Phosphoenzyme Conversion of the Sarcoplasmic Reticulum Ca\textsuperscript{2+}-ATPase

MOLECULAR INTERPRETATION OF INFRARED DIFFERENCE SPECTRA

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Time-resolved Fourier transform infrared difference spectra of the phosphoenzyme conversion and \(\text{Ca}^{2+}\) release reaction \((\text{Ca}_2\text{E}_1\text{P} \rightarrow \text{E}_2\text{P})\) of the sarcoplasmic reticulum \(\text{Ca}^{2+}\)-ATPase were recorded at \(\text{pH} 7\) and \(1\ °\text{C}\) in \(\text{H}_2\text{O}\) and \(\text{D}_2\text{O}\). In the amide I spectral region, the spectra indicate backbone conformational changes preserving conformational changes of the preceding phosphorylation step, \(\beta\)-sheet or turn structures (band at \(1685\ \text{cm}^{-1}\)) and \(\alpha\)-helical structures (band at \(1653\ \text{cm}^{-1}\)) seem to be involved. Spectra of the model compound EDTA for \(\text{Ca}^{2+}\) chelation indicate the assignment of bands at \(1570,\ 1554,\) and \(1559\ \text{cm}^{-1}\) to \(\text{Ca}^{2+}\) chelating Asp and Glu carboxylate groups partially shielded from the aqueous environment. In addition, an \(\text{E}_2\text{P}\) band at \(1638\ \text{cm}^{-1}\) has been tentatively assigned to a carboxylate group in a special environment. A Tyr residue seems to be involved in the reaction (band at \(1517\ \text{cm}^{-1}\) in \(\text{H}_2\text{O}\) and \(1515\ \text{cm}^{-1}\) in \(\text{D}_2\text{O}\)). A band at \(1192\ \text{cm}^{-1}\) was shown by isotopic replacement in the \(\gamma\)-phosphate of ATP to originate from the \(\text{E}_2\text{P}\) phosphate group. This is a clear indication that the immediate environment of the phosphoenzyme phosphate group changes in the conversion reaction, altering phosphate geometry and/or electron distribution.

Muscle relaxation is mediated by the removal of cytosolic \(\text{Ca}^{2+}\) by the \(\text{Ca}^{2+}\)-ATPase of the sarcoplasmic reticulum membrane. The ATPase couples active \(\text{Ca}^{2+}\) transport to the hydrolysis of ATP (1–4) in a reaction cycle that is shown in Scheme 1. In an essential reaction step, ATP reacts with Asp-351 to form a phosphoenzyme intermediate \((\text{Ca}_2\text{E}_1\rightarrow \text{Ca}_2\text{E}_1\text{P})\), which then converts from an ADP-sensitive form \((\text{Ca}_2\text{E}_1\text{P})\) to an ADP-insensitive form \((\text{E}_2\text{P})\) that is more rapidly hydrolyzed. This phosphoenzyme conversion is associated with \(\text{Ca}^{2+}\) release toward the sarcoplasmic reticulum lumen against the concentration gradient (1, 3, 5). The structural origin of the change of accessibility of the \(\text{Ca}^{2+}\) sites and of the essential reduction of \(\text{Ca}^{2+}\) affinity upon phosphoenzyme conversion \(\text{Ca}_2\text{E}_1\text{P} \rightarrow \text{E}_2\text{P}\) have yet to be elucidated.

The potentially large body of structural information regarding the ATPase reaction cycle provided by infrared spectroscopy has only recently begun to be exploited using the approach of effector molecule-induced infrared difference spectroscopy (6–13). The method uses the release of effector molecules from biologically “silent” photolabile derivatives, termed caged compounds (14–17), to generate high quality infrared difference spectra. (The reaction products and infrared difference spectra of caged ATP photolysis and of side reactions have been characterized (18–20).) The absorbance changes seen in these difference spectra give evidence for conformational changes of the polypeptide backbone and for alterations to the environment of amino acid side chains that take place in the reaction investigated.

Spectra of the phosphoenzyme conversion reaction in the 1800 to 1000 \text{cm}^{-1} spectral range have previously been described (8) and in that work were calculated by subtracting two difference spectra obtained with two different types of samples; a normalized difference spectrum of the \(\text{Ca}_2\text{E}_1\rightarrow \text{Ca}_2\text{E}_1\text{P}\) reaction was subtracted from a normalized difference spectrum of the \(\text{Ca}_2\text{E}_1\rightarrow \text{E}_2\text{P}\) reaction. The use of two different samples in the subtraction and the normalization to identical protein content limit the reliability of these spectra and make it desirable to obtain the phosphoenzyme conversion spectrum more directly. This is possible using time-resolved rapid scan Fourier transform infrared (FTIR) spectroscopy (10). Using this approach, we present here the spectra of the phosphoenzyme conversion reaction obtained in \(\text{H}_2\text{O}\) and \(\text{D}_2\text{O}\) after the release of unlabeled ATP or \(\gamma\text{-}[\text{18O}]\text{ATP}\). In particular, bands of the putative \(\text{Ca}^{2+}\) chelating carboxylate groups and of the phosphoenzyme phosphate group are discussed.

MATERIALS AND METHODS

Sample Preparation—Samples for time-resolved infrared spectroscopy of the \(\text{Ca}_2\text{E}_1\text{P} \rightarrow \text{E}_2\text{P}\) reaction were prepared as described previously (8, 10) by removal of free water from a sarcoplasmic reticulum suspension equilibrated in \(\text{H}_2\text{O}\) or \(\text{D}_2\text{O}\) buffer. Samples were immediately rehydrated with \(\text{H}_2\text{O}\) or \(\text{D}_2\text{O}\) with or without \(20\%\ \text{Me}_2\text{SO}\). This method resulted in active ATPase samples (6). Approximate concentrations of unlabeled ATP or \(\gamma\text{-}[\text{18O}]\text{ATP}\) were 0.7 mM ATPase, 300 mM imidazole, pH 7.0, 1 mM CaCl\(_2\), 20 mM glutathione, 20 mM caged ATP, 0.5 mM ADP, 2 mg/ml adenylate kinase, and \(20\%\ \text{Me}_2\text{SO}\) in approximately \(1\ \mu\text{l}\) of sample volume. Approximately 2–3 mM ATP was released per flash. Caged \(\gamma\text{-}[\text{18O}]\text{ATP}\) at \(91\%\) isotopic enrichment per oxygen atom was a gift of M. R. Webb and J. E. T. Corrie (National Institute for Medical Research, London).

FTIR Measurements—Time-resolved FTIR measurements of the \(\text{Ca}_2\text{E}_1\text{P} \rightarrow \text{E}_2\text{P}\) reaction were performed at \(1\ °\text{C}\) with a modified Bruker IFS 66 spectrometer as described previously (10). Difference spectra for the reaction were obtained by subtracting a spectrum representing predominantly \(\text{Ca}_2\text{E}_1\text{P}\), and to a small extent \(\text{E}_2\text{P}\) (recorded 3.3–11 s after photolysis of caged ATP in \(\text{H}_2\text{O}\) or after 11–19 s in \(\text{D}_2\text{O}\)), from a spectrum representing \(\text{E}_2\text{P}\) (recorded after 88 and 146 s in \(\text{H}_2\text{O}\) and \(\text{D}_2\text{O}\), respectively). The resulting spectrum was normalized as described (10) to the full amplitude of the absorbance changes associated with the phosphoenzyme conversion reaction.

1 The abbreviations used are: FTIR, Fourier transform infrared; \(\text{Ca}^{2+}\)-ATPase, \(\text{Ca}^{2+}\) transporting ATPase (EC 3.6.1.38); caged ATP, P\(^2\)-1-(2-nitrophenyl)ethyladenosine 5’-triphosphate; \(\text{Ca}_2\text{E}_1\text{P}\); ADP-sensitive phosphoenzyme; \(\text{E}_2\text{P}\); ADP-insensitive phosphoenzyme; \(\nu_s\), symmetric stretching vibration; \(\nu_{as}\), antisymmetric stretching vibration.

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As the difference spectra were obtained directly from the time-resolved measurements, a normalization of spectra to an identical protein concentration is in principle not necessary. However, for a better comparison of samples in \( \text{H}_2\text{O} \) and \( \text{D}_2\text{O} \) and to prevent the possible predominance of individual samples with high protein content in the averaged spectra, spectra were normalized to an identical protein concentration before averaging, as described (21).

Absorbance spectra in \( \text{H}_2\text{O} \) and \( \text{D}_2\text{O} \) of the model compound for \( \text{Ca}^{2+} \) release EDTA were recorded with and without \( \text{Ca}^{2+} \) at \( 20^\circ\text{C} \) and pH 12.8. Small variations in the path length between different samples were corrected by normalizing the water absorption of every sample to a single water spectrum that served as a standard for the path length. Sample concentrations were 84 mM.

**Band-narrowing Procedures**—Apart from second-derivative spectra, the following procedure was applied. The original spectrum was smoothed over 24 cm\(^{-1}\) and multiplied by 0.95. This spectrum was then subtracted from the original spectrum, which eliminates broad features of the original spectrum. The resulting spectrum is dominated by the fine-structure in the original spectrum, and thus we term the method “fine-structure enhancement” in the following text. When tested with protein absorbance spectra, this method gives results similar to Fourier self-deconvolution (data not shown). For a clearer presentation, fine-structure enhanced spectra were multiplied by 10 and second-derivative spectra by \( 2^{15} \).

To assess whether peaks in the fine-structure enhanced spectra correspond to “true” component bands or are the result of artifacts in the band-narrowing procedure (see Fig. 3), the original difference spectra were fitted, and the resulting fit was fine-structure enhanced and compared with the fine-structure enhanced original spectra (data not shown). Interestingly, it was found that the spectral region of 1700–1670 cm\(^{-1}\) can accurately be fitted by only three bands at 1693 (–), 1686 (–), and 1670 cm\(^{-1}\) (++) despite the plateau at 1678/1676 cm\(^{-1}\) in the fine-structure enhanced \( \text{H}_2\text{O} \) spectrum (solid line in Fig. 3B). It is not necessary to introduce an additional band in the fitting model to reproduce that plateau.

**RESULTS AND DISCUSSION**

**Model Spectra of \( \text{Ca}^{2+} \) Release**—From site-directed mutagenesis studies it is thought that Glu-309, Glu-771, and Asp-800 form part of the high affinity \( \text{Ca}^{2+} \) binding sites of \( \text{Ca}_2\text{E}_1 \) and \( \text{Ca}_2\text{E}_1^- \) (4). Thus, we studied the effect of \( \text{Ca}^{2+} \) binding on the infrared spectrum of carboxylate groups using the \( \text{Ca}^{2+} \) chelator EDTA.

Fig. 1A shows absorbance spectra in \( \text{D}_2\text{O} \) of free EDTA (solid line) and of the EDTA complex with \( \text{Ca}^{2+} \) (dotted line). The bands near 1590 and 1410 cm\(^{-1}\) can be assigned to the COO\(^{-}\) antisymmetric stretching vibration \( (v_{as}) \) and the symmetric stretching vibration \( (v_s) \), respectively. The shoulder at 1612 cm\(^{-1}\) indicates some heterogeneity of the COO\(^{-}\) groups, resulting from a small proportion of EDTA with protonated nitrogen atoms (22, 23). Upon \( \text{Ca}^{2+} \) release there is a downshift of 4–8 cm\(^{-1}\) for both of the main bands (at 1590 and 1414 cm\(^{-1}\)), which translates in the difference spectrum of \( \text{Ca}^{2+} \) release (absorbance of free EDTA minus absorbance of the \( \text{Ca}^{2+} \) complex) into the two minimum/maximum features at 1590/1568 and 1418/1400 cm\(^{-1}\).

When \( \text{D}_2\text{O} \) is replaced by \( \text{H}_2\text{O} \), the bands of the antisymmetric stretching vibration are downshifted by 6–8 cm\(^{-1}\) for free EDTA and for the complex (data not shown). The band of the symmetric stretching vibration is less sensitive and shifts...
downwards only for free EDTA (by 2 cm$^{-1}$). A similar behavior has been observed for sodium acetate (24) and for Asp and Glu (25, 26).

Thus, the model spectra have identified marker band pairs for Ca$^{2+}$ release from carboxylate groups near 1575 and 1410 cm$^{-1}$. The former is expected to be sensitive to H$_2$O $\rightarrow$ D$_2$O replacement with a higher frequency in D$_2$O.

The Phosphoenzyme Conversion Spectrum—Fig. 2A shows the difference spectrum of the phosphoenzyme conversion and Ca$^{2+}$ release reaction, Ca$_2$E$_1$P $\rightarrow$ E$_2$P (solid line), and the respective spectrum in D$_2$O buffer (dashed line). Positive bands are characteristic for E$_2$P, negative bands for Ca$_2$E$_1$P. Fig. 3A shows the same spectra on an expanded scale.

Ca$^{2+}$ Release from Carboxylate Groups—In the phosphoenzyme conversion spectrum, similar band profiles are observed as in the model difference spectrum for Ca$^{2+}$ release (Fig. 1B) at 1570/1554 and 1411/1399 cm$^{-1}$ (numbers for H$_2$O). The slight upshift upon H$_2$O $\rightarrow$ D$_2$O exchange characteristic for the $\nu_{\text{as}}$ vibration of carboxylate groups (see above) is also notable (1570 $\rightarrow$ 1572 cm$^{-1}$, 1554 $\rightarrow$ 1555 cm$^{-1}$) and is best seen in Fig. 3A. However, the shift is less than that observed above for completely exposed carboxylate groups. The lower wavenumber of the 1570/1554 cm$^{-1}$ band pair as compared with the model spectra (1586/1556 cm$^{-1}$ in H$_2$O and 1590/1568 cm$^{-1}$ in D$_2$O) is expected because the $\nu_{\text{as}}$ (COO$^-$) band position of free Asp and Glu residues (25, 26) is 2–4 cm$^{-1}$ (Asp) and 18 cm$^{-1}$ (Glu) lower than that of EDTA (in H$_2$O and D$_2$O). These observations support and substantiate the tentative assignment of the band pairs at 1570/1554 and 1411/1399 cm$^{-1}$ (numbers for H$_2$O) to Ca$^{2+}$ chelating residues of the ATPase (8). The relatively small shift observed for the 1570/1554 cm$^{-1}$ band pair in D$_2$O may indicate that these groups are not completely exposed to water.

The band at 1638 cm$^{-1}$ also shows the upshift characteristic of a $\nu_{\text{as}}$(COO$^-$) band upon H$_2$O $\rightarrow$ D$_2$O replacement (best seen in Fig. 3A). Other bands absorbing in that region, i.e. the amide I mode of $\beta$-sheet structures, and side chain modes of His, Arg, and Lys should either show downshifts of about 10 cm$^{-1}$ (backbone and His) (26–30), 50 cm$^{-1}$ (Arg) (25, 26), and 400 cm$^{-1}$ (Lys) (31) or should remain unchanged if the respective groups are inaccessible to H$^1$ $\rightarrow$ D$^1$ exchange. The position of the 1638 cm$^{-1}$ band is rather unusual for a $\nu_{\text{as}}$ band of a COO$^-$ group, which absorbs for model compounds in aqueous solution at 1574 cm$^{-1}$ (Asp) or at 1560 cm$^{-1}$ (Glu) (25). However, a strong salt bridge or hydrogen bond to one of the oxygens of a carboxylate group could shift the band well above 1600 cm$^{-1}$ (24, 32), and a reasonable scenario for the band at 1638 cm$^{-1}$ could be that the parting Ca$^{2+}$ is replaced by a strong hydrogen bond donor or a positive charge.

Bands at 1758 and 1710 cm$^{-1}$ have tentatively been assigned to the protonation of at least two chelating carboxylate groups (33), which is in line with the small downshift observed upon H$_2$O $\rightarrow$ D$_2$O replacement.

Protein Backbone: the Amide I Region—The largest absorbance changes (up to 0.5% of the total protein absorbance) are observed in the amide I region of the spectrum (1700–1610 cm$^{-1}$) as are the strongest effects of protein deuteration. In this region, the amide I mode of the polypeptide backbone as well as the side chains of Asn, Gln, Arg, His$^{+}$, and Lys absorb strongly (25, 26, 30). In the lower wavenumber part of that region, Asp and Glu COO$^-$ groups with strong interactions may contribute as discussed above.

Large downshifts ($\approx$ 30 cm$^{-1}$) of bands upon H$_2$O $\rightarrow$ D$_2$O replacement are characteristic for the side chain absorptions of Asn, Gln, Lys, and Arg, as are small upshifts as discussed above for COO$^-$ bands and downshifts of up to 10 cm$^{-1}$ for amide I and His$^{+}$ bands. No band shifts are expected for groups that are located in parts of the protein that are not accessible to deuteration.

The large number of alterations in this spectral region, when H$_2$O is replaced by D$_2$O, makes it difficult to arrive at a unique explanation for the band shifts, and additional experiments as well as data processing were necessary, as described below.

The recording of spectra as soon as possible after H$_2$O $\rightarrow$ D$_2$O replacement is important, as data processing were necessary, as described below.
The only clear exception is the minimum at ~1630 cm\(^{-1}\) that develops over a few hours of incubation. This position is characteristic of amide I modes of \(\beta\)-sheet structures and of the C=O group of deuterated Asn or Gln residues (26).

Band-narrowing procedures (see "Materials and Methods") were applied to identify band shifts upon \(^1\text{H} \rightarrow ^2\text{H}\) exchange. Fig. 3A shows the original difference spectra and Fig. 3, B and C, the resulting spectra after band narrowing. These reveal essentially the same peak positions in \(\text{H}_2\text{O}\) and \(^2\text{H}_2\text{O}\) (see Fig. 3, B and C), with one exception. The negative band at 1689 cm\(^{-1}\) in the unprocessed original \(\text{H}_2\text{O}\) spectrum (solid line in Fig. 3A) is composed of at least two bands, giving minima in the processed spectrum at 1693 and 1685 cm\(^{-1}\) (solid line in Fig. 3, B and C). The highest wavenumber component of the negative band seems to be nearly unaffected by protein deuteration and is observed in \(^2\text{H}_2\text{O}\) at 1692 cm\(^{-1}\). It could be caused by a conformational change of \(\beta\)-sheet or turn structures or an Asn, Gln, or Arg side chain that is located in the core of the protein and is inaccessible to deuteration.

The other component of the 1689 cm\(^{-1}\) band seems to shift from its position in the processed spectrum (Fig. 3B) at 1685 (\(\text{H}_2\text{O}\)) to 1677 cm\(^{-1}\) (\(^2\text{H}_2\text{O}\)) and to cancel part of the adjacent positive band observed at 1671 cm\(^{-1}\) in \(\text{H}_2\text{O}\) (Fig. 3A). This shift is characteristic of amide I modes of the polypeptide backbone, and the position then indicates a conformational change of \(\beta\)-sheet or turn structures. These structures are predicted to occur only in the extramembraneous domains of the protein (34, 35), and thus the observed conformational change is likely to take place in these protein domains.

As the positions of the other bands in the amide I region are hardly affected by deuteration, the isotope effects are the result either of intensity changes or of bands that are not evident after applying the band-narrowing procedures because they are broader than the ones detected. Narrow bands tend to dominate the processed spectra.

The bands hardly affected by \(^1\text{H} \rightarrow ^2\text{H}\) exchange are found at 1653, 1638, 1621, and 1607 cm\(^{-1}\). A band at 1653 cm\(^{-1}\) is often observed for \(\alpha\)-helical structures and is often hardly affected by \(\text{H}_2\text{O} \rightarrow ^2\text{H}_2\text{O}\) replacement as observed here. The band at 1638 cm\(^{-1}\), which shifts slightly upward, was tentatively assigned above to a COO\(^-\) group. The bands at 1621 and 1607 cm\(^{-1}\) seem to retain their position in \(^2\text{H}_2\text{O}\) with a possible overlap of additional bands in \(\text{H}_2\text{O}\) or \(^2\text{H}_2\text{O}\). The band at 1621 cm\(^{-1}\) may be assigned to a \(\beta\)-sheet structure or to the imide group of a Pro.
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residue in a helical or unordered conformation. The imide band is found approximately 20 cm⁻¹ lower than an amide band (36). Thus, the band at 1607 cm⁻¹, the position of which seems to be too low for an amide I band, could also be caused by a Pro imide group. Interestingly, there are three Pro residues in the putative Ca²⁺ binding transmembrane helices M4 and M6, mutation of which affects Ca²⁺ affinity and phosphoenzyme conversion (4). Alternatively, both bands may be caused by COO⁻ side chain groups not interacting with bulk water. This assumption relies on the fact that the bands do not show the upshift upon H₂O → D₂O replacement characteristic for carboxylate groups in water.

Protein Backbone: the Amide II Region—None of the three bands in the amide II region (1570–1530 cm⁻¹) shows the strong sensitivity toward H² → D² exchange expected for the amide II mode of the protein backbone. The amide II mode of backbone elements accessible to deuterium therefore does not seem to be affected by phosphoenzyme conversion and Ca²⁺ release. Two of the bands (at 1570 and 1554 cm⁻¹) have been tentatively attributed above to the ν(CH₃) vibration of COO⁻ groups.

Protein Backbone: the Amide III Region—The amide III mode absorbs between 1400 and 1200 cm⁻¹ and is sensitive to deuterium (37). This property is observed in the spectra for the bands at 1337 and 1318 cm⁻¹ (see Fig. 2A), which therefore might be attributed to amide III modes. The position of these bands is characteristic for turn structures (37) and is probably related to the amide I band at 1687 cm⁻¹, which has tentatively been assigned to turn or β-sheet structures. Alternatively, the bands at 1337 and 1318 cm⁻¹ may be caused by the δ(COH) mode of Ser, Asp, or Glu with a weakly bonded OH group.

Side Chain Modes Other than Carboxyl Modes—As mentioned above, bands of the side chains of Asn, Gln, Arg, and Lys in the amide I region show relatively large shifts upon deuteration. The extinction coefficient of the former three residues is relatively high (25, 26), whereas that of Lys is smaller. Thus, Lys bands may be masked by stronger bands. The HisH⁻ mode near 1631 cm⁻¹ (30) shows a 10 cm⁻¹ downshift in H₂O (26) and absorbs relatively strongly in H₂O (30). These characteristic shifts have not been observed in the spectra, and thus there is no clear evidence for the participation of Asn, Gln, Arg, and HisH⁻ in the phosphoenzyme conversion and Ca²⁺ release reaction. This statement holds only for those residues that are accessible to H² → D² exchange, which should include residues in the ATP binding site, the catalytic site, and at least part of the Ca²⁺ binding sites, because several bands that were tentatively assigned to the Ca²⁺ binding sites show an effect upon H₂O → D₂O replacement (see above).

The position of the band at 1517 cm⁻¹ and its slight downshift of 2 cm⁻¹ in D₂O is characteristic for a ring mode of protonated Tyr (25, 26, 30). Also, the band pairs at 1283 and 1264 cm⁻¹ may be attributed to the ν(C-O) mode of Tyr but also to a Trp mode, which is observed for indole at 1276 cm⁻¹ (38, 39). It has been suggested that Tyr-763 is involved in the cytoplasmic gate to the Ca²⁺ binding sites (4). The very small band at 1365 cm⁻¹ might be caused by a δ(CH₃) mode of aliphatic side chains.

Absorption of the Phosphate Group—Fig. 2B shows a comparison between the conversion spectrum obtained with unlabeled ATP and that obtained with [γ-¹⁵O]ATP. With the heavier isotope, phosphate bands are expected to be downshifted, thereby enabling the identification in the spectrum of alterations to the phosphate group. As the γ-phosphate is transferred to Asp-351 before phosphoenzyme conversion, differences between the spectra with the two isotopes will identify the absorbance of the phosphoenzyme phosphate group. As expected, the two spectra superimpose very well above 1250 cm⁻¹, where phosphate groups do not absorb. However, the band at 1192 cm⁻¹ is reduced upon isotopic substitution. Instead, the intensity is higher for [γ-¹⁵O]ATP than for the unlabeled phosphate bands (4). This identification of a phosphate band in the difference spectrum clearly shows that the conversion reaction considerably changes the environment of the phosphate group, thus affecting its electron density distribution and/or binding geometry. The band position at 1192 cm⁻¹ is rather unusual for a phosphate group and could be explained in two ways: (i) a widening of the P–O angles leading to a stronger coupling between the P–O vibrations and thus to a stronger splitting between the ν₁ and ν₃ modes; or (ii) an increase in electron density of some or all of the P–O bonds, either by breaking of the hydrogen bonds to all phosphate oxygens or by strong hydrogen bonding to one or two of the phosphate oxygen atoms, thus increasing the electron density in the other P–O bond(s). The band is also affected by H₂O → D₂O replacement, which seems to indicate that the phosphate oxygen interacts with either a water molecule or deuterated protein residues.

The active site of Eₐ-P was shown to be in a closed conformation (41) shielded from bulk water with a hydrophobic environment detected close to the ribose OH groups of a fluorescent ATP analogue (42, 43). This finding is in contrast to the properties of Ca₂⁺E₁-P, and it is thought that the decrease in the

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**Fig. 4.** Difference spectra of the Ca⁺⁺E₃-ATP → Ca⁺⁺E₃-P (solid line) and the Ca⁺⁺E₁-ATP → E₁-P (dotted line) reaction at pH 7 and 1 °C.
active site water activity upon the CaE₂P → E₃P conversion is responsible for the higher hydrolysis rate of E₃P as compared with CaE₁P⁻ (44). The infrared spectra presented here show that this conformational change has a direct effect on phosphate conformation and/or electron density distribution. Creation of a hydrophobic environment alone however, may not be sufficient to explain the phosphate band at 1192 cm⁻¹, because dehydration of the model compound acetyl phosphate shifts the νPO₃²⁻ band only from 1132 to 1177 cm⁻¹ at most (data not shown).

**Comparison with Spectra of ATPase Phosphorylation**—Fig. 4 shows spectra in ²H₂O of ATPase phosphorylation CaE₁ATP → CaE₁⁻P (solid line) (21) and of the overall reaction of phosphorylation and phosphenzyme conversion CaE₁ATP → E₃P (dotted line). These spectra were obtained from time-resolved measurements of the same set of samples. Negative bands are characteristic for CaE₁ATP, positive bands either for CaE₁⁻P (solid line) or E₃P (dotted line). (The spectrum of CaE₁ATP → CaE₁⁻P also shows a small contribution because of the E₃P marker band near 1750 cm⁻¹.) The spectral region shown includes the amide I region (1700–1610 cm⁻¹) that is sensitive to conformational changes of the protein backbone. The comparison indicates bands characteristic for CaE₁⁻P are still present in E₃P, which is especially obvious for the bands at 1682, 1630, and 1593 cm⁻¹. This finding indicates that at least some of the alterations to the protein conformation induced by ATPase phosphorylation seem to be preserved or even “enhanced” in the subsequent transition to E₃P. It seems as if the enzyme conformation on the way from CaE₁⁻P to E₃P goes further away from the pre-phosphorylation conformation of CaE₁ATP instead of returning to it.

**CONCLUSIONS**

Infrared difference spectra show that the protein conformation changes in the phosphenzyme conversion reaction, preserving conformational changes of the preceding step of enzyme phosphorylation. β-sheet or turn structures of the extramembrane domains and most likely α-helical structures seem to be affected. The net change of secondary structures, however, is small (10).

The release of Ca²⁺ proceeds from carboxylate groups that seem to be partly shielded from the aqueous environment. It is associated with the protonation of at least two carboxylate groups presumably involved in Ca²⁺ chelation. A change of environment of a Tyr residue was detected as well as a direct effect of phosphenzyme conversion on the geometry and/or electron density distribution of the phosphenzyme phosphate group. The latter is likely to be a prerequisite for the higher susceptibility toward hydrolytic attack of E₃P as compared with CaE₁⁻P. Interestingly, the currently assigned bands of the phosphate group and of Ca²⁺ release from carboxylate groups appear with the same reaction rate (10), showing that Ca²⁺ release and the change of phosphate environment proceed at the same time. This finding rules out a significant population of CaE₂E₂P, a postulated state (45) with phosphate properties of E₃P that still binds Ca²⁺.

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