Structural insights into the binding of cardiac glycosides to the digitalis receptor revealed by solid-state NMR

David A. Middleton*,†, Saffron Rankin*, Mikael Esmann§, and Anthony Watts*

†Department of Biochemistry, University of Oxford, South Parks Road, Oxford OX1 3QU, United Kingdom; and ‡Department of Biophysics, University of Aarhus, Ole Worms Alle 185, Universitetsparken, DK-8000 Aarhus C, Denmark

Communicated by Jens C. Skou, University of Aarhus, Aarhus C, Denmark, October 2, 2000 (received for review)

Several biologically active derivatives of the cardiotonic steroid ouabain have been made containing NMR isotopes (13C, 2H, and 19F) in the rhamnose sugar and steroid moieties, and examined at the digitalis receptor site of renal Na+/K+-ATPase by a combination of solid-state NMR methods. Deuteration NMR spectra of H-labeled inhibitors revealed that the sugar group was only loosely associated with the binding site, whereas the steroid group was more constrained, probably because of hydrogen bonding to residues around the K+-channel region. Crosspolarization magic-angle spinning NMR showed that chemical shifts of inhibitors 13C-labeled in the sugar group moved downfield by 0.5 ppm after binding to the digitalis site, suggesting that the sugar was close to aromatic side groups. 19F, 13C-rotational-echo double-resonance NMR strategy was used to determine the structure of an inhibitor in the digitalis receptor site, and it showed that the ouabain derivatives adopt a conformation in which the sugar extends out of the plane of the steroid ring system. The combined structural and dynamic information favors a model for inhibition in which the ouabain analogues lie across the surface of the Na+/K+-ATPase α-subunit with the sugar group facing away from the surface of the membrane but free to move into contact with one or more aromatic residues.

The Na+/K+-ATPase is found in the cells of higher eukaryotic organisms and is the well-known pharmacological receptor for digitalis compounds, including the cardiac glycosides (1, 2). Cardiac glycosides such as ouabain are plant-derived steroids that bind to the extracellular face of the Na+/K+-ATPase α-subunit with high affinity (kd = 5 nM) and remarkable selectivity, and thereby inhibit nucleotide hydrolysis and ion transport (3). Notwithstanding their toxicity, low levels of digitalis compounds have been used as inotropic drugs for over 200 years and remain a clinically recommended therapy for first-line treatment of congestive heart failure and cardiac arrhythmia.

The Na+/K+-ATPase exchanges three cytosolic Na+ for two extracellular K+, which is facilitated by the 1024-residue catalytic α-subunit and involves what are probably large-scale conformational changes of the protein, exposing, in turn, active sites to ions on each side of the membrane (4, 5). Results of topological analysis suggest that the Na+/K+-ATPase α-subunit contains 10 membrane-spanning regions, H1–H10, which is substantiated by the two-dimensional electron density maps of the homologous Ca2+-ATPase and Neurospora H+-ATPase (6, 7). Affinity labeling experiments and site-directed mutagenesis studies have given information about the digitalis receptor site. These have shown that ouabain interacts with both the N-terminal and C-terminal regions of the α-subunit (8–10) and suggest that the drug may exclude K+ from the ligation site within the membrane-spanning regions H4, H5, H6, and H8 (11, 12). To date, however, there exists no detailed model for cardiac glycoside binding that reconciles both drug structure and binding location.

Solid-state NMR spectroscopy has emerged in recent years as a powerful technique for examining the structure and dynamics of small molecules at their binding sites within membrane proteins (13, 14), such as retinal in rhodopsin (15) and inhibitors of the H+/K+-ATPase (16). Here, a number of ouabain derivatives containing NMR-visible reporter groups have been tested for inhibitory activity against renal Na+/K+-ATPase and examined at the digitalis binding site by solid-state NMR methods. Ouabain was modified either by forming an acetonide bridge between two hydroxyl groups in the ouabagenin steroid (ouabain monoacetonide, OMA), by forming acetonide bridges in the steroid and rhamnose sugar moieties (ouabain diacetonide, ODA), or by forming a fluorodiacetoni diolone bridge in the steroid and an acetonide bridge in the sugar group (ouabain fluorodiacetoni diolone, OFDA). Ouabain derivatives (I–IX) containing combinations of NMR-visible isotopes (H, 13C, and 19F) in the steroid group, the sugar group, or both are shown in Table 1. Isotope labeling enabled us to examine the molecular conformation, dynamics, and location of cardiac glycoside analogues in the native digitalis receptor site.

Materials and Methods

Materials. Ouabain hexahydrate, fluoroacetone, and [13C]acetone were obtained from Sigma. [3H]Ouabain was purified by chromatography on Na+/K+-ATPase as described by Hansen (17). Na+/K+-ATPase was prepared from pig kidney as described by Jørgensen (18) by using SDS as the activating detergent.

Synthesis of Ouabain Acetonide Derivatives. OMA, ODA, and OFDA derivatives (I–IX) were synthesized from ouabain (Table 1). Ouabain hexahydrate (0.05 g) was dissolved in acetone (1 ml), and copper sulfate (0.2 g) was added. The solution was stirred in a sealed flask in the dark at room temperature. After 2 days, TLC (3:1 ethyl acetate-methanol) confirmed that all of the starting material had reacted and two products had formed in an approximate ratio of 1:1, with Rf values of 0.5 and 0.75. The solution was filtered to remove copper sulfate and the filtrate evaporated to dryness. The mixture was dissolved in methanol and purified by silica chromatography using 3:1 ethyl acetate-methanol as the eluting solvent. The two products were identified as OMA (II) (0.025 g; 58%) and ODA (III) (0.016 g; 35%) by NMR and mass spectrometry.

The deuterated compound [steroid-2H6]ODA (IV) was prepared as described above using 1 ml [2H6]acetone with the
Table 1. Chemical structures and numbering system (I–IX) for the acetonide derivatives of ouabain used in these studies, showing the positions of $^2$H,$^1$H,$^19$F, and $^{13}$C isotopes

<table>
<thead>
<tr>
<th>compound</th>
<th>number</th>
<th>functional groups</th>
</tr>
</thead>
<tbody>
<tr>
<td>monoacetonide (OMA)</td>
<td></td>
<td>R R' X</td>
</tr>
<tr>
<td>I</td>
<td>CH$_3$</td>
<td>C</td>
</tr>
<tr>
<td>II</td>
<td>CH$_3$</td>
<td>$^{13}$C</td>
</tr>
<tr>
<td>diacetonide (ODA)</td>
<td></td>
<td>R R' X</td>
</tr>
<tr>
<td>III</td>
<td>CH$_3$</td>
<td>CH$_3$ C</td>
</tr>
<tr>
<td>IV</td>
<td>CD$_3$</td>
<td>CH$_3$ C</td>
</tr>
<tr>
<td>V</td>
<td>CH$_3$</td>
<td>CD$_3$ C</td>
</tr>
<tr>
<td>VI</td>
<td>CH$_3$</td>
<td>CH$_3$ $^{13}$C</td>
</tr>
<tr>
<td>steroid-fluorodiacetonide (OFDA)</td>
<td></td>
<td>R R' X</td>
</tr>
<tr>
<td>VII</td>
<td>CF$_3$</td>
<td>C</td>
</tr>
<tr>
<td>VIII</td>
<td>CF$_3$</td>
<td>$^{13}$C</td>
</tr>
<tr>
<td>IX</td>
<td>CHF$_2$</td>
<td>$^{13}$C</td>
</tr>
</tbody>
</table>

following modification. When the starting material had all converted to the [steroid-$^2$H$_2$]OMA, the product was isolated and dissolved in a suspension of 0.2 g copper sulfate and 1 ml acetone. The reaction proceeded as above until the diacetonide had fully formed. Synthesis of [rhamnoside-$^2$H$_2$]ODA (V) was carried out similarly, but using acetone as the starting solvent and then dissolving the monoacetonide in $[^2$H$_6$]acetone. Synthesis of the $^{13}$C-labeled OFDA compounds [rhamnoside-$^{13}$C$^{-19}$F$_3$]OFDA (VIII) and [rhamnoside-$^{13}$C$^{-2}$H$_3$OFDA (IX) was carried out as for [steroid-$^2$H$_2$]ODA, substituting fluoroacetone in place of $[^2$H$_6$]acetone and [2-$^{13}$C]acetone in place of acetone.

Na$^+/K^+$-ATPase Activity and Ligand Binding. The specific Na$^+/K^+$-ATPase activity was about 1,700 μmol/mg/h at 37°C (see ref. 19 for details).

The inhibitory potencies of ouabain, OMA (I), ODA (III), and OFDA (VII) were determined as the IC$_{50}$ (the inhibitor concentration causing 50% enzyme inactivation) for ATP hydrolysis. Na$^+/K^+$-ATPase (51 μg protein/ml) was preincubated in incubation medium (3 mM MgCl$_2$, 3 mM Pi, 40 mM Tris, pH 7.3) and 0–100 μM of ouabain or diacetonide derivative. After the incubation at 20°C for 60 min, an aliquot was transferred to hydrolytic assay medium, whereby the enzyme and glycoside was diluted 40-fold and incubated for a further 10 min before determining the liberated Pi.

The selectivity and binding affinity of the ouabain derivatives were determined from competition experiments using [H]$^1$H]ouabain (17). Na$^+/K^+$-ATPase membranes (25 μg protein/ml) were added to incubation medium containing 5–280 nM ouabain with 3,000 Bq/ml [H]$^1$H]ouabain and either I or III to a final concentration of 0–5 μM. The mixture was incubated for 37°C for 3 h and filtered to determine the amount of free ouabain left in the solution. The amount of bound ouabain was calculated as the difference between the radioactivity of the initial incubation medium and the filtrate. Displacement of [H]$^1$H]ouabain by OMA and ODA was followed by incubating Na$^+/K^+$-ATPase (23 μg protein/ml) in incubation medium containing 30 nM [H]$^1$H]ouabain for 60 min at 37°C. Ouabain, OMA, or ODA was added to a final concentration of 50 μM, and the release of bound [H]$^1$H]ouabain was followed for up to 25 h by filtration as described above.

Sample Preparation for NMR. Na$^+/K^+$-ATPase membranes (30 mg protein) were prepared as a pellet by centrifugation at 100,000 × g for 30 min at 4°C, resuspended in 1 ml incubation medium, and incubated with 0–300 μM of labeled inhibitor (II, IV–VI, VIII, and IX) for 60 min at 25°C. It was assumed that 1 mg of total protein contained 2.5–3.0 nmol ouabain binding sites (19). The suspension was centrifuged (100,000 × g at 4°C) for 30 min, and the pellet was transferred to a 6-mm external diameter zirconia MAS rotor fitted with Kel-F inserts to confine the sample to the center of the rotor. In some experiments, the membranes were first prepared as a pellet and the desired concentrations of the ouabain derivative were added.

NMR Experiments. All solid-state NMR experiments were performed on a Varian Infinity spectrometer operating at a magnetic field of 11.7 T at temperatures between 5°C and ~50°C. Wide-line $^2$H NMR experiments were carried out in a probehead fitted with single-tuned 7-turn 5-mm coil, using the quadrupole echo sequence with a pulse length of 3.5 μs. In crosspolarization magic-angle spinning (CP-MAS) experiments, samples were rotated at a MAS frequency ($\nu_m$) of 3–6 kHz. Hartmann–Hahn crosspolarization from $^1$H to $^{13}$C was achieved over a 1-ms contact time at a field of 65 kHz for both nuclei, and protons were decoupled during signal acquisition at a field of 65 kHz. Rotational-echo double-resonance (REDOR) experiments (13C-observe, $^{19}$F-dephase) were conducted at a MAS frequency of 4.5 kHz using a standard pulse sequence (20). The number of rotor cycles ($N_m$) was 8 or 16 at a spinning frequency $\nu_m$ of 4500 Hz, and the $\pi$ pulse length for both $^{19}$F and $^{13}$C frequencies was 9 μs in both cases.

Results and Discussion

Binding and Activity of Ouabain Derivatives. The inhibitory potencies of ouabain and its derivatives were determined from curves of Na$^+/K^+$-ATPase hydrolytic function over a range of drug concentrations (Fig. 1A). Ouabain is a highly potent inhibitor and inactivated the enzyme with an IC$_{50}$ of <0.08 μM, whereas OMA (I), ODA (III), and OFDA (VII) all exhibited IC$_{50}$ values of between 5 and 15 μM. Hence, modification of the steroid group of ouabain led to a reduction in inhibitory potency compared with that of the parent compound, whereas modification of the sugar had little additional effect on inhibitory potency. The acetonide fluoromethyl groups had no extra consequences for activity (Fig. 1A), indicating that the electronegative properties of fluorine were not important characteristics for inhibitor binding. It is known that the sugar group is not involved in the site of action (Fig. 1). The role of the sugar, which alone has no inhibitory effect, is unclear, although it increases the water solubility of cardiac glycosides relative to aglycones and may influence uptake and distribution of the inhibitor.

The selectivity of ouabain derivatives for the digitalis site was influenced by the $^2$H and $^{13}$C isotopes in the acetonide derivatives of ouabain used in these studies, showing the positions of $^2$H,$^1$H,$^19$F, and $^{13}$C isotopes.
and by ODA followed the same time course, a strong indication that the derivative binds to the same site as ouabain in a fully competitive manner. Dissociation constants for the derivatives, calculated from Scatchard plots of [3H]ouabain binding (not shown), were on the order of 600 nM compared with 10 nM for the parent compound. Hence, the derivatives retained considerable inhibitory potency despite having a lower binding affinity than ouabain.

Formation of acetonide bridges across the steroid and rhamnose hydroxyl groups reduces the hydrogen-bonding capacity of the derivatives; moreover, mutation of transmembrane residues having hydrogen-bonding potential has been shown to alter the sensitivity of Na+/K+-ATPase to ouabain binding (22). Hence, the diminished affinity of the ouabain analogues may arise from a reduction in the ability of the steroid group to form hydrogen bonds with conserved residues. Lowering the hydrogen-bonding capacity of the functionally less important rhamnose group appears to have less of an effect on inhibitory potency.

Dynamics of Inhibitors in the Digitalis Site. Information on the binding characteristics of ouabain analogues was obtained from 2H NMR studies on the dynamics of 2H-labeled ODA (compounds IV and V) titrated into Na+/K+-ATPase membranes. In the case of small deuterated ligands bound to large proteins undergoing slow anisotropic reorientation in the membrane, protein rotation alone is insufficient to completely average the 2H quadrupolar interaction, and the 2H spectral line shape is influenced by the local dynamics of the deuterated ligand (e.g., ref. 23).

The 2H NMR spectra of ODA with deuterons placed either in the methyl groups of the steroid acetonide (IV) or in the rhamnose acetonide methyls (V) are shown in Fig. 2. Spectra of powder samples of the two inhibitors were very similar to each other, and their line shapes and quadrupole couplings indicated that rapid unrestricted deuteriomethyl rotation was the dominant motion in the solid state (Fig. 2A). The spectral similarity was lost, however, when IV and V were titrated into the Na+/K+-ATPase membranes (Fig. 2B). The spectrum from the rhamnoside deuterons of V consisted of a single narrow line (Fig. 2B Right), indicating that the 2H quadrupolar interaction was reduced to close to the isotropic limit by molecular motion. The sugar group therefore must interact only very loosely with the digitalis site to be consistent with such a high degree of motional freedom.

By contrast, the 2H NMR spectrum from the steroid deuterons of IV in the membranes exhibited a broad powder line shape superimposed on a narrow central line (Fig. 2B Inset). The broad component disappeared after titrating an excess of ouabain into the membranes (Fig. 2B Left), indicating that the 2H quadrupolar interaction was reduced to close to the isotropic limit by molecular motion. The sugar group therefore must interact only very loosely with the digitalis site to be consistent with such a high degree of motional freedom.

The 2H NMR spectrum from the steroid deuterons of IV in the membranes exhibited a broad powder line shape superimposed on a narrow central line (Fig. 2B). The broad component disappeared after titrating an excess of ouabain into the membranes (Fig. 2B Inset) and hence corresponded to inhibitor in the digitalis site, whereas the central line was attributed to a much smaller fraction of nonbound IV in the aqueous medium. The residual quadrupole splitting of about 17 kHz observed for the bound inhibitor indicates that the steroid fragment of the molecule was much more restrained than the...
resulted from the accumulation of 8192 scans with a repetition rate of 2 s. Titration of a molar equivalence (with respect to the digitalis sites) of [13C]OMA (II) into the Na⁺/K⁺-ATPase membranes was marked in the spectrum by the emergence of a resonance peak at 100.4 ppm (Fig. 3B). From the results of the 2H NMR experiments (Fig. 2), it was expected that over 70% of the peak intensity corresponded to OMA bound to the digitalis site. The resonance frequency of II titrated into the Na⁺/K⁺-ATPase membranes was identical to that of an equal concentration of OMA in buffer solution (Fig. 3C), indicating that binding of the inhibitor did not perturb the electronic distribution around the steroid [13C] acetonide group.

Titration of [13C2]ODA (VI) into the digitalis site was marked by the appearance of two resonance lines, one at 100.4 ppm from the steroid [13C]-acetonide as before, the other at 110.9 ppm from the rhamnose [13C]-acetonite (Fig. 3D). The steroid-[13C] chemical shift from VI in the binding site was again identical to that of the inhibitor alone in buffer solution (Fig. 3F), whereas the rhamnose [13C] peak had moved downfield to 110.9 ppm from its solution chemical shift of 109.1 ppm. After titration of a 2-fold excess of VI into the Na⁺/K⁺-ATPase membranes (Fig. 3E), a new resonance peak from the rhamnose-[13C] emerged at the same position as that for VI in buffer solution (Fig. 3F). Hence, the two peaks were assigned to inhibitor VI free in solution (at 109.4 ppm) and at the digitalis site (110.9 ppm), and the spectra therefore indicated that all of the inhibitor was bound when present in molar equivalence to the binding sites (Fig. 3D).

The downfield shift of the rhamnose-[13C] peak occurring after binding of VI to the digitalis site can be ascribed to the deshielding effects of local electron-withdrawing groups of polar residues, ring currents of aromatic side-chains, or contributions from both mechanisms. It is likely that the polarity of the acetonide substituent on the sugar group is too low to favor an interaction with polar residues, and it is more probable that the sugar group comes into contact with aromatic residues.

**Structures of Inhibitors in the Digitalis Site.** Fluorinated [13C]-labeled OFDA was examined by using the CP-MAS NMR method REDOR (20). Distances between isotope labels are measured by following the loss of signal intensity from the observed nuclear spin (e.g., [13C]) after reintroducing its distance-dependent dipolar coupling with a heteronuclear spin (e.g., 19F). After N2 rotor cycles at a given sample spinning frequency (v), the distance between the observed and the coupled atomic nuclei (up to 10 Å for 19F-[13C]) can be calculated from the fraction of signal remaining (20).

A strategy was used in which fluorinated analogues with a [13C] label placed in the rhamnose moiety, and either a -CF3 group (VIII) or -CH3F group (IX) in the steroid acetonide, were examined at the digitalis receptor site. Comparing the dipolar dephasing of the rhamnose [13C] signal by the combined effect of three equivalent [19F] nuclei and also by the weaker effect of a single [19F] permitted two independent constraints to be placed on the inhibitor binding conformation. In both experiments, the distance separating the rhamnose [13C] and the centroid defining the time-averaged position of the axially rotating fluorine atom(s) is related to two flexible torsional angles (φ1 and φ2) in the molecule (Fig. 4A). All other angles and bond lengths are known, and hence, by measuring the distance-dependent dipolar coupling between [13C] and [19F], it is possible to deduce combinations of φ1 and φ2 that are consistent with the experimental data.

The results of REDOR experiments on VIII and IX in the digitalis site are shown in Fig. 4B. Two [13C] spectra were obtained, one a full-echo spectrum that serves as a control, and the other
a 19F dephased-echo spectrum after reintroduction of the 13C-19F dipolar interaction that carries the distance information. The intensity of the full-echo spectrum divided by the intensity of the dephasaged echo spectrum (S/S0) is proportional to the 13C-19F interatomic distance. In Na+/K+-ATPase membranes containing VIII, which has a CH3F group, the intensities of the full-echo and dephasaged-echo (red) spectra from either VIII (Left) or IX (Right) in the digitalis site. Spectra resulted from the accumulation of 54,000 scans. (C) Third, molecular conformations were determined by comparing the calculated S/S0 values with the experimental values. Torsional angles \( \phi_1 \) and \( \phi_2 \) giving the best fit of the calculated S/S0 to the experimental dephasing are identified from the light-colored regions of contour plots for VIII (Left) and IX (Right).

The molecular conformation of OFDA in the digitalis site was determined by comparing the measured S/S0 with values computed (28) for combinations of angles \( \phi_1 \) and \( \phi_2 \) covering the entire conformational space of the sugar group (Fig. 4). Combinations of angles consistent with the observed dephasing for VIII and IX in the digitalis site are given by the white regions of the contour plots shown in Fig. 4C. In the case of the monofluoro-compound (IX), angle \( \phi_1 \) was found to range between 10° and 290°, whereas angle \( \phi_2 \) could take all possible values between 0° and 360° (Fig. 4C Left). These angles rule out a significant number of conformations. In the case of VIII, the combined coupling of three equivalent fluorine nuclei to the sugar 13C produced greater dephasing, consistent with two clusters of \( \phi_1 \) and \( \phi_2 \) values. The clusters corresponded to two groups of closely related low-energy molecular conformations (Fig. 4C Right). In both clusters, the inhibitor adopts a conformation with the sugar group extending approximately 5 Å out of the plane defined by the steroid hydroxy groups (Fig. 5A).

Models for the Docking of Cardiac Glycoside Analogues. Current understanding of the digitalis site is owed largely to phenomenological observations of mutated residues conferring ouabain resistance to HeLa cells (10). Here, direct observation of the dynamics, structure, and local environment of labeled ouabain derivatives, using a combination of solid-state NMR methods, has shed light on the dimensions of the cardiac glycosides and the surface of the \( \alpha \) subunit, and a possible docking orientation.

Fig. 5. Structural features of cardiac glycosides and the digitalis site. (A) Representatives of the two groups of closely related structures of OFDA at the digitalis site determined by 13C,19F-REDOR NMR. Carbon atoms are shown in green, oxygen atoms in red, and hydroxyl groups are represented as spheres for clarity. (B) The 10 putative transmembrane regions of the Na\(^+/K\(^{+-}\)ATPase \( \alpha \) subunit were fit to the electron density map of Ca\(^{2+}\)-ATPase (6), showing in red the mutation sites conferring ouabain resistance to HeLa cells. One possible structure of OFDA is shown alongside the protein model to illustrate the comparative dimensions of the cardiac glycosides and the surface of the \( \alpha \) subunit, and a possible docking orientation.
has been proposed, therefore, that the N-terminal H1-H2 region forms a recognition site for cardiac glycosides, whereas the C-terminal region forming the putative K+ channel (H4-H6, H8) is the inhibitory site, because K+ antagonizes ouabain binding (10, 24).

The seemingly disperse distribution of ouabain-sensitive residues form a feasible interaction site if the putative membrane helices H1-H10 are arranged according to the structure of skeletal muscle Ca2+-ATPase (Fig. 5B and ref. 6). In this arrangement, the binding site stretches across the extracellular face of the enzyme, and the helices forming the outer extremities of the site (H1 and H5) are separated by about 20 Å. This distance is approximately the same as the separation of the two ends of the ouabain derivatives in the digitals site (Fig. 5B).

The ouabain derivatives exhibit rather less inhibitory potency than the parent compound (Fig. 1A), but bind to the Na+/K+-ATPase competitively with respect to ouabain (Fig. 1B) and are thereby valid probes of the digitals site. At this stage it would be ambitious to use the information on the analogues to draw many firm conclusions about natural cardiac glycosides in the binding site, because of the lower binding affinity of the analogues and their unique chemical structures. Some suggestion of how natural cardiac glycosides are oriented within the digitals site might be provided by the analogues, however. The high mobility of the sugar group of bound ODA (Fig. 2B) indicates that this part of the inhibitor molecule is much less restrained by interactions with the protein than is the steroid moiety. From the binding conformation of OFDA (Fig. 5A), it is anticipated that the mobility of the sugar would be highest if the orientation of the steroid ring system allows the sugar to extend away from the surface of the protein. The steroid moiety then could be restrained by hydrogen bonds between the hydroxyl groups (shown in red in Fig. 5A) and the binding surface. If, on the other hand, the steroid group were flipped over so that the sugar faces toward the protein surface, the sugar would penetrate into the transmembrane region of H1 and H2. This would allow interaction of the sugar with ouabain-sensitive transmembrane residues (Cys104, Tyr108, and Glu111), but also would confer extra stability on the sugar that is inconsistent with the observed high mobility. Hence, the 3H NMR data argues in favor of a sugar group in close association with the periphery of the protein, and against a mechanism in which protein residues secure the sugar.

The downfield shift of the rhamnoside 13C resonance peak upon binding of ODA to the digitals site (Fig. 3) gives more indication of the sugar group environment. Such a shift might be caused by the ring currents of neighboring aromatic residues and one candidate is Tyr108, because substitutions at this position are known to reduce ouabain affinity (22). Tyr108 is located in the transmembrane region H1 according to the hypothetical membrane topology of Na+/K+-ATPase (31), and to attain a close proximity to the aromatic side-group the sugar therefore must be oriented toward the membrane interior. As discussed above, however, this inhibitor orientation is not consistent with the high mobility of the sugar, and it is necessary to seek alternative candidates in the extracytoplasmic regions. These include Trp310 in the H3-H4 loop, which has been labeled with a photoactivatable ouabain derivative (8), and a number of aromatic residues in the H7-H8 loop, although the latter possibility would necessitate the sugar group facing toward the C-terminal end of the protein.

In conclusion, using a combination of solid-state NMR methods and existing site-directed mutagenesis information (10), we have gained insights into the digitals receptor site and its interaction with active ouabain analogues. This work has laid important foundations for further experiments to measure direct interactions between labeled inhibitors and the digitals site.

We thank Birthe Bjerring Jensen and Angelina Damgaard for expert technical assistance and Dr. Richard Kemp-Harper for assistance with NMR instrumentation. The Biotechnology and Biological Sciences Research Council is acknowledged for a Senior Research Fellowship (to T.A.K.) and for equipment awards (with Higher Education Funding Council for England) under the Joint Research Equipment Initiative. This work was supported by Grant 51195 M from the Human Frontier Science Program.