Characterization of the Protonation and Hydrogen Bonding State of the Histidine Residues in IIA\textsuperscript{mut}, a Domain of the Phosphoenolpyruvate-Dependent Mannitol-Specific Transport Protein\textsuperscript{+}

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ABSTRACT: The A domain of the mannitol-specific EI, IIA\textsuperscript{mut}, was subcloned and proven to be functional in the isolated form (Van Weeghel et al., 1991). It contains a histidine phosphorylation site, the first of two phosphorylation sites in the parent protein. In this paper, we describe the characterization of the three histidine residues in IIA\textsuperscript{mut} with respect to their protonation and hydrogen bonding state, using $^1$H [$^{15}$N] heteronuclear NMR techniques and protein selectively enriched with [6,2-15N] histidine. The active site residue has a low $pK_a$ (<5.8) and shows no hydrogen bond interactions. The proton in the neutral ring is located at the Ne2 position, which also proved to be the site of phosphorylation. The phosphorylation raises the $pK_a$ of the active site histidine considerably but does not change the hydrogen bond situation. The other two histidine residues, one of which is probably located on the surface of the protein, were also characterized. Both show hydrogen bond interactions in the unphosphorylated protein, but these are disturbed by the phosphorylation process. These observations, combined with small changes in $pK_a$ and titration behavior, indicate that the IIA\textsuperscript{mut} changes its conformation upon phosphorylation.

EII\textsuperscript{mut} is the membrane component of the mannitol-specific phosphoenolpyruvate (PEP)-dependent phosphotransferase system (PTS)\textsuperscript{2} in Escherichia coli. It takes care of the transport of mannitol across the inner membrane and its concomitant phosphorylation. The phosphoryl group in this process is donated initially by PEP and is shuttled to the EII\textsuperscript{mut} via EI and HPr. Phospho-HPr phosphorylates a histidine residue (His554) of EII\textsuperscript{mut}; the phosphoryl group is then transferred to a cysteine residue, Cys384, and finally to the incoming sugar. (Pas & Robillard, 1988; Pas et al., 1988, 1991). EII\textsuperscript{mut} consists of three domains, A, B, and C. The A and B domain are cytoplasmic and contain the phosphorylation sites, the C domain is located in the membrane. The A domain, IIA\textsuperscript{mut}, containing the His554 phosphorylation site was recently subcloned, overexpressed, and purified by Van Weeghel et al. (1991). The separated domain is able to perform the same tasks as when integrated in the parent enzyme. The general characterization of IIA\textsuperscript{mut} and its interaction with HPr will be treated in a separate study. This paper treats the hydrogen bond structure at the three histidine residues in IIA\textsuperscript{mut} and the changes that take place upon phosphorylation of the protein.

MATERIALS AND METHODS

Bacterial Strains and Growth. Escherichia coli JC411 (his, arg, leu, mer) was transformed with the plasmid pMccI, described by Van Weeghel et al. (1991) using standard procedures. The strain was grown at 37°C on the mineral medium described by Van Dijk et al. (1990), containing 6,2-15N-labeled histidine (20 µg/mL) as the sole histidine source and chloramphenical (20 µg/mL) instead of ampicillin as antibiotic. Cells were allowed to grow for 2-3 days after which they were harvested by centrifugation (10000g, 5 min, 4°C).

Purification of IIA\textsuperscript{mut}. IIA\textsuperscript{mut} was purified at 4°C by a procedure slightly different from the one described by Van Weeghel et al. (1991). Cells were resuspended in buffer 1 (20 mM Tris-HCl (pH 7.6), 1 mM DTT, 1 mM EDTA, 1 mM NaN\textsubscript{3}; 10 mL/g wet weight). DNase and RNase (0.03 mg/mL) were added, and the cells were disrupted in a French press (10 000 psi). Cell debris was removed by centrifugation (48000g, 2 h) and the supernatant was collected. The pellet was resuspended in buffer 1 (2 mL/g wet weight) and re centrifuged as described. The supernatant was collected and added to the first supernatant. The resulting solution was dialyzed against 2 L of buffer 2 (20 mM Tris-HCl (pH 9.5), 1 mM DTT, 150 mM NaCl, 1 mM NaN\textsubscript{3}). The dialyzed solution was loaded on a Q-Sepharose column (2×20 cm; 0.7 mL/min), equilibrated in buffer 2. After the column was washed with at least an additional 200 mL of buffer 2, the IIA\textsuperscript{mut} was eluted with a linear NaCl gradient (2×150 mL; 150–300 mM NaCl). Fractions were collected and assayed for IIA\textsuperscript{mut} activity with the IIA\textsuperscript{mut} complementation assay (Van Weeghel et al., 1991). IIA\textsuperscript{mut}-containing fractions were pooled and dialyzed overnight against 3 L of buffer 3 (10 mM Tris-HCl (pH 7.6), 1 mM NaN\textsubscript{3}) after which the solution was concentrated to 30 mL on an Amicon ultrafiltration apparatus, using a FILTRON Omega 3 filter. The concentrated solution was loaded on a Sephadex G75 column (5×70 cm, 2 mL/min) equilibrated in buffer 3. Fractions were collected and

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3 Abbreviations: EI, enzyme I; glc, glucose; HMOC, heteronuclear multiple quantum coherence; lac, lactose; man, mannoside; mtl, mannitol; PEP, phosphoenolpyruvate; PTS, phosphoenolpyruvate-dependent phosphotransferase system; TPII, time-proportional phase incrementation; xylitol.
assayed for IIA<sup>mtl</sup> activity as described. The IIA<sup>mtl</sup>-containing fractions were pooled and subjected to polyacrylamide gel electrophoresis. The protein migrated as a single homogeneous band.

Other Enzyme Purifications and Determination of Enzyme Concentration. HPr was isolated as described by Van Dijk et al. (1991) and EI was isolated according to Robillard et al. (1979). Enzyme concentrations were determined by the pyruvate burst method, described by Robillard and Blawuw (1987).

Gel Electrophoresis. Polyacrylamide gel electrophoresis was performed as described by Laemmli (1970) with 12.5% acrylamide gels. Gels were stained by Coomassie staining or by silver staining (Wray et al., 1981).

Preparation of IIA<sup>mtl</sup> and PIIA<sup>mtl</sup> Samples for NMR Measurements. IIA<sup>mtl</sup> (2–4 mM) solutions were brought to the appropriate pH by dialyzing 0.5 mL of the solution against 20 mL of KPi buffer (50 mM in D<sub>2</sub>O, 2 mM EDTA, 1 mM Na<sub>2</sub>N<sub>3</sub>) at the desired pH. The pH was measured before and after each series of spectra, and values agreed to within 0.05 unit. PIIA<sup>mtl</sup> was generated in situ in the NMR tube by first dialyzing the IIA<sup>mtl</sup> solution against 5 mL of D<sub>2</sub>O buffer containing 50 mM KPi, 500 mM PEP, 5 mM MgCl<sub>2</sub>, 5 mM EDTA, 1 mM DTT, and 1 mM Na<sub>2</sub>N<sub>3</sub>. After 2 h of dialysis, 50 µL of a solution containing EI (250 µM) and HPr (360 µM) was added, resulting in the rapid phosphorylation of IIA<sup>mtl</sup>. NMR measurements were started 10 min after the addition of EI and HPr. The presence of characteristic PEP proton resonances in the NMR spectra of PIIA<sup>mtl</sup> confirmed the presence of excess PEP throughout the NMR measurement.

NMR Measurements. All NMR spectra were recorded at 26 °C with a VARIAN VX500 NMR spectrometer, operating at a 1H frequency of 500 MHz and equipped with a SUN 3/160 computer. 1H[15N] heteronuclear multiple quantum coherence (HMQC) spectra were recorded using the pulse sequence described by Bendlall et al. (1983). Spectra were recorded with ω1[15N] = 10 000 Hz and ω2(1H) = 6666 Hz, with 256 increments in ω1. For each ω1 value 128, 192, or 256 transients were recorded, the number of scans depending on the experiment. The acquisition time in t2 was 0.154 s using 2048 data points. The J-value was set to 30 Hz, which in pilot experiments proved to give the best signal to noise ratio. During acquisition, the 15N was decoupled using a broadband WALTZ decoupling. Spectra were recorded using time-proportional phase incrementation (TPPI) to allow for the discrimination between positive and negative frequencies in ω1 and ω2. Spectra with different ω1 were stored separately to allow for small corrections in phase and amplitude after Fourier transformation in t2 (1H). Data processing was performed on a Convex C1-XP computer, using the program SNARF developed in our lab by F. van Hoesel. FIDs were Fourier transformed using a Lorentzian to Gaussian filter in both time domains and a Hanning window in t1 (15N) and sometimes in t2 (1H). A simple baseline correction was performed in t1 after Fourier transformation. 1H chemical shifts are referenced to internal 2,2-dimethyl-2-silapentane-1-sulfonate (DSS); 15N chemical shifts (absolute value) are relative to external 1H[15NO]<sub>3</sub>.

Appearance and Interpretation of 1H[15N] HMQC Spectra. 15N NMR provides insight into the hydrogen bond structure at imidazole ring systems. On the basis of a series of model studies, Bachovchin (1986) proposed a division of imidazole nitrogens into three types, α, α+, and β. They have characteristic chemical shifts, reflecting the protonation state of the individual nitrogens and the entire ring (see Table I). Hydrogen bond interactions at the imidazole nitrogens also influence the 15N chemical shift positions. The α-type and α+-type nuclei can function as H-donors; the β-type can function as H-acceptors using their lone electron pair. These interactions result in small but distinct changes in the resonance positions of the imidazole nitrogens, the α or α+ moving maximally 10 ppm downfield, the β moving maximally 10 ppm upfield, (Bachovchin, 1986). These effects are also given in Table I. This classification of 15N NMR chemical shifts has been used to characterize the protonation and hydrogen bonding state of histidine residues in a number of proteins (Bachovchin & Roberts, 1978; Bachovchin, 1985, 1986; Van Dijk et al., 1990). In this study, 1H[15N] HMQC spectra of the histidine residues in IIA<sup>mtl</sup> were recorded. The J<sub>NNH</sub> couplings of the N61 and N2 nuclei to the C2 and C4 protons allow one to determine the resonance frequencies of all four nuclei in a single HMQC spectrum. The protons that are directly attached to the nitrogens are not observed because they exchange with water too quickly to be detected. When dealing with a neutral imidazole ring system, two extreme situations can occur. The first is when the hydrogen is firmly located on one of the nitrogen nuclei, resulting in an imidazole ring with an α-type and a β-type nitrogen. One then observes a triangle of cross-peaks in the HMQC spectrum as in Figure 1a with the C2H proton reporting two 15N frequencies, one at 210 ppm corresponding to an α-type nitrogen and one at 128 ppm corresponding to a β-type nitrogen. The C4H reports only one nitrogen frequency corresponding to a β-type nitrogen. The coupling between the N61 and C4H is too small (1–2 Hz) to result in an observable cross-peak. The cross-peak pattern enables the unambiguous assignment of the nitrogen and proton frequencies to specific nuclei and at the same time specifies which nitrogen is protonated. In Figure 1a, the N61 is protonated; if the N2 had been protonated, the C4H cross-peak would have been located at an 15N frequency of 210 ppm.

The second extreme occurs when the proton is not located on one of the nitrogens but exchanges rapidly between the two. If the proton shows no preference for the N61 or N2, the nitrogens are magnetically equivalent and resonate at the same frequency of 169 ppm, the average between the α and β resonance position, resulting in a HMQC spectrum with two cross-peaks as in Figure 1b. Intermediate situations exist, for example, when there is a slow tautomeration as shown in Figure 1c. This results in four cross-peaks instead of three. Both the N61 and the N2 have a character part of the time and β character the rest of the time. When the N61 has α character, it resonates at the α position of 210 ppm; when it has β character it resonates at the β position of 128 ppm. The same holds for the N2 nucleus. Consequently, both the C2H

| Table I: 15N Chemical Shift Positions of Imidazole Nitrogens in the NMR Spectrum, As Described by Bachovchin (1986) |
|---|---|---|
| 15N type | chemical shift position (ppm) | maximal H-bond effect (ppm) |
| α | 210 | -10 |
| α+ | 201 | -10 |
| β | 128 | +10 |

* Chemical shift positions are relative to external 1 H[15NO]<sub>3</sub> (absolute values).
Characterization of Histidine Residues in IIA\textsuperscript{mut}

RESULTS

\textsuperscript{1}H\{\textsuperscript{15}N\} HMQC spectra of [\textsuperscript{\delta1,\textepsilon2,\textsuperscript{15}N] histidine-labeled IIA\textsuperscript{mut} recorded at several pH values provided the data for the titration curves for the \textsuperscript{1}H and \textsuperscript{15}N nuclei in the histidine imidazole rings. The data for the \textsuperscript{1}H titration curves were extended with 1D \textsuperscript{1}H NMR measurements at several pH values using a nonlabeled IIA\textsuperscript{mut} preparation.

Relaxation Behavior of the Histidine Resonances

Figure 2 shows the HMQC spectrum for IIA\textsuperscript{mut} at pH = 7.5. The three histidine spin systems are indicated by the lines drawn. The HisB spin system shows fast tautomeration as is evident from its nitrogen chemical shift positions (184 ppm and 191 ppm for N\textepsilon1 and N\textepsilon2, respectively); it is comparable to the situation depicted in the model spectrum in Figure 1d. The other two histidine spin systems clearly show a proton localization at the N\textepsilon1 (HisC) or the N\textepsilon2 (HisA). In addition to different tautomeration states, the histidines also show very different relaxation behavior. This is evidenced by the intensity of the signals of the resonances in the HMQC spectrum. The resonances of HisA are the weakest in the spectrum; the ones of HisC are stronger but they are still much less intense than the signals of HisB. The intensity of a cross-peak in the HMQC spectrum is defined by two processes, the buildup of the signal via the J-coupling and its decay via relaxation. The intensity of the cross-peaks in the absence of any relaxation is described by eq 1, using the product operator notation of Soerensen et al. (1983).

\begin{equation}
M(t) = -I_p \sin^2 \left(2 \pi J \tau \right) \cos \left(\omega_{15N} \tau \right)
\end{equation}

The 'H titration curves for the three histidine residues in IIA\textsuperscript{mut} are given in Figure 3. The data are taken from 1D \textsuperscript{1}H measurements and \textsuperscript{1}H\{\textsuperscript{15}N\} HMQC spectra. Assignment of the resonances was done by using the HMQC spectra, as has been discussed. The data points for each resonance were fitted to eq 2, using a least-squares fitting procedure. \textit{K}_a is the acid dissociation constant, \delta_{obs} is the observed chemical shift position, \delta_a and \delta_b are the chemical shift positions of the protons in the fully protonated and deprotonated imidazole ring respectively, and \textit{x} is the \textit{H}^+ concentration (moles per liter). The solid lines in the figure result from the fit. The results of the analysis are summarized in Table II. The \textit{pK}_a values for HisA could not be determined because when the HisA starts titrating at pH = 5.8, its resonances broaden and become undetectable; at lower pH values the protein also rapidly precipitates. HisB shows normal titration behavior but HisC is peculiar in that its C4H shows almost no titration behavior at all (Table II). It titrates only 0.04 ppm whereas 0.4 ppm is normal for such a proton.
FIGURE 2: $^1$H($^{15}$N) HMQC spectrum of II$^{\text{m}}$ (3 mM, pH = 7.4) selectively enriched with [81,c2-$^{15}$N]histidine. Our interpretation of the spectra in terms of the dominant tautomeric state of histidine residues A, B, and C is shown at the top of the figure.

Table II: Characterization of the Histidine Residues in II$^{\text{m}}$ with Respect to Their $^1$H Titration Parameters

<table>
<thead>
<tr>
<th>Residue</th>
<th>Residue</th>
<th>pK</th>
<th>$\delta_\text{H}$ (ppm)</th>
<th>$\delta_\text{N}$ (ppm)</th>
<th>$\Delta\delta$ (ppm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HisA</td>
<td>C2H</td>
<td>=5.80</td>
<td>7.58</td>
<td>n.o.</td>
<td>&gt;8.50</td>
</tr>
<tr>
<td></td>
<td>C4H</td>
<td>=5.80</td>
<td>6.82</td>
<td>n.o.</td>
<td>nd</td>
</tr>
<tr>
<td></td>
<td>N81</td>
<td>=5.80</td>
<td>129</td>
<td>n.o.</td>
<td>nd</td>
</tr>
<tr>
<td></td>
<td>N82</td>
<td>=5.80</td>
<td>209</td>
<td>n.o.</td>
<td>nd</td>
</tr>
<tr>
<td>HisB</td>
<td>C2H</td>
<td>7.30</td>
<td>7.75</td>
<td>8.72</td>
<td>0.97</td>
</tr>
<tr>
<td></td>
<td>C4H</td>
<td>7.31</td>
<td>7.22</td>
<td>7.63</td>
<td>0.41</td>
</tr>
<tr>
<td></td>
<td>N81</td>
<td>7.32</td>
<td>201</td>
<td>169</td>
<td>32</td>
</tr>
<tr>
<td></td>
<td>N82</td>
<td>7.30</td>
<td>203</td>
<td>178</td>
<td>25</td>
</tr>
<tr>
<td>HisC</td>
<td>C2H</td>
<td>6.58</td>
<td>7.62</td>
<td>8.64</td>
<td>1.07</td>
</tr>
<tr>
<td></td>
<td>C4H</td>
<td>nd</td>
<td>6.88</td>
<td>6.92</td>
<td>0.04</td>
</tr>
<tr>
<td></td>
<td>N81</td>
<td>nd</td>
<td>205</td>
<td>n.o.</td>
<td>nd</td>
</tr>
<tr>
<td></td>
<td>N82</td>
<td>nd</td>
<td>131</td>
<td>n.o.</td>
<td>nd</td>
</tr>
</tbody>
</table>

*The parameters were obtained from the experimental data by a least-squares fitting procedure as is described in the text. n.o. = not observed; nd = not determined.

Apparently, the C4H experiences pH-dependent changes in its environment which influence its titration behavior.

$^{15}$N Titration Curves. HisA: The $^{15}$N titration curve for HisA is shown in Figure 4. For reasons described in the previous section, the titration curve for this residue is incomplete. Consequently, the hydrogen bond structure at this residue can only be determined for the deprotonated imidazole ring. For this ring, two nitrogen resonances are observed, one at 129 ppm and one at 209 ppm. From the cross-peak pattern, Figure 2, it is clear that the proton is located at the N81 position. Neither nitrogen shows significant hydrogen bonding since they both resonate at positions which are very close to the ones expected for pure type $\alpha$ and $\beta$ nuclei, 210 and 128 ppm, respectively.

HisB: This histidine shows an imidazole ring with a fast tautomeration in the neutral state. The titration curves are shown in Figure 4b. In the protonated state, both nitrogens resonate at high $\delta$-values (201 and 203 ppm), indicating two type $\alpha$ nitrogens not significantly involved in hydrogen bond formation. In the neutral imidazole ring, the N81 is found at 169 ppm and the N82 at 178 ppm, the center between these two being at 174 ppm. The assignment of N81 and N82 is based on the cross-peak intensity pattern. The C2H shows cross-peaks to both N81 and N82; tautomeration will result in cross-peaks with more or less equal intensities, assuming that all nuclei show similar relaxation behaviors. The C4H shows a cross-peak to only N82; in a tautomeric equilibrium, the intensity of this cross-peak will be divided over two positions so that its intensity will always be lower than that of the corresponding C2H cross-peaks. The intensity of the C4H cross-peaks reflects the preferred protonation state of the N82 nucleus, which in HisB is the protonated state. This is reflected
in the higher intensity of the cross-peak at a higher field (Figure 2). On the basis of the 9 ppm chemical shift difference between N61 and Nε2, an equilibrium value \( K = N(61)/N(ε2) \) of 0.78 was calculated for the tautomeric process in the neutral imidazole ring. The calculation was performed as described in the previous section, using a δ-value for the β nucleus of 138 ppm instead of 128 ppm because of hydrogen bonding effects. Despite the rapid tautomization, there is an indication that the neutral imidazole ring of HisB is hydrogen bonded. This can be derived from a center position of 174 ppm between N61 and Nε2. If no hydrogen bond were present, this average position would be 169 ppm, the middle between the α-type and β-type nitrogen resonance positions. A higher average value means that either the α-type nitrogen resonance position has moved to values higher than 210 ppm or the β-type nitrogen position has moved to values higher than 128 ppm or that both positions have shifted to higher values. A higher value for the α-type nitrogen position cannot be explained in terms of H-bond formation. Hydrogen bonding would only move the resonance position to lower values (Table I), and this value would explain the position of the center between N61 and Nε2 at 174 ppm. This indicates that the deprotonated nitrogen of both tautomers is acting as a hydrogen bond acceptor. Such a situation requires that the hydrogen bond donating group be in close proximity to both imidazole nitrogens. This would be possible if the ring rotated rapidly so that the N61 and Nε2 were pointing toward the H-bond donor. Rapid rotation is in agreement with the slow relaxation observed for this histidine. A fast tautomization on the \(^{15}\text{N}\) NMR time scale implies an exchange rate constant higher than about \(1 \times 10^5 \text{ s}^{-1}\) (calculated using a chemical shift difference between the two nitrogens of 72 ppm). Rate constants of up to \(10^9 \text{ s}^{-1}\) have been reported for water molecules that were rather strongly hydrogen bonded to the deprotonated nitrogen in imidazole (Ralph & Grunwald, 1969). High exchange rates are also evidenced by the fact that the resonances of this histidine are observed over the complete titration range. Sudmeier et al. (1980) showed from theoretical considerations that the histidine nitrogen resonances are expected to broaden strongly during a titration at the high fields used in the present study. This should lead to the disappearance of the \(^{15}\text{N}\) resonances from the NMR spectrum. The fact that the resonances are observed during the complete titration indicates that the exchange of protons
is catalyzed. In the case of HisB, this can be done by the
H-bond donating group at the imidazole ring or by buffer
ions. Similar fast exchange equilibria have been reported
before (van Dijk et al., 1990).

HisC: Only a partial $^{15}$N titration curve could be measured
for this residue. (Figure 4c). Since the proton titration could
be completely followed over the entire pH range and no
significant line broadening was observed, the disappearance
of the HMQC cross-peaks for HisC must be caused by line
broadening of the nitrogen resonances. As just described,
line broadening is expected for $^{15}$N NMR titrations of histidine
residues with normal protonation rates due to the 73 ppm
line broadening is expected for $^{15}$N NMR titrations of histidine
residues. Similar fast exchange equilibria have been reported
before (van Dijk et al., 1990). However, these considerations do
not explain the disappearance of the $\alpha^+\rightarrow\alpha$-type resonance
unless very slow (de)protonation rates or an alteration in the
tautomeric equilibrium as a function of pH is involved.
Observation of the $^{15}$N resonances at high pH only allows the
hydrogen bond structure to be determined for the neutral
imidazole ring. The N$\delta$1 resonates at 205 ppm, which is 5
ppm lower than expected for a pure type $\alpha$ nitrogen (Table
I) and indicates the involvement of this nitrogen in a hydrogen
bond of intermediate strength. The N$\epsilon$2, resonating at 131
ppm, may be involved in weak hydrogen bond interactions,
since it resonates 3 ppm higher than expected for a pure type
$\beta$ nitrogen (Table I). The hydrogen bond characterization
is summarized in Table III, part a.

**Alternative Conformation for One of the Histidine
Residues**

The HMQC spectrum of some preparations showed four
instead of three histidine spin systems (see Figure 5). As
there are only three histidines present in IIA$^{mut}$, this must
mean that one of the histidines occurs in two alternate states.
The occurrence of two states for a histidine residue is not
uncommon, and it has been described for the active site
histidine in ribonuclease T1 (Schmidt et al., 1991) and for the
histidine residues in HPr (Van Dijk et al., 1990). The "extra"
histidine spin system has a protonated N$\epsilon$2 nucleus and a
deprotonated N$\delta$1 nucleus. The appearance of this extra spin
system is accompanied by a decreased peak intensity of the
C4H/N$\epsilon$2 cross-peak of HisC (compare Figures 2 and 5). We
will come back to this in the Discussion section.

**$^1H$ Titration Curves for the Histidine Residues in PIIA$^{mut}$**

One-dimensional $^1H$ NMR spectra of PIIA$^{mut}$ were recorded
at several pH values. The assignment of resonances to a
particular histidine was done using the HMQC spectrum for
IIA$^{mut}$ (Figure 6). The assignment of the histidines as HisA,
HisB, or HisC was done by comparison with Figures 2 and
5, and by comparison of the $^1H$ titration data for IIA$^{mut}$ and
PIIA$^{mut}$. Similar cross-peak pattern in the HMQC spectrum
combined with similarities in the $pK_a$ values and proton
chemical shifts led to the assignments, presented in Table II.
The assignment of HisA is not straightforward from this
comparison, but it will be discussed in the next section. The
$^1H$ titration curves for PIIA$^{mut}$ are given in Figure 7 and were
analyzed as described previously by fitting to eq 2. The results
are summarized in Table II. The most dramatic change is
observed for HisA, whose $pK_a$ is raised from below 5.8 to
above 8.6. Above pH 8.6, the resonance of the C2H of HisA
broadens and disappears from the 1D $^1H$ spectrum. The C4H
resonance of this residue could not be unambiguously iden-
tified. The large change in $pK_a$, induced by the phosphory-
lation, make this residue the most likely candidate for the
phosphorylation site, since active site histidine residues in other
PTS proteins show similar changes upon phosphorylation. The
conclusion will be confirmed by the HMQC spectrum of
PIIA$^{mut}$, to be discussed in the next section. HisB shows a
small change in $pK_a$ value (from 7.30 to 7.46); the proton
titration behavior is almost the same as before the phos-
phorylation (Table II). HisC is more affected by the phosphory-
lation than HisB, but the changes are not as dramatic as
for HisA. The $pK_a$ is raised from 6.58 to 7.15 upon phosphorylation,
and the total shift upon titration of the C2H is reduced from 1.07 to 0.43 ppm (Table I); the C4H titration
is unchanged. These results indicate local changes in the
surroundings of HisC when the protein becomes phosphory-
lated.

**$^1H$/$^{15}$N HMQC Spectrum of PIIA$^{mut}$**

A pH of 8.1 was chosen for the HMQC spectrum for PIIA$^{mut}$
(Figure 6) because the hydrolysis of the phosphoprotein is not
too high at this pH, allowing for sufficient data acquisition.
The phosphorylation of the protein was checked at several
time intervals during the data acquisition by looking at
characteristic frequencies in the resulting 2D spectra, indi-
cating dephosphorylation.

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### Table III: Characterization of the Hydrogen Bonding State of the Histidine Residues in IIA$^{mut}$ before Phosphorylation and Changes in the Hydrogen Bonding State upon Phosphorylation of IIA$^{mut}$, Performed at pH = 8.1

<table>
<thead>
<tr>
<th>Residue</th>
<th>$pK_a$</th>
<th>$^{15}$N Resonance</th>
<th>Hydrogen Bond State</th>
<th>$pK_a$</th>
<th>$^{15}$N Resonance</th>
<th>Hydrogen Bond State</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>a. Characterization before Phosphorylation</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HisA ($pK_a &lt; 5.80$)</td>
<td>N$\delta$1</td>
<td>type $\beta$, no hydrogen bond</td>
<td>not observed</td>
<td>N$\delta$1</td>
<td>no hydrogen bond</td>
<td>not observed</td>
</tr>
<tr>
<td></td>
<td>N$\epsilon$2</td>
<td>type $\alpha$, no hydrogen bond</td>
<td>not observed</td>
<td>N$\epsilon$2</td>
<td>no hydrogen bond</td>
<td>not observed</td>
</tr>
<tr>
<td>HisB ($pK_a = 7.30$)</td>
<td>N$\delta$1</td>
<td>fast tautomeration, proton preference for N$\epsilon$2, strong</td>
<td>type $\alpha^+$, no hydrogen bond</td>
<td>N$\delta$1</td>
<td>fast tautomeration, strong</td>
<td>type $\alpha^+$, no hydrogen bond</td>
</tr>
<tr>
<td></td>
<td>N$\epsilon$2</td>
<td>H bond to the $\beta$-type nitrogen</td>
<td>type $\alpha^+$, no hydrogen bond</td>
<td>N$\epsilon$2</td>
<td>H bond to the $\beta$-type nitrogen</td>
<td>type $\alpha^+$, no hydrogen bond</td>
</tr>
<tr>
<td>HisC ($pK_a = 6.58$)</td>
<td>N$\delta$1</td>
<td>type $\alpha$, moderate hydrogen bond</td>
<td>not observed</td>
<td>N$\delta$1</td>
<td>type $\alpha$, moderate hydrogen bond</td>
<td>not observed</td>
</tr>
<tr>
<td></td>
<td>N$\epsilon$2</td>
<td>type $\beta$, no hydrogen bond</td>
<td>not observed</td>
<td>N$\epsilon$2</td>
<td>type $\beta$, no hydrogen bond</td>
<td>not observed</td>
</tr>
<tr>
<td><strong>b. Changes upon Phosphorylation</strong></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>HisA</td>
<td>$pK_a &lt; 5.80$</td>
<td>N$\delta$1</td>
<td>no hydrogen bond</td>
<td>$pK_a &gt; 8.50$</td>
<td>N$\delta$1</td>
<td>no hydrogen bond</td>
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<tr>
<td></td>
<td>N$\epsilon$2</td>
<td>no hydrogen bond</td>
<td>N$\epsilon$2</td>
<td>phosphorylated</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HisB</td>
<td>$pK_a = 7.30$</td>
<td>N$\delta$1</td>
<td>fast tautomeration, strong hydrogen bond</td>
<td>$pK_a = 7.47$</td>
<td>N$\delta$1</td>
<td>fast tautomeration, strong hydrogen bond</td>
</tr>
<tr>
<td></td>
<td>N$\epsilon$2</td>
<td>strong hydrogen bond</td>
<td>N$\epsilon$2</td>
<td>no hydrogen bond</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HisC</td>
<td>$pK_a = 6.70$</td>
<td>N$\delta$1</td>
<td>moderate hydrogen bond</td>
<td>$pK_a = 7.15$</td>
<td>N$\delta$1</td>
<td>moderate hydrogen bond</td>
</tr>
<tr>
<td></td>
<td>N$\epsilon$2</td>
<td>no hydrogen bond</td>
<td>N$\epsilon$2</td>
<td>no hydrogen bond</td>
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</table>
HisA: This residue is assigned as is indicated in Figure 6. No resonances are observed at the positions where HisA resonances occurred in the nonphosphorylated protein. The spin system, indicated in Figure 6 as HisA was also observed in the 1D $^1$H NMR spectra. The N61 resonance of this spin system is found at 202 ppm, indicative of an $\alpha^+$-type nitrogen, or an $\alpha$-type nitrogen, involved in strong H-bond formation. As the pKa of this residue is above pKa $= 8.6$ (Table I), the observed nitrogen obviously is $\alpha^+$-type and is not involved in H-bond formation. The Nc2 of HisA is found at 166 ppm, a value typical for a phosphorylated imidazole nitrogen (Van Dijk et al., 1990). Because the Nc2 resonances are not intense in the contour plot, the $^1$H trace at this $^{15}$N frequency is plotted in Figure 6b, and the relevant resonances are indicated. The data thus indicate that HisA is the active site histidine and that phosphorylation takes place at the Nc2 position.

HisB: This residue was easily identified in the spectrum by its appearance and by its $^1$H titration parameters, which are almost unaffected by the phosphorylation. A comparison of Figure 6 with Figures 2 and 5 shows, however, that the tautomeric equilibrium of this residue is affected by the phosphorylation process. Before phosphorylation, four resonances were observed for this residue, indicative of a fast tautomerization with the proton having a slight preference for the Nc2 position. In Figure 6 only two cross-peaks are observed for this residue at $\delta_{15N} = 175$ ppm, one at each proton frequency. The peaks appear broad in the $^{13}$N domain because the plot is contoured at low levels to detect the HisA and HisC cross-peaks. However, when the nitrogen trace at the C2H frequency is plotted, we clearly see only one resonance (Figure 6c). The HMQC spectrum of HisB in Figure 6a is analogous to the model spectrum in Figure 1b; they are both representative of a histidine showing a fast tautomerization with the proton being equally distributed over the two nitrogen positions. Furthermore, the resonance position at 175 ppm indicates that HisB is no longer hydrogen bonded. Calculations, using eq 2 and a pKa of 7.47 for HisB in IIA$^{\text{mli}}$ (Table II) give a theoretical chemical shift position of 175.1 ppm at $pH = 8.1$, the same as the observed resonance position. Before phosphorylation, there was a strong hydrogen bond at this residue which is apparently disturbed when the protein becomes phosphorylated.

HisC: The $^1$H titration curves for this residue show that it is somewhat affected by the phosphorylation process (Table II). For clarity, we show the proton trace at the Nc2 frequency (133 ppm, Figure 6d) to indicate the expectation of the Nc2/C4H cross-peak, which is not observable at the contour level used in Figure 6a and only barely visible in Figure 5. This is the position found for the C4H in the nonphosphorylated protein (see Table II). The histidine still shows the same proton localization at the N61 position, but its $^{15}$N resonances at 206 and 133 ppm are at positions which are close to the ones calculated (204 and 134 ppm, respectively) for a non-hydrogen-bonded imidazole ring using the estimated pKa for this residue (Table II). While there was a moderate hydrogen bond to the N61 in IIA$^{\text{mli}}$, phosphorylation disturbs the H-bond at HisC just as it did at HisB. The interpretations of the hydrogen bond changes are summarized in Table III, part b.

Remaining Cross-Peaks. It is clear from Figure 6 that more cross-peaks are present in the spectrum than expected. One of them is the "extra" spin system that was already observed in Figure 5 and which we interpreted as an alternative conformation of HisA, HisB, or HisC. Then, there remain some cross-peaks which all have a more or less diffuse appearance. They are found at positions that do not correspond with previously observed positions for HisA, HisB, or HisC. These cross-peaks may originate from IIA$^{\text{mli}}$ that cannot be properly phosphorylated. Gel filtration analysis showed the
FIGURE 6: (a) $^1$H$^{15}$N-HMQC spectrum of PIIA. The dominant protonation state of the histidine residues is shown at the top of the figure. (b) Proton trace from panel a at $\delta_{1H} = 166$ ppm. (c) Nitrogen trace from panel a at $\delta_{1H} = 7.87$ ppm. (d) Proton trace from panel a at $\delta_{1H} = 133$ ppm.

FIGURE 7: $^1$H titration curves for the histidine residues in PIIA$^{\text{dil}}$. Symbols: ($\Delta$) C2H (HisA), (O) C2H (HisB), (>) C4H (HisB), (X) C2H (HisC), (V) C4H (HisC). The C4H of HisA could not be unambiguously identified.

The presence of some dimeric and higher oligomeric forms of the protein which we showed to be inactive. At present, however, we cannot unambiguously explain the origin of the diffuse cross-peaks.

DISCUSSION

We have described the characterization of the three histidine residues in IIA$^{\text{dil}}$ with respect to their protonation state and their hydrogen-bonding interactions using 1D $^1$H NMR and $^1$H$^{15}$N-HMQC spectra. The $^{15}$N-chemical shift data provide the information necessary to deduce the hydrogen bond interactions with the histidine imidazole rings. The data are summarized in Tables II and III. HisA proved to be the active site residue, and thus is equivalent to His554 in the intact EIImtl. The phosphorylation takes place at the Nε2 position of the imidazole ring, which is analogous to that observed for other proteins with a function similar to that of IIA$^{\text{dil}}$ (Deutscher et al., 1982a; Dörschug et al., 1984).
increase in pKₐ of this residue upon phosphorylation is similar to what has been observed for the active site residues in other PTS proteins like HPr, IIA⁶C, and IIA⁸C, be it that the shift in the case of IIA⁶C is very large (from below 5.8 to above 8.6). There are several points to be stressed concerning the active site histidine. First, the imidazole ring is not hydrogen bonded in the phosphorylated or dephosphorylated state. Similar studies on HPr (Van Dijk et al., 1990) showed hydrogen bonding to the imidazole ring of the active site histidine that seemed to maintain the imidazole ring in a fixed position. The ring orientation was thought to be important for the phosphorylation of HPr, and we expected that a similar situation would occur in IIA⁶C. Second, there is a change in tautomeric state upon phosphorylation. The proton is situated at the Ne2 position in the nonphosphorylated protein (at pH > 5.8). This is also the place of attachment of the phosphorous group. One would expect that the site of phosphorylation would be deprotonated so that the electrophilic phosphorous group would be easily directed to the nucleophilic deprotonated imidazole nitrogen. This has been found for HPr (Van Dijk et al., 1990). After phosphorylation, the imidazole ring remains protonated as witnessed by its high pKₐ (>8.6). Consequently, the proton shifts before or during the phosphorylation process from the Ne2 to the N6₁ position. It is likely that the PHPr/IIA⁶C complex formed prior to phosphoryl transfer induces the change in the tautomeric state expected for an efficient phosphorylation reaction pathway. Evidence for such a PHPr/IIA⁶C complex has been found in 3¹P NMR studies and will be reported elsewhere.

The phosphorylation process affects not only the active site residue, but also the other two histidine residues in IIA⁶C. Similar effects have been observed for the histidine residues in IIA⁶C (Kalbitzer et al., 1981). Both are hydrogen bonded in the nonphosphorylated protein, but not in the phosphorylated IIA⁶C. Furthermore, changes in pKₐ for both histidines and changes in the tautomization state of HisB were observed, indicating that the phosphorylation induces overall conformational changes in the protein, a suggestion that is supported by the observation of a number of changes in the aliphatic region of the ¹H NMR spectrum (data not shown). In this respect, IIA⁶C behaves like IIA⁶C, IIA⁸C, and IIA⁸C which also show overall conformational changes upon phosphorylation (Dörschug et al., 1984; London & Hausman, 1983; Deutscher et al., 1982b). Our data suggest that one of the histidine residues experiences an alternative conformation. This has been assigned to HisC, since it is the only residue that shows a change in its relaxation behavior when the alternative histidine appears in the spectrum (compare Figures 2 and 5). High-resolution NMR studies are now in progress which will reveal the sequence assignments of HisB and HisC and hopefully provide a structural basis for understanding the significance, if any, of the alternative conformation observed for HisC.

The first high-resolution structures of IIA domains, IIA⁶C from *Bacillus subtilis* and *Escherichia coli*, have recently been published (Liao et al., 1991; Worthylake et al., 1991). Since we expect IIA domains to function similarly, it is worth examining the data presented here in light of the X-ray structures of IIA⁶C. It is essential to note, however, that there is no significant homology between the primary structures of IIA⁶C and IIA⁶C from either source. Furthermore, the tertiary structures of IIA⁶C will probably be very different from those of the IIA⁶C since FTIR data in H₂O and D₂O show approximately 45% α-helix and 27% β-sheet for IIA⁶C from *E. coli* where both IIA⁶C structures are dominated by β-sheet. The following comparison is based on the assumption that the arrangement of residues at the active site of IIA⁶C is similar to that of the two IIA⁶C structures. The *B. subtilis* IIA⁶C possesses two histidines in the active site region: His83 which becomes phosphorylated and His68. HisA in IIA⁶C corresponds to His83. Since HisB appears to be situated on the exterior of the protein and since HisC is more strongly affected by the phosphorylation, HisC probably corresponds to His68. Both *B. subtilis* IIA⁶C histidines are associated with phenylalanine residues, and these are thought to be important for the relative orientation of the two histidine imidazole rings; no clear hydrogen bonds at the imidazole rings have been observed, although Liao et al. (1991) point out that there are indications for the presence of such structures. The absence of hydrogen bonds at the active site His83 in IIA⁶C would be similar to what we observed for the HisA in IIA⁶C. Furthermore, the presence of a phenylalanine near HisC, the other suggested histidine in the active site of IIA⁶C, could explain the abnormally small pH-dependent shift of the C4H of this residue. A change in ring current shift due to movement of the two residues relative to one another upon (de)protonation is all that is necessary. In their model of the phosphorylated IIA⁶C, Liao et al. (1991) suggest that the phosphoryl oxygen atoms would interact with the N42 of His68 while the N6₁ of this residue would hydrogen bond to a side chain oxygen of a threonine residue. This is contrary to our finding that neither nitrogen atom of HisC in IIA⁶C is hydrogen bonded in the phosphorylated enzyme. The weak intensity of the N42/C4H cross-peak for HisC in IIA⁶C may arise from intermediate exchange dynamics with other species or intermediates which we cannot detect. Finally, the N42 of HisC is a β-type nitrogen (i.e., not protonated) and not able to serve as a H-bond donor to the phosphoryl oxygens.

In the case of IIA⁶C from *E. coli*, Worthylake et al. (1991) report that the N6₁ nitrogens of both histidines at the active site are hydrogen bonded; that of His90, the phosphorylation site, to the carbonyl oxygen of Gly92 (2.7 Å) and that of His71 to Oγ₁ of Thr73 (3.1 Å). His90 would correspond to our HisA. As summarized in part a of Table III, the N6₁ of HisA is not protonated at neutral pH, in contrast to the His90 N6₁ in IIA⁶C, and it is not involved in hydrogen bond formation. The N6₁ of HisC which may correspond to His71 is hydrogen bonded.

In conclusion, there are some similarities between the proposals for the active site configuration arising from the X-ray data for IIA⁶C from *B. subtilis* or *E. coli* and the NMR data for IIA⁶C from *E. coli* in the unphosphorylated enzymes, but there is not full agreement. Furthermore, the proposals for the phosphorylated domains of IIA⁶C versus those of *B. subtilis* IIA⁶C differ significantly. All of these differences may reflect differences in the atomic details at the phosphorylation sites of the three enzymes if not differences in phosphorylation mechanism itself.

Presper et al. (1989) proposed a possible role for a second histidine as a migration site from the phosphoryl group on its way to the IIB domain. The finding, in both crystal structures, of two histidines with their ε₂ nitrogens close enough together to accommodate such a transition is enticing. But again, the present NMR data do not support such a proposal for IIA⁶C; phosphorylation dramatically affects only one histidine, HisA. Neither of the other two histidines is that strongly affected. It is possible, however, that such a migration will only occur when stoichiometric amounts of PHPr or the B domain are present. Experiments are in progress to test these suggestions.
REFERENCES


Registry No. His, 71-00-1.