The Binding Site of the V-ATPase Inhibitor Apicularen Is in the Vicinity of Those for Bafilomycin and Archazolid

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Background: Apicularen is a specific V-ATPase inhibitor that binds to the VO complex of the holoenzyme.

Results: Apicularen binds at the interface of the VO subunits a and c.

Conclusion: The binding site for apicularen is in the vicinity of those for bafilomycin and archazolid.

Significance: We propose the first model of binding site arrangement for these three classes of V-ATPase inhibitors.

The investigation of V-ATPases as potential therapeutic drug targets and hence of their specific inhibitors is a promising approach in osteoporosis and cancer treatment because the occurrence of these diseases is interrelated to the function of the V-ATPase. Apicularen belongs to the novel inhibitor family of the benzolactone enamides, which are highly potent but feature the unique characteristic of not inhibiting V-ATPases from fungal sources. In this study we specify, for the first time, the binding site of apicularen within the membrane spanning VO complex. By photoaffinity labeling using derivatives of apicularen and of the plecomacrolides bafilomycin and concanamycin, each coupled to 14C-labeled 4-(3-trifluoromethyldiazirin-3-yl)benzoic acid, we verified that apicularen binds at the interface of the VO subunits a and c. The binding site is in the vicinity to those of the plecomacrolides and of the archazolids, a third family of V-ATPase inhibitors. Expression of subunit c homologues from Homo sapiens and Manduca sexta, both species sensitive to benzolactone enamides, in a Saccharomyces cerevisiae strain lacking the corresponding intrinsic gene did not transfer this sensitivity to yeast. Therefore, the binding site of benzolactone enamides cannot be formed exclusively by subunit c. Apparently, subunit a substantially contributes to the binding of the benzolactone enamides.

Vacuolar-type ATPases (V-ATPases) are found in the endomembrane system of all eukaryotic cells and in the plasma membranes of many animal cells. They energize multiple transport processes and regulate the pH in cells and organelles by coupling ATP hydrolysis to proton pumping (1, 2). Their heteromultimeric structure includes two main complexes, the catalytic V	extsubscript{i} complex with a subunit composition of A	extsubscript{3}B	extsubscript{3}CDE	extsubscript{3}FG	extsubscript{3}H and the proton translocating VO complex with the subunits a, c, and c’ (in fungi additional subunit c’), d, and e (2, 3). ATP hydrolysis takes place at the hexameric headpiece built by three A catalytic subunits and three B regulatory subunits. The emerging free energy is converted into a rotation of the central stalk subunits D, F, and d and the ring formed by the proteolipid subunit c (2). For transmembrane transport, protons enter the proteolipid ring via the cytosolic hemichannel of subunit a and bind reversibly to the conserved essential glutamate in each c subunit. After rotation of the c-ring, protons leave via the luminal hemichannel of subunit a (2, 4).

By regulating the intracellular or intracellular pH, organelar V-ATPases operate in diverse processes such as receptor-mediated endocytosis, protein processing, and degradation as well as intracellular transport. Plasma membrane V-ATPases are involved in bone resorption, extracellular acidification, or energization of secondary active transport processes (2, 5). The connection of V-ATPase function to diseases such as osteoporosis or cancer and the verification of the V-ATPase as a suitable therapeutic target certainly require a comprehensive investigation of the V-ATPase, its inhibitors, and the intermolecular interactions. In the 1980s the plecomacrolide bafilomycin was identified as the first specific V-ATPase inhibitor, exhibiting nanomolar IC	extsubscript{50} values (6). Until now, a reasonable number of other inhibitory compounds from different sources have been discovered (7). Among them are the macrolactone archazolid and the benzolactone enamide apicularen, both of which show a similar inhibitory efficacy as the popular plecomacrolides bafilomycin and concanamycin (8–10). However, the benzolactone enamides exhibit a unique feature as they do not, in contrast to the other inhibitors, affect V-ATPases from fungi, and therefore they are the first source specific V-ATPase inhibitors (7, 11). In consequence, this class of compounds may be one of the most promising groups of candidates for a therapeutic use of V-ATPase inhibitors.

Concerning the inhibitor-binding sites and the mechanisms of V-ATPase inhibition, two discrete approaches using plecomacrolides have led to first insights as follows. Labeling experiments with the purified V	extsubscript{i}V	extsubscript{O} holoenzyme of Manduca sexta and a 125I-labeled derivative of concanamycin (Fig. 1) revealed the binding of plecomacrolides to the V	extsubscript{O} subunit c (12). Simul-
taneously, mutational analysis of the V$_c$ subunit c in Neurospora crassa disclosed that certain single amino acid exchanges in the sequence of this subunit altered the affinity of bafilomycin to the V-ATPase (13). Later on, additional single amino acid exchanges in subunit c of the V-ATPases from N. crassa and Saccharomyces cerevisiae resulted in a more precise localization of the pleacomacrolide-binding site, which accordingly resides at the interface between helices 1 and 2 of one subunit c and helix 4 of an adjacent subunit c in the ring (14, 15). Interestingly, the c-ring did not appear to contain the whole pleacomacrolide-binding site because mutations in subunit a of the yeast V-ATPase also conferred resistance to bafilomycin (16). In our previous photoaffinity labeling (PAL) studies using the concanamycin derivative mentioned above, the photoactivatable cross-linking diazirinyl group was bound to the macrocyclic ring of the inhibitor that led to an exclusive label at subunit c (12). Labeling of merely subunit c was surprising regarding the length (6.4 Å) and flexibility of the attached diazirinyl. However, with respect to the mutational analysis and modeling of the binding site within the c-ring, this was a strong indication that position C9 of concanamycin may be deeply buried between two adjacent c subunits. In this study, we used derivatives of bafilomycin and concanamycin modified with the newly developed $^{14}$C-labeled 4-(3-trifluoromethyl-diazirin-3-yl)benzoic acid (17). By repositioning the diazirinyl moiety to the opposite side of the pleacomacrolide structures (Fig. 1), we anticipated labeling not only of subunit c but now also of subunit a. For the modification at C23, we did not expect strong influence on the inhibitory efficacy, as in previous studies it had already been shown that this position has only a negligible effect and that it does not seem to belong to the major pharmacophore (18–20).

The binding site of the archazolids originally had been presumed to overlap to a large extent with that of the pleacomacroldes as archazolid prevented binding of a concanamycin derivative (10). However, the binding site for archazolids is relocated to the equatorial region of the c-ring and therefore overlaps with the pleacomacrolide-binding site to a minor extent than previously thought (21). This revision has been derived from recent site-directed mutagenesis of the yeast V-ATPase subunit c and labeling of the M. sexta V-ATPase using a radioactive derivative of archazolid A as well as the fluorescent dicyclohexylcarbodiimide derivative NCD-4.

Up to now, information concerning the binding site of the benzolactone enamides is rare. For the benzolactone enamide salicylihalamide A, it has been reported that it binds to a different site than the pleacomacrolides, although it inhibits proton translocation through the V$_O$ complex (12, 22). Recent labeling experiments in the presence of apicularen revealed no interference of pleacomacrolide, archazolid, or NCD-4 binding to subunit c (10, 21). Yet it was not possible to elucidate where apicularen binds within the V$_O$ complex. The development of the $^{14}$C-labeled 4-(3-trifluoromethyl-diazirin-3-yl)benzoic acid mentioned above now provided a convenient way to prepare an apicularen derivative that irreversibly cross-links to the protein upon UV exposure and therefore could be used to identify the interacting V-ATPase subunit(s) (17).

Furthermore, we used the radioactive derivatives of apicularen, bafilomycin, and concanamycin as well as nonradioactive compounds in competition assays to gain new insights into the interaction of the inhibitors. Considering the fact that the fungal V-ATPases are insensitive to benzolactone enamides, we used yeast deletion mutants deficient in subunit Vma3, Vph1, or Stv1 for the heterologous expression of their human or insect homologues to prove whether it is possible to transfer sensitivity against apicularen to the yeast-human or yeast-insect hybrid V-ATPase. Summing up all available information, we provide a model predicting the arrangement of the binding sites for pleacomacrolides, archazolids, and benzolactone enamides within the V$_O$ complex.

**EXPERIMENTAL PROCEDURES**

**Inhibitors**—Bafilomycin A$_1$, concanamycin A, 21-deoxyconcanamycin A, apicularen A, archazolid A, and saliphenyllahalamide were isolated or prepared as published (8, 9, 18, 23). 21-0-[4-(3-Trifluoromethyl-diazirin-3-yl)benzoyl]bafilomycin A$_1$ (D-bafilomycin), 23-O-[4-(3-trifluoromethyl-diazirin-3-yl)benzoyl]-21-deoxyconcanamycin A (D-concanamycin), and their $^{14}$C-labeled derivatives were prepared according to Ref. 17. Based on previous evaluation of apicularen analogues, we favored position C3 to attach the diazirinyl moiety (7, 24). For synthesis of 3-O-[4-(3-trifluoromethyl-diazirin-3-yl)benzoyl]apicularen (D-apicularen), to a solution of apicularen A (1.5 mg, 3.4 μmol) in CH$_2$Cl$_2$ (5 ml) was added 4-(dimethylamino)-pyridine (0.9 mg, 7 μmol), 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide (1.1 mg, 7 μmol), and 4-(3-trifluoromethyl-diazirin-3-yl)benzoic acid (1.1 mg, 4.8 μmol), and the solution was stirred for 2.5 h in the dark at 25°C. Column chromatography of the mixture on silica gel (CHCl$_3$/MeOH 25:1) furnished 1.4 mg (63%) of D-apicularen (R$_f$ = 0.17, CHCl$_3$/MeOH 50:1). NMR data of D-apicularen is as follows: $^1$H NMR (600 MHz, [D$_6$]-acetone): δ = 0.99 (t, J = 7.5 Hz, 3H, 25-H$_3$), 1.46–1.56 (m, 3H, 14-H$_8$, 10-H$_{10p}$, 12-H$_{12p}$), 1.65 (dd, J = 5.3, 7.8, 13.1 Hz, 1H, 12-H$_{12a}$), 1.78 (dt, J = 10.9, 14.7 Hz, 1H, 14-H$_{14}$), 1.95 (dt, J = 4.6, 13.0 Hz, 1H, 10-H$_{10}$), 2.00–2.06 (m, 2H, 16-H$_{16}$), 2.27 (dqunt, J = 1.6, 7.5 Hz, 2H, 24-H$_{24}$), 2.60 (dd, J = 1.5, 14.6 Hz, 1H, 8-H$_{8}$), 3.44 (dd, J = 10.4, 14.6 Hz, 1H, 18-H$_{18}$), 3.83–3.89 (m, 2H, 9-H, 11-OH), 3.98 (m, 1H, 11-H), 4.23 (m, 1H, 13-H), 4.97 (dt, J = 7.4, 14.5 Hz, 1H, 17-H), 5.39 (m, 1H, 15-H), 5.69 (dd, J = 11.5 Hz, 1H, 20-H), 5.79 (m, 1H, 23-H), 6.71 (dd, J = 10.4, 14.5 Hz, 1H, 18-H), 6.85 (dt, J = 1.1, 11.5 Hz, 1H, 21-H), 7.23 (m, 2H, 4-H, 6-H), 7.44 (dd, J = 7.7, 8.2 Hz, 1H, 5-H), 7.50 (d, J = 8.5 Hz, 2H, Ar-H), 7.51 (m, 1H, 22-H), 8.23 (d, J = 8.7 Hz, 2 H, Ar-H), 8.92 (d, J = 10.4 Hz, 1 H, NH).$^{13}$C NMR (150.8 MHz, [D$_6$]-acetone) is as follows: δ = 14.3 (C-25), 21.0 (C-24), 29.9 (CN$_2$, hidden), 35.8 (C-16), 38.5 (C-14), 39.0 (C-10), 39.8 (C-8), 40.1 (C-12), 64.7 (C-11), 67.5 (C-13), 73.7 (C-9), 74.6 (C-15), 89.1 (C-275 Hz, C-F), 125.4 (C-22), 126.4 (C-18), 127.8 (C-Ar), 129.0 (C-4), 130.5 (C-5), 130.6 (C-7), 131.4 (C-Ar), 131.7 (C-2) 134.7 (C-Ar), 135.2 (C-25), 164.1 (C-1’), 167.9 (C-1).

Using [1-$^{14}$C]-4-(3-trifluoromethyl-diazirin-3-yl)benzoic acid, the $^{14}$C-labeled apicularen derivative was prepared accordingly with a specific activity of 37.9 mCi/mmol as determined with a
binding site of the V-ATPase inhibitor apicularen

liquid scintillation counter. All inhibitors were dissolved in DMSO and stored in stock solutions (10 mM) at −80 °C.

**Photoaffinity Labeling**—Thirty µg of M. sexta V1/V0 holoenzymes, 20 µg of V1, or 10 µg of V0 complex, respectively, were incubated with 52 µM 21-O-[4-(3-trifluoromethyl)diazirin-3-yl]-[14C]-benzoyl] fadilomycin A1, 14C-D-bafilomycin, 52 µM 23-O-[4-(3-trifluoromethyl)diazirin-3-yl]-[14C]-benzoyl]-21-deoxyconcanalide A1 (14C-D-concanalide), or 100 µM 3-O-[4-(3-trifluoromethyl)diazirin-3-yl]-[14C]-benzoyl]apicularen (14C-D-apicularen) for 5 min at 25 °C.

In initial experiments, the presence of ATP apparently led to a slightly better label efficacy (data not shown). This may be the consequence of an increased number of V-ATPase inhibitor complexes forced into these specific conformations in which the inhibitor is, as Bowman and co-workers suggested, bound “like a stone in the gears” (14). Therefore, 1 mM ATP in 1.5 mM MgCl2 was added to a final volume of 40 µL. For competition assays, 52 µM of the 14C-diazirinylbenzoyl-labeled inhibitors were used after a preincubation with a 10-fold excess of a non-radioactive inhibitor for 5 min at 25 °C. Cross-linking was induced by irradiating the samples for 1 min with UV light (366 nm) on ice. Subsequently, the samples were separated by SDS-PAGE (T 17%, C 0.4%) and stained with Coomassie. The gels were then dried on Whatman paper, exposed to a phosphor screen for 72 h, and analyzed with a phosphorimager (GE Healthcare).

**Minimum Energy Calculation for D-Concanalide**—For a theoretical insight from MM2-force field calculations on isolated systems in the gas phase the diazirinyl-labeled concanalides were investigated as example for nanomolar V-ATPase inhibitors. We calculated in respect to the possible positions which the flexible diazirinyl group may occupy in comparison to the go fV O complex, resulted to pH 5.5, with 100 µM 3-O-[4-(3-trifluoromethyl)diazirin-3-yl]-[14C]-benzoyl|apicularen (14C-D-apicularen) for 5 min at 25 °C.

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**Yeast Strains and Growth Conditions**—The yeast strains used in this study were BMA64-1B (MATa, ura3-52; trplΔ2, leu2-3,112; his3-11, ade2-1, can1-100) (Euroscarf, Frankfurt, Germany) and BMA64-1B/vma3 (MATa, ura3-52, trplΔ2, leu2-3,112, his3-11, ade2-1, can1-100, vma3::HIS3) (21). The complemented strain BMA64-1B/vma3+ VMA3 was obtained by transformation of the plasmid carrying the VMA3 gene pRS415/VMA3 (21) into the vma3 deletion mutant by electroporation.

The mutant strain BY4741Δvph1Δstv1 was obtained by crossing the haploid strains BY4741Δvph1 and BY4742Δstv1 (both Euroscarf, Frankfurt, Germany). Each 500 µL of the appropriate overnight cultures were mixed and pelleted and afterward incubated overnight at 30 °C on agar plates containing YPDA, pH 5.5. Small amounts of the resulting zygotes were resuspended in 100 µL of H2O, applied on a new agar plate, separated by using a micromanipulator (Singer MSM), and again incubated at 30 °C. To obtain haploid double deletion mutants, the diploid strains were sporulated on agar plates containing 2% potassium acetate for 8 days at 30 °C. Again, a small amount of spores were resuspended in 100 µL of H2O, treated with 100 µg of zymolyase (100 units/mg, MP Biomedicals) for 10 min, and applied on an agar plate (YPD, pH 5.5), and the tetrads were separated by using a micromanipulator (Singer MSM). By selection on plates containing a micromanipulator (Singer MSM). By selection on plates containing 250 µg/µL G418, those tetrads of which two of the colonies grew on media with G418 and two colonies that did not grow on G418 were used for a PCR analysis of the genotype. By using two different primer combinations, (a) TACTCAAGCCGCGCTGTAATCCAG-TTGCAGACG and ATCGCGAGCCCATTTATACC and (b) GACATAGGCCCACAGTGATG and ATCGCGAGCCCAT-TTATACC, as well as the isolated genomic DNA of the four tetrad colonies as templates, the deletions of the genes vph1 and stv1 were confirmed.

Cells were usually grown on YPD, pH 5.5, additionally containing 0.02% adenine (YPD, pH 5.5). The medium was buffered to pH 5.5 with 50 mM MES, 50 mM MOPS. For YPD, pH 7.5, the pH was adjusted using NaOH. YPD, pH 5.5, medium with 0.1 M CaCl2 was obtained by adding 10% of a sterile 1 M CaCl2 stock solution after autoclaving the medium. For plates, 2% agar was added. Selection of cells was carried out by using plates with SD medium containing 0.67% yeast nitrogen base without amino acids, 2% glucose, 1.5% agar, and 10% drop out amino acid solution (21) lacking the respective amino acid. The medium was buffered with 50 mM MES, 50 mM MOPS and adjusted to pH 5.5 using HCl.

**Plasmid Construction**—The plasmid pXJ40-KKO-16k containing the coding sequence of the gene ATP6VOC of the human V-ATPase subunit c was kindly provided by M. A. Skinner (25). The ATP6VOC gene was amplified by PCR using the plasmid pXJ40-KKO-16k as template and the forward primer TACTCAAGCGCTTATGCGTCCAGACG and the reverse primer TACTCAAGCGCTTCAAGCGTAACTCGG-TACGCGATGTTGAGAGATGAG. Notably, the second primer additionally contains the coding sequence for a single C-terminal HA tag. The PCR product was cloned into the yeast expression vector pGAD7 by using the HindIII restriction sites resulting in a plasmid called pGAD7/VATP6VOC. The gene of the M. sexta proteolipid subunit c was amplified by PCR using a Bluescript plasmid containing the accordant coding sequence (26) as template as well as the forward primer TACCTAAAGCGCTTATGCGTCCAGACG and the reverse primer TACTCAAGCGCTTATGCGTCCAGACG and the reverse primer TACTCAAGCGCTTATGCGTCCAGACG.

The PCR product was cloned into the vector pGAD7 by using the HindIII restriction sites. The resulting plasmid pGAD7/Msc and the plasmid pGAD7/VATP6VOC were transformed into the vma3 deletion mutant BMA64-1B/vma3+ by electroporation, and the cells were selected on SD medium without leucine.

The construction of a plasmid carrying the coding sequence for a hybrid subunit containing the N-terminal half of yeast VPH1 and the C-terminal half of the human a4 subunit was performed by homologous recombination in the yeast wild type strain BY4741 (Euroscarf, Frankfurt, Germany). For this purpose, the C-terminal half of the human a4 subunit encoded by the amino acids 396–840 and additional homologous regions were amplified by PCR using the primers CGGTATTGCTCATTGAAACAGAGCCGCGCTGTAATCCAG-TTGCAGACG and GGTGGATTGGATTGCCAAGAGGAGATGAG. Notably, the second primer additionally contains the coding sequence for a single C-terminal HA tag. The PCR product was cloned into the vector pGAD7 by using the HindIII restriction sites. The resulting plasmid pGAD7/Msc and the plasmid pGAD7/VATP6VOC were transformed into the vma3 deletion mutant BMA64-1B/vma3+ by electroporation, and the cells were selected on SD medium without leucine.

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Karet) as template. For the plasmid pRS415/VPH1, the yeast VPH1 gene and in addition 400 bp upstream and downstream implying its native promoter and terminator was cloned into the yeast CEN vector pRS415. Afterward, the plasmid was linearized by the restriction enzyme BsgI. This open pRS415/VPH1 as well as the C-terminal half of the human a4 subunit were transformed into the yeast wild type strain BY4741. By homologous recombination, the C-terminal half of VPH1 encoded by the amino acids 411–840 was replaced by the C-terminal half of the human a4 subunit resulting in a yeast human hybrid subunit a. The plasmid was called pRS415/VPH1/a4.

Serial Drop Dilution Assay—To analyze the V-ATPase function, serial drop dilution assays were carried out. Yeast strains were grown overnight in 5 ml of YPDA, pH 5.5, and diluted to 10^5 cells/ml in distilled water. Serial 10-fold dilutions from 10^4 cells/ml to 10^0 cells/ml were prepared, and each 5 µl was dropped onto plates with different media. After 3 days of incubation at 30 °C, pictures were taken.

Other Methods—The V1VO-ATPase holoenzyme, the VO complex, and the V1 complex were isolated from M. sexta midgut as published previously (12, 27). Yeast vacuolar membranes were purified as described by Bockelmann et al. (21).

V-ATPase assays using the M. sexta V1VO holoenzyme or V1 complex or yeast vacuolar membranes were carried out as already reported (21, 27, 28). Protein content was determined using Amido Black 10B (28). SDS-PAGE, Western blotting on nitrocellulose, and immunostaining were performed as described previously (12, 29). For immunodetection, the primary antibodies against the V-ATPase subunits A (30) and e (31) from M. sexta, Vma1 (A6422, Invitrogen), and Vma6 (gift from Christian Ungermann, University of Osnabrück) from yeast, the HA tag (Y-11) (sc-805, Santa Cruz Biotechnology), and the secondary antibodies against rabbit (A3687, Sigma) and mouse (A3562, Sigma) were used.

RESULTS AND DISCUSSION

Labeling of the V-ATPase with Radioactive PAL Inhibitors—Before the diazirinyl-labeled derivatives of bafilomycin, concanamide, and apicularen (for structures see Fig. 1) were used for photoaffinity labeling, we tested in enzyme activity assays whether the modifications had an influence on V-ATPase inhibition. Compared with the natural products, the labeled derivatives exhibited reductions of their inhibitor potential by a factor of only 5–100 (Fig. 2) and therefore appeared to be highly suitable for the labeling assays.
To determine which subunits of the V-ATPase interact with the inhibitors, the V₁V₀ holoenzyme, the V₀ complex, or as a control, the V₁ complex were incubated with the ¹⁴C-diazirinyl-labeled derivatives of the inhibitors, and the cross-link reactions were induced by the exposition to UV light. The results of autoradiography after SDS-PAGE are shown in Fig. 3. Generally speaking, labeling patterns appeared to be rather similar for all three inhibitors. The comparison of the exposed (lanes with even numbers, 4–20) and unexposed samples (lanes with odd numbers, 5–21) revealed a clear labeling of the VO subunits a and c in the V₁V₀ holoenzyme as well as in the V₀ complex.

To date, subunit c was the only polypeptide of the VO complex that could be labeled successfully by inhibitors such as concanamycin, archazolid, or NCD-4 (12, 21). Yet the appearance of a label at subunit a was not unexpected as its participation in plecomacrolide binding had already been shown by site-directed mutagenesis studies in the yeast V-ATPase (16). Additionally, it was anticipated that placement of the diazirinyl in the concanamycin derivative at position C23 instead of position C9 as in the previous ¹²⁵I-concanolide used in 2002 (12, 17) would lead to a different labeling pattern. This concept is based on the assumption that the diazirinyl attached to the C9 of the inhibitor is, as already mentioned above, buried in the binding site between two adjacent c subunits, whereas the diazirinyl at position C23 probably is localized at the interface between subunits a and c. For a rough estimation of the space-compassing manner of the diazirinyl derivatives and how they label two opposing subunits, we exemplarily performed a minimum energy calculation for D-concanolide. The model provides a view on the theoretical positions and spanned distances of the diazirinyl group, which is, with respect to the pharmacophore, flexible around its single bonds of the ester moiety. The results of the calculations pointed to two preferred conformations, one with the diazirinyl moiety above and one below the macrolactone ring close to the molecule (Fig. 4) and a small distance of the label. These theoretical data support the hypothesis derived from the functional assays that labeling of both subunits a and c is conclusive even though the inhibitor is bound in one certain orientation in the binding site.

The irradiated samples of the V₀ complex exhibited a further band below subunit a, which was most probably a degradation product of subunit a and also visible in the Coomassie staining of the V₀ complex. Labeling of the V₁ subunits in the V₁V₀ holoenzyme and in the V₁ complex are considered unspecific as it occurred in both the irradiated and nonirradiated samples. In addition, we may exclude specific inhibitory interactions with the V₁ complex as it was already shown previously that the ATPase activity of the V₁ complex of M. sexta and yeast is not affected by the plecomacrolide concanamycin and that salicylhalamide, another benzolactone enamide, did not inhibit the V₁ complex of bovine clathrin-coated vesicles (22, 27, 32). In this study, we confirmed these results for the V₁ complex from M. sexta using either 30 μM of D-bafilomycin, D-concanolide, D-apicularen, or their parent compounds. None of these inhibitors had an effect, neither on the Ca²⁺-ATPase nor Mg²⁺-ATPase activity of the isolated V₁ complex (data not shown).

FIGURE 2. Inhibition of the ATPase activity of the M. sexta V₁V₀ holoenzyme by bafilomycin, concanamycin, apicularen, and their derivatives. Values represent the means ± S.D. of experiments with three different preparations. The specific enzyme activity of the controls without inhibitors was 1.6 ± 0.4 μmol-mg⁻¹-min⁻¹.

FIGURE 3. Photoaffinity labeling of M. sexta V₁V₀ holoenzyme, V₀ complex, and V₀ complex with the ¹⁴C-labeled derivatives of D-bafilomycin, D-concanolide, and D-apicularen. For the labeling assays, 30 μg of V₁V₀ holoenzyme, 20 μg of V₀ complex, or 10 μg of V₀ complex were first incubated for 5 min at 25 °C with 52 μM ¹⁴C-D-bafilomycin, 52 μM ¹⁴C-D-concanolide, or 100 μM ¹⁴C-D-apicularen, respectively. For the labeling assays, 30 μg of V₁V₀ holoenzyme, 20 μg of V₀ complex, or 10 μg of V₀ complex were first incubated for 5 min at 25 °C with 52 μM ¹⁴C-D-bafilomycin, 52 μM ¹⁴C-D-concanolide, or 100 μM ¹⁴C-D-apicularen, respectively. Before the samples were exposed to UV light (366 nm) for 1 min (+) or kept in the dark (−), 1 mM Mg-ATP was added. Afterward, the protein subunits were separated by SDS-PAGE, stained with Coomassie Blue, and exposed to a phosphor screen. Lanes 1–3, typical staining with Coomassie Blue; lanes 4–21, readout of the phosphor screen.
Additional labeling of V1 subunits may thus be interpreted as a consequence of unspecific binding of the derivatives due to the high protein and inhibitor concentrations in the assays, as they appeared in both the exposed and unexposed samples and even in the purified V1 complex. A further label close to the V1 subunits F and G was found in the V1VO holoenzyme and in the VO complex but not in the V1 complex. To check the origin of this unexpected label, pieces containing subunits F and G as well as the areas above, in between, and below these subunits were excised from a gel for further analysis. Scintillation counting revealed no radioactivity in pieces containing subunits F and G, but in contrast, radioactivity was detected at the interspace between the subunits and below subunit F (data not shown). In addition, mass spectrometric analysis (ESI-MS) verified subunits F and G, but no protein was detected in the areas above, in between, and below the subunits. Therefore, we suppose that these labels result from cross-links of the diazirinyl-derivatives with lipid or detergent molecules that remain bound specifically to the VO complex during purification. This assumption is supported by the results of von Ballmoos et al. (33) who observed cross-links between lipids and diazirinyl-labeled carbodiimide inhibitors interacting with the F1 subunit c of the ATP synthase.

FIGURE 4. Minimum energy calculation for D-concanolide. For 21-deoxyconcanolide A, the space of the connected diazirinyl moiety is placed preferably at the east hemisphere. The length of the benzoyl-diazirinyl label can be calculated to 6.4 Å and its distance from the macrolactone ring to 5.4 Å (A). Minimum energy calculations suggest that the diazirinyl group is situated preferably above and below the macrolactone ring very close to the molecule (A and B). A, benzoyl diazirinyl system is on the other side of the carbonyl group of the macrolactone ester. B, benzoyl diazirinyl system is on the same side of the carbonyl group of the macrolactone ester.

FIGURE 5. Western blot analysis of the V-ATPase with 14C-D-apicularen. For the identification of the 14C-D-apicularen-labeled band with the approximate molecular mass of subunit e 30 μg of the V1VO holoenzyme were incubated with 100 μM 14C-D-apicularen, and before the samples were exposed to UV light (366 nm) or kept in the dark, 1 mM Mg-ATP was added. Afterward, the protein subunits were separated by SDS-PAGE. Lane 1, staining of the SDS-gel with Coomassie Blue; lane 2, staining of the proteins with Ponceau S after electro-transfer onto a nitrocellulose membrane; lane 3, immunostaining with monoclonal anti-A (221-9) and anti-e (224-3) antibodies; lane 4, autoradiography of the irradiated sample; lane 5, autoradiography of the nonirradiated sample.

Treatment of the V1VO holoenzyme as well as the VO complex with 14C-D-apicularen exhibited an additional radioactively labeled band with an approximate molecular mass of 20 kDa (Fig. 3, lanes 16 and 20). This size suggests that two polypeptides of the VO complex could be labeled subunit c and/or subunit e. Although a gene encoding a putative subunit c was identified in the M. sexta genome project with a calculated molecular mass of 27 kDa, we have no indication that a product of this gene is an integral part of the purified V-ATPase derived from the plasma membrane of M. sexta. Extensive analysis of all proteins of the purified V1VO holoenzyme, the purified VO complex, and chloroform/methanol extracts of both by MALDI-MS and ESI-MS supplemented by N-terminal sequencing revealed so far no evidence for the presence of a subunit c (12, 34). Based on this state of knowledge, we so far do not see any evidence for the presence of subunit c in the plasma membrane V-ATPase of M. sexta. However, Western blot analysis of the 14C-D-apicularen-labeled VO holoenzyme using monoclonal antibodies against subunit e clearly identified this labeled band as subunit e (Fig. 5, lane 3). Being a part of the VO complex probably associated with subunit a (35), it is very likely that subunit e gets labeled due to its close proximity to subunit a, its position opposite to the c-ring, or even due to a direct contribution to the binding site. Anyhow, the

3 M. Huss and H. Wieczorek, unpublished results.
result that D-apicularen is the first inhibitor also labeling subunit e indicates again that the binding site of the benzolactone enamides is indeed different from the binding sites for the plecomacrolides and the archazolids and that the mechanism by which the benzolactone enamides inhibit V-ATPases may also be a different one. Unfortunately, nearly nothing is known about the exact localization or the function of subunit e in the V-ATPase, except that it is an integral and essential part of the VO complex (12, 31, 36, 37). Although subunit e is positioned next to subunit a in some models of the VO complex, until now there are no experimental data supporting a direct interaction between subunit e and any other subunit of the VO complex (36, 38). Therefore, it appears difficult to integrate subunit e into a concise model of the localization of the inhibitor-binding sites for which reason we will draw our model without subunit e.

Specification of the Arrangement of the Binding Sites for Bafilomycin, Concanamycin, and Apicularen by Competitive Labeling—To investigate to what extent the binding sites for apicularen, concanamycin, and bafilomycin overlap within the VO complex, we performed photoaffinity labeling assays with the 14C-diazirinyl-containing derivatives in the presence of a 10-fold excess of the natural compounds, i.e. the V1VO holoenzyme was preincubated with DMSO, bafilomycin, concanamycin, archazolid, apicularen, or saliphenylhalamide, respectively. Then samples were incubated for 5 min at 25 °C with 52 μM 14C-D-bafilomycin. Before the samples were incubated, protein subunits were separated by SDS-PAGE, stained with Coomassie Blue, and exposed to a phosphor screen. Analysis of the pixel intensity of the labeled bands with ImageQuant. Values show the means ± S.D. of two independent preparations, except for DMSO (n = 6) and apicularen (n = 5).
archazolid was suppressed in a similar manner (10, 12, 21). Evidently, the interaction with the \( V_O \) subunit a also was significantly reduced by an excess of these compounds. The same effect can be seen in the \( ^{14}\text{C}-\text{D}-\text{concanolide} \) series, where bafilomycin, concanamycin, and archazolid impeded labeling of both \( V_O \) subunits a and c (Fig. 7, \textit{lanes} 4, 6, and 8). Notably, in neither series did the benzolactone enamides apicularen and saliphenylhalamide prevent labeling by \( ^{14}\text{C}-\text{D-plecomacrolides} \) (Figs. 6, \textit{lanes} 10 and 12, and 7, \textit{lanes} 10 and 12). This result supports the previous assumption that the binding site for the benzolactone enamides is significantly different from that for the plecomacrolides (10, 12, 22).

Competition assays using \( ^{14}\text{C}-\text{D-} \text{apicularen} \) led to the unexpected result that the native apicularen itself did nearly not prevent the labeling (Fig. 8, \textit{lane} 10). A more effective reduction of labeling was achieved by preincubation with the nonradioactive D-apicularen and with saliphenylhalamide, another benzolactone enamide (Fig. 8, \textit{lanes} 12 and 14). Therefore, we suggest that the less bulky native apicularen might be too small to cover the complete binding site of \( ^{14}\text{C}-\text{D-} \text{apicularen} \). By contrast, the nonradioactive D-apicularen has the same size, and saliphenylhalamide has a comparable size (Fig. 1), and therefore both compounds are more suitable to displace \( ^{14}\text{C}-\text{D-} \text{apicularen} \). Furthermore, we may exclude binding of D-apicularen to an alternative binding site due to the sheer presence of the attached diazirinyl group, because a control compound modified with a diazirinyl was, in contrast to D-apicularen, not able to inhibit the V-ATPase (39). Surprisingly, an excess of the plecomacrolides bafilomycin or concanamycin also reduced labeling by \( ^{14}\text{C}-\text{D-} \text{apicularen} \) (Fig. 8, \textit{lanes} 4 and 6), whereas apicularen did not reduce labeling by \( ^{14}\text{C}-\text{D-} \text{bafilomycin} \) as described above. Yet, in experiments using D-apicularen and \( ^{14}\text{C}-\text{D-} \text{bafilomycin} \), a partial reduction of the radioactive label was detectable (Fig. 6, \textit{lane} 14). Based on these results, we suspect the diazirinyl group of D-apicularen to extend into the plecomacrolide-binding site leading to a displacement by the plecomacrolides or the other way around. Yet the binding site for the native apicularen is obviously different from that for the plecomacrolides (10, 12, 22).

The macrolactone archazolid did not prevent labeling of the V-ATPase by \( ^{14}\text{C}-\text{D-} \text{apicularen} \) (Fig. 8, \textit{lane} 8). This was expected from our previously published results with apicularen, NCD-4, and archazolid that had elucidated that archazolid binds to subunit c of the \( V_1V_O \) holoenzyme and has an overlapping but different binding site as bafilomycin (21). Taken together, our results now show that apicularen has a binding site within the subunit a or c of the \( V_O \) complex, which differs
Binding Site of the V-ATPase Inhibitor Apicularen

from those for archazolid and for the plecomacrolides, but a slight overlap of the binding sites for the plecomacrolides and apicularen cannot be excluded.

Exploration of the Apicularen-binding Site by Genetic Manipulation of the Yeast V-ATPase—To find out whether subunit a or c harbors the apicularen-binding site, we took advantage of the facts that bakers’ yeast can easily be genetically manipulated and furthermore that apicularen does not inhibit the fungal V-ATPases. The expression of the apicularen-sensitive homologues of either human V-ATPase subunit a or c in appropriate yeast deletion mutants should result in a hybrid V-ATPase, which is active and sensitive to apicularen. This would elegantly prove to which of these two subunits apicularen binds. The possibility to complement a yeast vma3 deletion mutant strain with c subunits of other organisms has already been reported, e.g. for the 16-kDa proteolipid from Nephrops norvegicus or Drosophila melanogaster (40, 41). In our approach, we expressed the human ATP6VOC in a yeast Δvma3 mutant strain that resulted in a partially restored vma– phenotype as this strain grew on media with an elevated CaCl2 concentration, whereas growth on media with alkaline pH failed (Fig. 9A) (42, 43). This moderately recovered V-ATPase activity was also confirmed by Western blot analysis of purified vacuolar membranes that revealed an appropriate amount of assembled V1VOC holoenzyme by immunodetection of the V1 subunit A as well as the VOC subunits d and human c (Fig. 9B). From ATPase activity assays with isolated vacuolar membranes, a specific enzyme activity of 0.064 ± 0.005 µmol·mg⁻¹·min⁻¹ was determined for the ATP6VOC-expressing strain, which is about 16% of the wild type activity and, according to Leng et al. (44), nearly enough activity to exhibit wild type growth. Even though this hybrid V-ATPase containing ATP6VOC was highly sensitive to bafilomycin A1, it still was not sensitive to apicularen (Fig. 10). In parallel, we also expressed the proteolipid subunit c of M. sexta in a yeast Δvma3 mutant strain. The analysis of the growth of the resulting strain on media with an elevated CaCl2 concentration or an alkaline pH as well as the Western blot analysis of the V-ATPase assembly in isolated vacuolar membranes and ATPase activity assays revealed the same results (data not shown) as for the expression of the human ATP6VOC. Similarly, this hybrid V-ATPase expressing the M. sexta Vc subunit c was not sensitive to apicularen. Therefore, subunit c of the Vc complex obviously does not harbor the binding site of apicularen.

The next step was to express the human isoform a4 of the V-ATPase in a Δvph1Δstv1 mutant strain to check whether this hybrid V-ATPase is sensitive to apicularen, and therefore subunit a may host its binding site. Unfortunately, this approach did not lead to an active V-ATPase (data not shown). Furthermore, the expression of a yeast-human hybrid subunit c containing the N-terminal half of yeast Vph1, which is important for targeting and regulation of the V-ATPase (45), connected with the C-terminal half of the human a4 subunit did not lead to the restoration of V-ATPase activity (data not shown). Nevertheless, we can conclude that subunit c does not exclusively form the binding site for apicularen and that subunit a contributes substantially to the binding of apicularen.

FIGURE 10. Inhibition of the V-ATPase activity of vacuolar membranes from the wild type strain BMA64-1B (WT) and the ATP6VOC-expressing strain (hybrid) by bafilomycin A1 and apicularen A. Values represent the means of two independent preparations ± S.D. The specific enzyme activity of the controls was 0.064 ± 0.005 µmol·mg⁻¹·min⁻¹ (ATP6VOC-expressing strain) and 0.41 ± 0.1 µmol·mg⁻¹·min⁻¹ (wild type strain BMA64-1B).

FIGURE 11. Model of the inhibitor-binding site arrangement in the membrane. Left, side view; right, top view from cytosol. For simplicity, the N-terminal part of subunit a is drawn as a single sphere, and only two c subunits of the c-ring are shown. The four transmembrane helices of the c subunits are numbered (1–4). The suggested locations of the inhibitor-binding sites of apicularen, bafilomycin, and archazolid are colored in red, blue, and yellow, respectively. D, the position of the diazirinyl group attached to apicularen. E, the position of the essential glutamate in helix 4.
Modeling of the Binding Site for Apicularen in the Vicinity of Those for Plecomacrolides and Archazolid—Taken together, the information from our results presented here and from previously published data on the binding sites for archazolid and for bafilomycin can be integrated in a concise model arranging the binding sites for three different classes of V-ATPase inhibitors within the \( V_{0} \) complex (Fig. 11). Based on mutagenesis studies of the c subunit performed in \( N. \) crassa and \( S. \) cerevisiae, the binding site for the plecomacrolides is localized between two adjacent c subunits in the cytosolic half of the membrane bilayer (15, 21), indicated in blue in our model. The archazolid-binding site, displayed in yellow, has been shown to reside in the equatorial region of one single subunit c covering the essential glutamate (21). Now we suggest that the binding site for apicularen, indicated in red, is located at the interface of subunits a and c, above the equatorial region and in the cytosolic half of the membrane bilayer. Our assumption is based on the following results: (i) apicularen labels both subunits a and c (ii) it does not interfere with archazolid binding, and (iii) it does not interfere with bafilomycin binding but may interact in the vicinity of bafilomycin, because D-apicularen, with its attached diazirinyl group, obviously interferes with the binding of the competing inhibitor (Fig. 6, lane 14). As already mentioned above, we decide to not include subunit e into our model as there is no reliable information on its localization available.

To gain further insights into the binding site of apicularen, it would be illuminative to construct a benzolactone enamide-sensitive subunit a in yeast by exchanging amino acids using the sequences of the human and insect subunits a as a guideline. This effort should be accompanied by elucidating the peptides cross-linked to D-apicularen by mass spectrometric analysis. In this context, modified derivatives with the diazirinyl group placed at another position or one of the improved diazirinylbenzoic acids with perfluorobutyl and perfluoroocetyl chains (F-PAL) (39) could also be applied.

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