k_{rev} (\circ)) was calculated at each Ca²⁺ concentration by the equation; $k_{rev} =$ $k_{\rm h}F_{\rm s}/(1 - F_{\rm s})$. Here, $F_{\rm s}$ is the fraction of EP in the second phase, and $k_{\rm h}$ is the forward hydrolysis rate of E2P without Ca^{2+} . For comparison with the wild type in the absence of K⁺ (•) and Y122A in the presence of K⁺ (Δ), their lumenal Ca²⁺ access rates (the rates of the lumenal Ca²⁺-induced reverse E2P decay via E1PCa₂ in the presence of ADP) obtained in Fig. 8A were plotted.

FIGURE 10. Strength of the mutational effects of seven residues in Tyr¹²-hydrophobic cluster on *E2P* hydrolysis and lumenal Ca²⁺ affinity. The detailed structure at Y122-HC is shown with $E2 \cdot AlF_4^-$ (the analog for $E2 \sim P$, the transition state of the *E2P* hydrolysis (21), PDB code: 1XP5 (15)). The seven residues involved in Y122-HC (Tyr¹²²/Leu¹¹⁹, Ile¹⁷⁹/Leu¹⁸⁰, Ile²³², and Val⁷⁰⁵/Val⁷²⁶), AlF₄⁻ bound at the phosphorylation site Asp³⁵¹, and the bound potassium ion are shown by van der Waals spheres. The seven residues in Y122-HC are colored differently based on the strength of the retardation of the *E2P* hydrolysis rate (*lower panel*) and that of the increase in the lumenal Ca²⁺ affinity (*upper panel*). The color changes gradually from *red* for strongest effects to *blue* as becoming weaker.

FIGURE 11. **Bound K⁺ and Tyr¹²²-hydrophobic cluster in crystal structures.** The part of structures $E1Ca_2 \cdot AIF_4 \cdot ADP$ ($E1 \sim PCa_2 \cdot ADP$ analog, *left*) and $E2 \cdot AIF_4 \cdot (E2 \sim P \text{ analog,$ *right* $})$ around Y122-HC and the bound K⁺ ion are shown in cartoon models (PDB codes: 1T5T and 1XP5 (12, 15)). The two structures were manually aligned with M8-M10 helices, which do not move virtually in the two. K⁺ bound in these structures is shown by *yellow* van der Waals sphere. Gln²⁴⁴ on the A/M3-linker at the immediate vicinity of the bound K⁺ in $E2 \cdot AIF_4^-$ is indicated by ball and stick model.

TABLE 1

Parameters obtained for Ca²⁺ dependence of accumulation of ADP-insensitive EP in Y122-HC

mutants

As shown in Fig. 5, the lumenal Ca^{2+} -induced change in the steady-state accumulation of ADPinsensitive *E*P of the seven Y122-HC mutants in the presence of 0.1 M K⁺ were fitted to the Hill equation. The parameters thus obtained by the least squares fit are listed here. $K_{0.5}$ is the Ca^{2+} concentration giving the half maximum change in the fraction of ADP-insensitive *E*P among the total amount of *E*P, therefore the apparent affinity for lumenal Ca^{2+} . The highest fraction of the ADPinsensitive *E*P at the low Ca^{2+} concentration range (0 ~ 10 μ M) and its lowest fraction at the high Ca^{2+} concentration range (over ~mM) are also listed as the obtained parameters in the fitting (see Fig. 5). The value $n_{\rm H}$ is the Hill coefficient.

Mutant	Fraction of ADP-inser and lowest Ca ²⁺ con	$K_{0.5}$	$n_{ m H}$	
	High Ca ²⁺	Low Ca ²⁺		
	% of total am	тM		
L119A	11	78	0.12	2.1
Y122A	6	86	0.16	1.9
I179A	12	92	0.25	2.3
L180A	9	44	0.11	1.6
I232A	10	89	0.32	2.0
V705A	39	80	0.27	1.6
V726A	23	65	0.28	2.0

${\rm TABLE}\,2$

Affinities of E2P for lumenal Ca^{2+} estimated by kinetic analyses and those determined at steady state

analyses

As described in the Fig. 8*B*, the lumenal Ca^{2+} affinities ($K_{0.5}$) of *E*2P of the wild type and the mutant Y122A were estimated by the kinetic analyses of the lumenal Ca^{2+} -induced *E*2P to *E*1PCa₂ reverse transition and of the forward *E*1PCa₂ to *E*2P transition. The $K_{0.5}$ values and the Hill coefficients (n_H) thus estimated kinetically in Fig. 8 in the absence and presence of 0.1 M K⁺ at pH 7.3 are summarized here. Listed together are those determined by the steady-state analyses of the lumenal Ca^{2+} -induced change in the accumulated fraction of the ADP-insensitive *E*P (*E*2P) under otherwise the same conditions in Figs. 2 and 6 for the mutant Y122A with and without K⁺ and the wild type without K⁺.

	K^+	Estimated by analyses	kinetic	Determined b state analyses	y steady-
	(0.1 M)	$K_{0.5}$	n _H	$K_{0.5}$	$n_{ m H}$
		тM		mМ	
WT	+	1.48	2.2	_a	_ ^a
	_	0.45	1.7	0.40	1.5
Y122A	+	0.15	1.9	0.16	1.9
	_	0.22	1.7	0.22	2.2

^a Not determined because the accumulation of ADP-insensitive *E*P was very low at all the Ca^{2+} concentrations examined and therefore possible change was not revealed.



Scheme 1



Figure 1



Figure 2



Figure 3



Figure 4



Figure 5



Figure 6



Figure 7



Figure 8



Figure 9



Figure 10



Figure 11

SUPPLEMENTAL MATERIAL

for the manuscript by

Kazuo Yamasaki, Guoli Wang, Takashi Daiho, Stefania Danko, and Hiroshi Suzuki

Roles of Tyr¹²²-Hydrophobic Cluster and K⁺ Binding in Ca²⁺-releasing Process of ADP-insensitive Phosphoenzyme of Sarcoplasmic Reticulum Ca²⁺-ATPase



Supplemental Figure 1. ATPase activities of SERCA1a wild type and mutant Y122A at various high concentrations of Ca^{2+} . The ATPase activities of the microsomes expressing SERCA1a wild type (WT) or mutant Y122A, and that of the control microsomes (no SERCA1a expression) were determined at various Ca^{2+} concentrations in the presence of A23187 and 7 mM MgCl₂ as described under "EXPERIMENTAL PROCEDURES." The points at zero Ca^{2+} indicate the activities determined without the CaCl₂ addition. Note that the Ca^{2+} -ATPase activity was completely inhibited in Y122A at all the Ca^{2+} concentrations. In the wild type, the reduction of the activity at the high Ca^{2+} concentrations involves not only the lumenal Ca^{2+} -induced feedback inhibition but also the marked retardation of the Ca^{2+} -ATPase cycle by the substrate CaATP produced at the high Ca^{2+} concentrations. In fact, the affinity of the enzyme for CaATP is approximately 10 times higher than that for MgATP, and the turnover rate of *EP* formed from CaATP (thus Ca^{2+} bound at the catalytic site) is far slower than that formed with MgATP (41). Therefore, even with the wild type, it is not possible to determine the lumenal Ca^{2+} affinity by this experimental design.



Supplemental Figure 2. Ca^{2+} dependence of accumulation of ADP-insensitive *EP* in the steady state in wild type. The fraction of ADP-insensitive *EP* in the total amount of *EP* was determined with the wild type at various Ca^{2+} concentrations as described in Fig. 2 for the mutant Y122A.



Supplemental Figure 3. Lumenal Ca²⁺- and ADP-induced reverse *E2P* decay of Tyr¹²²hydrophobic cluster mutants in the presence of K⁺. With all the seven Y122-HC mutants, *E2P* was first formed from ${}^{32}P_{i}$ in the absence of Ca²⁺, and the *E2P* decay time courses upon the addition of various concentrations of Ca²⁺ and ADP were determined as described in Fig. 7*A* for the representative mutant Y122A. The single exponential decay rates thus obtained were plotted *versus* Ca²⁺ concentrations as in Fig. 7*B*.



Supplemental Figure 4. Different extents of mutational effects of seven residues in Y122-HC on *E2P* hydrolysis and lumenal Ca²⁺ affinity. The *E*2P hydrolysis rates of each of the seven mutants for Y122-HC (*open bars*) were determined in the absence of Ca²⁺ and ADP and presence of 0.1 M K⁺ otherwise as in Fig. 7. The results agree with our previous study (22). The apparent affinities of *E*2P for lumenal Ca²⁺ in the presence of 0.1 M K⁺ at pH 7.3 ($K_{0.5}$, gray bars) were determined with the seven Y122-HC mutants in Fig. 6 and estimated with the wild type in Fig. 8*B*. These values are shown as % of those of the wild type.





Supplemental Figure 5B

Supplemental Figure 5. Inclination and rotational movements of P domain with twisting-like motion of M4 and M5 in $E1PCa_2 \rightarrow E2P + 2Ca^{2+}$. Three structures $E1Ca_2$ (PDB code; 1SU4 (10)), $E1Ca_2 \cdot AIF_4 \cdot ADP$ (the analog for transition state $E1\sim PCa_2 \cdot ADP$ (1T5T (12))), and $E2 \cdot MgF_4^{-2-}$ ($E2 \cdot P_i$ analog (1WPG)) are fitted manually with M8-M10 helices.

(A) Side view of the whole molecule and Top view of the P and A domains, M4, and M5 from the cytoplasmic side are shown by semitransparent mode. The seven residues involved in Y122-HC (Leu¹¹⁹/Tyr¹²², Ile¹⁷⁹/Leu¹⁸⁰, Ile²³², and Val⁷⁰⁵/Val⁷²⁶) are shown by van der Waals spheres. The Val²⁰⁰ loop and T¹⁸¹GES loop are depicted by ribbon diagrams. The alpha carbons of Asp³⁵¹ (located approximately at the center of the P domain) and Glu⁶⁸⁰ (at the outermost part of this domain) were depicted by green spheres to show the motion of the P domain. The three thin red arrows on Top view of E1Ca₂·AlF₄·ADP show the movements of Tyr¹²², Leu¹⁸⁰, and Val⁷²⁶, respectively, from E1Ca₂ to E1Ca₂·AlF₄·ADP. The three *thin red arrows* on the *Side* and *Top views* of E2·MgF₄²⁻ show the movements of these three residues from $E1Ca_2 \cdot AIF_4 \cdot ADP$ to $E2 \cdot MgF_4^{2-1}$ (as indicated by Y122, L180, and V726) to form Y122-HC. The *solid blue arrow* on $E2 \cdot MgF_4^{2-}$ shows the movement of Asp³⁵¹ from $E1Ca_2 \cdot AlF_4 \cdot ADP$ to $E2 \cdot MgF_4^{2-}$. The *dotted yellow arc arrows* on $E2 \cdot MgF_4^{2-}$ show the rotational motion of the P domain around Asp³⁵¹ in the change from $E1Ca_2 \cdot AlF_4 \cdot ADP$ to $E2 \cdot MgF_4^{2-2}$; note that the *dotted vellow arc arrow* placed on Glu⁶⁸⁰ (green sphere) actually indicates the approximate movement of this residue in the change. In E2·MgF₄²⁻, Glu⁶⁸⁰ is located in the immediate vicinity of the Val²⁰⁰ loop and actually involved in the ionic interactions with this loop. The *dotted blue arc arrow* on $E2 \cdot MgF_4^{2-}$ shows the rotational movement of A domain from $E1Ca_2 \cdot AlF_4^{-} \cdot ADP$ to $E2 \cdot MgF_4^{2-}$ around the T¹⁸¹GES-loop region. The *dotted pink* and *light blue ribbons* superimposed on $E2 \cdot MgF_4^{2-}$ indicate the locations of the Val²⁰⁰- and T¹⁸¹GES-loops in $E1Ca_2$ ·AlF₄·ADP, respectively; the Val²⁰⁰ loop, the outermost loop of the A domain, largely moves and comes to the interface with the P domain at Glu⁶⁸⁰ as the A domain largely rotates around the T¹⁸¹GES-loop region.

(*B*) M4 and M5 (*solid ribbons*), M2 (*trace of its alpha carbons*), and the alpha carbons of Asp³⁵¹ and Glu⁶⁸⁰ (*van der Waals spheres*) are exhibited in the three structures; $E1Ca_2$ (*green*), $E1Ca_2 \cdot AIF_4^-$ ·ADP (*red*), and $E2 \cdot MgF_4^{2-}$ (*blue*). The three structures are fitted manually with M8-M10. The *Side view* and *Top view* from the cytoplasmic side are shown. The two Ca^{2+} ions bound in $E1Ca_2 \cdot AIF_4^-$ ·ADP are shown by *yellow van der Waals spheres*, and its release into lumen is roughly shown by *dotted salmonpink arrow* on *Side view*. On *Top view*, the three *thin solid lines* are drawn from Glu⁶⁸⁰ (the outermost part of the P domain) through Asp³⁵¹ (the approximate center of the P domain) to show their alignment in each of the three structures and thus to depict the rotational motion of the P domain (*brown solid arc arrow*) in the change from $E1Ca_2$ to $E1Ca_2 \cdot AIF_4^- \cdot ADP$ and further to $E2 \cdot MgF_4^{2-}$. The three *broken thin lines* are drawn from the cytoplasmic top part of M5 (Phe⁷⁴⁰) through that of M4 (Lys^{329}), and three *dash-dotted thin lines* are from the lumenal end of M5 at the cytoplasmic part (*black broken arc arrow*) and at the lumenal end (*magenta dash-dotted arc arrow*) in the change

from $E1Ca_2/E1Ca_2 \cdot AlF_4 \cdot ADP$ to $E2 \cdot MgF_4^{2-}$. The *orange solid arrow* shows the movement of Tyr¹²² on the cytoplasmic top part of M2 in the change from $E1Ca_2/E1Ca_2 \cdot AlF_4 \cdot ADP$ to $E2 \cdot MgF_4^{2-}$.



Supplemental Figure 6. Schematic model for rearrangement of P and A domains for *EP* isomerization and Ca²⁺ release. Depicted is our schematic model for the motions of the A and P domains in the successive two steps (*large yellow arrows*); $E1PCa_2 \rightarrow E2PCa_2$ (loss of the ADP-sensitivity) and $E2PCa_2 \rightarrow E2P + 2Ca^{2+}$ (Ca²⁺ release). The model is made on the basis of the observed mutational effects of Y122-HC and Val²⁰⁰ loop (22-24). The top view from the cytoplasmic side is shown (as in *Top view* of Supplemental Figure 3A). The P and A domains, Val²⁰⁰ loop, T¹⁸¹GES loop, and Asp³⁵¹ in *E1PCa*₂ and *E2P* are positioned according to the structures of $E1Ca_2 \cdot AIF_4$ ·ADP (1T5T) and $E2 \cdot MgF_4^{-2-}$ (1WPG), respectively. *E2PCa*₂ is the postulated intermediate state. The positions of the seven residues involved in Y122-HC in the three regions are shown by the representative ones (Y122, L180, and V705). In the pictures placed for the two *large yellow arrows*, the structure (*broken lines*) before its change was superimposed on that after the change, and the motions of the A and P domains are indicated by *dotted blue arrow* and *dotted red arrow*, respectively.

$E1PCa_2 \rightarrow E2PCa_2$; the loss of ADP-sensitivity:

The A domain rotates largely (probably due to the strain of the A/M3-linker) around the T¹⁸¹GES loop and the P domain inclines to some extent toward the A domain, thereby the T¹⁸¹GES loop docks onto the Asp³⁵¹-region producing hydrogen-bonding interactions (*dotted blue line*), hence causing the loss of the ADP-sensitivity (the T¹⁸¹GES loop blocks the access of ADP β -phosphate to Asp³⁵¹- acylphosphate). In this state, the two interaction networks at Y122-HC and at the Val²⁰⁰ loop are not yet fully produced. Our previous mutation study on the interaction network between Arg³³⁴/Arg³²⁴ of the P domain and top part of M2 (Glu¹²³/Tyr¹²²/Glu¹²¹/Glu¹¹⁷/Asn¹¹⁴/Glu¹¹³/Asn¹¹¹/Glu¹⁰⁹/Gln¹⁰⁸) indicated (22) that the inclination of the P domain occur at least to some extent along this interaction network for the loss of the ADP-sensitivity, *E*1PCa₂ \rightarrow *E*2PCa₂.

 $E2PCa_2 \rightarrow E2P + 2Ca^{2+}$; Ca^{2+} release after the loss of ADP-sensitivity:

The P and A domains further rotate slightly in the opposite directions and the top part of M2 (Tyr^{122}/Leu^{119}) likely moves more (*green solid arrow*), thereby they produce the interactions fully at Y122-HC and at Val²⁰⁰ loop (*dotted yellow circles*) so as to realize and stabilize the Ca²⁺-released structure of *E*2P. The native and appropriately short length of the A/M1-linker (its strain) critically contributes to the structural changes for Ca²⁺-deocclusion/release and formation of Y122-HC in $E2PCa_2 \rightarrow E2P + 2Ca^{2+}$; in this case, especially for the vertical motions of the P and A domains and M4/M5, and the inclination of M2. These vertical factors are not depicted in this top view.

Upon these changes, the catalytic site is rearranged to produce its appropriate configuration for the subsequent Asp³⁵¹-acylphosphate hydrolysis, *e.g.* the T¹⁸¹GES loop comes to the position where the Glu¹⁸³-coordinated water molecule can attack Asp³⁵¹-acylphosphate. Hence in this structural mechanism, the possible hydrolysis without releasing Ca²⁺ is avoided, and the Ca²⁺ release and the subsequent *E*2P hydrolysis are accomplished as the ordered sequence.

In the wild type, all these successive structural events in $E1PCa_2 \rightarrow E2PCa_2 \rightarrow E2P + 2Ca^{2+}$ occur in coordinated manner and thus rapidly; therefore $E2PCa_2$ does not accumulate stably in the wild type.