Biochemical and Structural Insights into Carbonic Anhydrase XII/Fab6A10 Complex

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Abstract

6A10 is a CA XII inhibitory monoclonal antibody, which was demonstrated to reduce the growth of cancer cells in vitro and in a xenograft model of lung cancer. It was also shown to enhance chemosensitivity of multiresistant cancer cell lines and to significantly reduce the number of lung metastases in combination with doxorubicin in mice carrying human triple-negative breast cancer xenografts. Starting from these data, we report here on the development of the 6A10 antigen-binding fragment (Fab), termed Fab6A10, and its functional, biochemical, and structural characterization. In vitro binding and inhibition assays demonstrated that Fab6A10 selectively binds and inhibits CA XII, whereas immunohistochemistry experiments highlighted its capability to stain malignant glioma cells in contrast to the surrounding brain tissue. Finally, the crystallographic structure of CA XII/Fab6A10 complex provided insights into the inhibition mechanism of Fab6A10, showing that upon binding, it obstructs the substrate access to the enzyme active site and interacts with CA XII His64 freezing it in its out conformation. Altogether, these data indicate Fab6A10 as a new promising therapeutic tool against cancer.

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Introduction

Human carbonic anhydrases (hCAs) represent a family of enzymes that catalyze the reversible hydration of CO₂ to HCO₃⁻ and protons [1,2]. There are 12 catalytically active isoforms known in humans (CAs I–IV, VA and VB, VI, VII, IX, XII–XIV) that differ with respect to tissue and organ distribution, as well as catalytic efficiency and cellular localization. Despite these differences, hCAs share a high level of three-dimensional similarity, a zinc ion in the active site, and the same two-step catalytic mechanism described by Eqs. (1) and (2) [1]:

\[
\begin{align*}
\text{E}^{2+} + \text{OH}^- + \text{CO}_2 & \rightleftharpoons \text{E}^{2+} + \text{HCO}_3^- + \text{H}_2\text{O} \\
\text{E}^{2+} + \text{H}_2\text{O} & \rightleftharpoons \text{E}^{2+} + \text{OH}^- + \text{BH}^+ 
\end{align*}
\] (1)

The first step of this mechanism involves the nucleophilic attack of a Zn²⁺-bound hydroxide on a CO₂ molecule bound in a hydrophobic pocket within...
the active site, with consequent formation of \( \text{HCO}_3^- \), which is subsequently displaced by a water molecule with the generation of the catalytically inactive form of the enzyme \( \text{EZn}^{2+}-\text{H}_2\text{O} \) (Eq. (1)). The second step, which is rate-limiting, regenerates the \( \text{Zn}^{2+} \)-bound hydroxide through a proton transfer reaction from the \( \text{Zn}^{2+} \)-bound water molecule to the bulk solvent (Eq. (2); B indicates an exogenous proton acceptor from solvent) [1,3]. In most human isoforms, His64 residue, positioned in the middle of the active site cavity, assists this step by acting as a proton shuttle [4–6]. Indeed, His64 is generally observed in two different conformations, termed in and out [7–13]. In the in conformation, this histidine is orientated toward the interior of the active site to accept a proton from a network of water molecules that connects His64 to the zinc-bound water molecule, whereas in the out conformation it is orientated toward the external of the active site to deliver the proton to the bulk solvent (Fig. S1) [7–9]. Several studies suggested that the His64 conformational mobility is an essential requirement for proton transfer [3].

Catalyzing a simple yet fundamental reaction, human CAs are involved in many physiological processes; consequently, their abnormal expression levels and/or activities have been associated to different human diseases, including glaucoma, epileptic seizures, altitude illnesses, obesity, pain, and cancer [2]. For these reasons, CAs represent an interesting therapeutic target for the development of inhibitors or activators with biomedical applications [1]. Among the 12 catalytically active human isoforms, the two membrane-associated isoforms, hCA IX and hCA XII, have been shown to be overexpressed in many tumors and associated with cancer progression and metastases [14,15]. However, whereas hCA IX has been extensively characterized for drug design studies [16,17] which culminated in the progress of the sulfonamide inhibitor SLC-0111 and the monoclonal antibody RENCAREX® to clinical studies [18–23], hCA XII has been marginally investigated so far.

hCA XII is a transmembrane protein consisting of an extracellular N-terminal CA domain, a transmembrane (TM) region, and a small intracytoplasmic (IC) tail containing potential phosphorylation sites [24]. Structural studies revealed for the catalytic domain a dimeric quaternary structure and a typical \( \alpha \)-CA fold, characterized by a central twisted \( \beta \)-sheet fold by helical connections and additional \( \beta \)-strands [25]. The active site is located in a large, conical cavity, which spans from the protein surface to the center of the molecule and contains on its bottom the catalytic zinc ion coordinated by three conserved histidine residues [25].

Recently, a monoclonal antibody (mAb) towards CA XII, termed as 6A10, has been developed which is able to efficiently inhibit CA XII enzymatic activity in vitro [26] and in intact cells [27]. Notably, 6A10 reduces the growth of cancer cells in vitro and in a xenograft model of lung carcinoma, with a postulated mode of action directly dependent from inhibition of CA XII catalytic activity [27]. 6A10 was also shown to interfere with P-glycoprotein (P-GP) activity in chemoresistant cancer cell lines, resulting in enhanced chemosensitivity [28], and to significantly reduce the number of lung metastases in doxorubicin-treated mice, carrying human triple-negative breast cancer xenografts [26]. These data demonstrated that inhibition of CA XII by 6A10 could be effectively used to reduce chemoresistance of cancer cells and to interfere with the metastatic process in a clinical setting [28].

Due to the promising features of 6A10, a recombinant antigen-binding fragment (Fab) of this antibody, hereafter termed Fab6A10, has been developed. Indeed, the usage of antibody fragments has several advantages compared with full IgG monoclonal antibodies including a lower immunogenicity [29,30] and a faster penetration of tissues [31]. Here, we report an extensive biochemical and structural characterization of Fab6A10. Binding and inhibition assays demonstrated that Fab6A10 is highly specific toward CA XII, inhibiting its catalytic activity as the parental mAb 6A10. In addition, the crystallographic structure of the CA XII/Fab6A10 complex has been solved providing molecular insights into the inhibition mechanism of this Fab. Our data clearly indicate Fab6A10 as a new promising tool for the adjuvant treatment of different types of CA XII-positive cancer.

**Results**

**Fab6A10 production, affinity, and specific binding to hCA XII**

The variable immunoglobulin sequences of 6A10 were obtained by rapid amplification of cDNA ends (RACE)-PCR and fused to the constant Fab part of a hIgG1 molecule by gene synthesis. Fab6A10 was produced in CHO cells stably transfected with expression plasmids encoding the heavy and light Ig gene fragments and subsequently purified from the supernatant by affinity chromatography.

The capability of Fab6A10 to bind CA XII-positive human A549 lung cancer cells was investigated by flow cytometry incubating cells with serial dilutions of Fab6A10 (Fig. 1A and B). A concentration-dependent increase of the fluorescent signal was observed, and a \( K_d \) value of 3.4 nM was assessed (Fig. 1B and Table S1).

The specificity of Fab6A10 was assessed by comparing its binding to CA XII-positive parental ASPC1 pancreatic cancer cells with a CA XII-negative knockout subclone. Although binding of Fab6A10 to parental ASPC1 cells was clearly detectable (Fig. 1C), the CA XII-negative subclone was completely negative (Fig. 1D).
To provide further evidence of the specific binding between Fab6A10 and CA XII, surface plasmon resonance (SPR) experiments were carried out using the two purified recombinant proteins, and results were compared with those obtained using mAb 6A10. In details, either Fab6A10 or mAb 6A10 were amine-coupled to the sensor chip and subsequently incubated with serial dilutions of recombinant human CA XII. A $K_D$ of 12.8 nM was measured for the Fab compared with a $K_D$ of 5.8 nM for the entire antibody (Fig. S2 and Table S1). This difference is likely due to the monovalent character of the Fab compared with the bivalent nature of full-length 6A10 [32].

Finally, the binding capability of Fab6A10 to primary tumor tissues was evaluated by immunohistochemistry. Here again, experiments were carried out with mAb 6A10 for comparison. In detail, human glioblastoma and astrocytoma, both reported to express high levels of CA XII [33], stained positive with 6A10 (Fig. 2A and B). Similar results were obtained with Fab6A10 (Fig. 2C). In the infiltration zone, only the tumor cells show a distinct reaction in contrast to the negative infiltrated tissue (Fig. 2D). Taken together, these data demonstrate the capacity of Fab6A10 to bind CA XII in vitro.

Inhibition properties of Fab6A10 on hCA XII catalytic activity

Because mAb 6A10 was reported to inhibit in vitro CA XII enzymatic activity [26], the inhibitory effect of Fab6A10 on both the esterase and the CO$_2$ hydration activity of CA XII was evaluated here. Esterase inhibition assays (Fig. S3) showed that Fab6A10 likewise its parental antibody is an efficient inhibitor of this activity.

CO$_2$ hydration inhibition assays (Table 1) showed that Fab6A10 blocks also this activity very efficiently with a $K_I$ of 6.6 nM, comparable with that of the original, full-length antibody ($K_I$ of 6A10 = 3.1 nM) [26]. Interestingly, Fab6A10 does not inhibit the off-target isoforms CA I, CA II, and CA IX.
Biochemical and structural characterization of CA XII/Fab6A10 complex

A detailed biochemical and structural characterization of the complex formed by Fab6A10 and the CA XII catalytic domain was undertaken to elucidate the molecular basis of CA XII binding and inhibition by Fab6A10.

Before complex preparation, the oligomerization state of the recombinant CA XII was assessed by light scattering analysis showing that in our experimental conditions, differently from what reported by Whittington and coworkers [25], the protein exists as a monomer of 31 KDa (Fig. S4, red line). Concurrently, also Fab6A10 alone was investigated by light scattering, eluting as a monomer of 48 KDa (Fig. S4, green line). Complex formation was carried out by incubating CA XII and Fab6A10 in a 1:1 M ratio at 20 °C for 18 h and further purifying it by size exclusion chromatography in order to remove any excess of uncomplexed proteins. As attended, the complex eluted as a unique peak with a molar mass of 79 KDa as shown by SEC-MALS-QELS (Fig. S4, blue line).

Crystals of the CA XII/Fab6A10 purified complex were obtained by the hanging drop method using ammonium sulfate as precipitant. They belonged to the I222 space group and diffracted to 2.8 Å resolution (Table 2). The structure was solved by molecular replacement using the crystallographic coordinates of CA XII in complex with a benzene-sulfonamide inhibitor (PDB code 4WW8) [34] and the Fab of the antibody 13G5 (PDB code 3FO0) [35] as starting models, and refined to Rwork and Rfree values of 20.3% and 23.0%, respectively (see Table 2). The asymmetric unit contained two copies of the complex, which were refined with NCS restraints. The final model included for each complex residues from 3 to 262 of CA XII, residues 1-127 and 134-215 of the Fab heavy chain, and residues from 1 to 213 of the Fab light chain.

### Table 1. Inhibition data of hCA I, II, IX, and XII with Fab6A10. Inhibition data of the standard inhibitor acetazolamide (AAZ) are also reported for comparison.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Kᵢᵦ (nM)</th>
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<tr>
<td></td>
<td>hCA I</td>
</tr>
<tr>
<td>Fab6A10</td>
<td>&gt;1000</td>
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<tr>
<td>AAZb</td>
<td>250</td>
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* a Mean from three different assays, by a stopped flow technique.
  b Data are taken from Battke et al.[26].

### Biochemical and structural characterization of CA XII/Fab6A10 complex

Fig. 2. Frozen sections of human tissues immunostained with either mAb 6A10 (A, B) or Fab6A10 (C, D). Brown staining shows positivity of cells; negative cells are blue through counterstaining with hematoxylin. (A) Glioblastoma (highly malignant brain tumor). Strong staining of the tumor cells (scale bar: 20 μm). (B) Infiltration zone of a diffuse astrocytoma (low-grade brain tumor). Positive staining of infiltrating tumor cells (red arrows), surrounded by CA XII negative brain tissue (scale bar: 20 μm). (C) Glioblastoma. Strong staining of the tumor cells (scale bar: 10 μm). (D) Infiltration zone of a glioblastoma. Positive staining of infiltrating tumor cells (red arrows), surrounded by CA XII negative infiltrated brain tissue (scale bar: 10 μm).

Table 1. Inhibition data of hCA I, II, IX, and XII with Fab6A10. Inhibition data of the standard inhibitor acetazolamide (AAZ) are also reported for comparison.
The two complexes in the asymmetric unit revealed only minor differences, with an r.m.s.d. for the superposition of the corresponding Cx of only 0.3 Å. For this reason, only one complex was arbitrarily chosen for the following discussion.

Analysis of the complex structure (Fig. 3) revealed that both the heavy and light chains of the Fab are involved in the binding to CA XII, contributing to 77% and 23%, respectively, of the buried surface area of the complex which is of about 1100 Å² on each side of interaction. A total of 11 potential hydrogen bonds (Table S2), four salt bridges (Table S3), and a large number of van der Waals contacts stabilize the binding.

Fab6A10 binds CA XII utilizing all complementarity-determining regions (CDRs) (H1, H2, H3, L1, L2, and L3) (Fig. S5). However, whereas L1, L2, and L3 interact with a protein region adjacent to the active site cleft, H1, H2, and H3 strongly interact with the rim of the catalytic cavity, acting as a plug on it (Fig. 3B and C). H3 is also able to penetrate in the upper side of the cavity, with residue Tyr98 at only 6.5 Å from the catalytic zinc ion (Fig. 3D). Interestingly, Fab6A10 does not interact directly with the three histidines coordinating the catalytic zinc ion (His94, His96, and His119), the gatekeeper residues Thr199 and Glu106, and the catalytic zinc ion, but a salt bridge interaction is observed between Asp54 of the Fab H2 loop and CA XII His64 (Fig. 3A). This interaction freezes His64 in its out conformation, thus not allowing conformational flexibility necessary for its action as proton shuttle (see Introduction) [3].

The structural superposition of the unbound CA XII with the same enzyme in complex with Fab reveals that the binding of Fab6A10 to the enzyme does not cause any rearrangement either in the backbone or in the conformation of the residues within the active site cavity (Fig. S6). Interestingly, also in the unbound hCA XII, His64 is in out conformation, but it is not forced in this position by any strong interaction as it happens for CA XII His64 in complex with Fab.

Finally, because binding and inhibition experiments demonstrated that FabA610 is very specific for CA XII, a comparison between CA XII and the other isoforms here evaluated (hCA I, hCA II, and hCA IX) was carried out to clarify the molecular basis of this high specificity. In particular, a structure-based sequence alignment was performed showing that the majority of CA XII residues involved in Fab binding are not conserved among the four isozymes (Fig. S7). Indeed, as described earlier, Fab6A10 binds the whole rim of the active site cavity, which is the most variable active site region among all hCAs as reported in literature [1,18,36–38].

### Discussion

Differently to CA IX, whose expression mainly occurs in hypoxic tumors [14], CA XII beyond being overexpressed in various human tumors such as RCC [24], breast [39,40], colorectal [41,42], gastrointestinal [41], ovarian [43], and pancreatic [44] carcinoma and brain [33,45], is also present in several normal tissues [46]. However, the participation of CA XII into the extracellular acidification and maintenance of a more alkaline intracellular pH in hypoxic tumor cells [47] make this enzyme relevant to tumor research. In particular, the evidence that CA IX and CA XII knockout gives a remarkable reduction of xenograft tumor volume compared with knockout of CA IX alone [47], and that the simultaneous silencing of ca9 and ca12 combined with radiotherapy strongly decreased LS174Tr tumor progression in vivo [48], highlights how inhibition of both the proteins in tumors can have
Fig. 3. Fab6A10/CA XII complex structure. (A) Ribbon representation. Fab6A10 is reported in blue (heavy chain) and cyan (light chain), whereas CA XII is reported in yellow. The CA XII zinc ion and its coordinating histidines are shown to indicate the position of the active site. hCA XII residues buried at the interface with Fab6A10 heavy chain and light chain are reported in red and violet, respectively, whereas those at interface with both chains are in light blue. Fab6A10 residues buried at complex interface are green (heavy chain) or orange (light chain). (B, C) Fab6A10/CA XII epitope and paratope as open-book representation: (B) Epitope of Fab6A10 on CA XII. Color code is as in (A). The zinc ion at the bottom of the active site cavity is shown as a gray sphere. It is evident that the residues of CA XII at the interface with Fab6A10 heavy chain (residues in red) outline the whole rim of the active site cavity. (C) CA XII contact region on Fab6A10. Color code is...
a great antineoplastic potential [49]. Nevertheless, differently from CA IX, only few molecules specifically targeting CA XII have been developed and preclinically investigated so far [26–28,49]. Among these molecules, mAb 6A10 is one of the most promising [26–28].

Recent progress in the field of antibody engineering has allowed the development of Fabs as an interesting alternative to mAbs because they are easier to produce and purify, are less immunogenic, and show better tissue penetration and reduced toxicity [50]. Thus, based on these considerations and on the very encouraging results described for 6A10, in this paper, we reported on the development of Fab6A10 and its functional, biochemical, and structural characterization. We demonstrated the capability of Fab6A10 to bind selectively CA XII in vitro, to stain immunohistochemically malignant glioma cells in contrast to the surrounding infiltrated brain tissue, and to inhibit human CA XII. The obtained structural data are in agreement with these findings. Indeed, the crystallographic structure of the Fab6A10/CA XII complex showed that Fab6A10 strongly binds CA XII by interacting with the rim of the active site cavity, which is the least conserved active site region among all the human CA isoforms, thus explaining its strong specificity towards CA XII. Moreover, both the occlusion of the active site entrance and the limited conformational flexibility of His64 observed in the complex structure concur to the high inhibitory properties of Fab6A10. Interestingly, both mechanisms of inhibition have been described for small molecule CA inhibitors [38]. In particular, the occlusion of the enzyme active site is the mechanism utilized by coumarins, a class of small molecules acting as isoform-selective CA inhibitors [51,52], whereas some carboxylic acid derivatives have been reported to inhibit CAs by freezing His64 in its out conformation [53]. It is worth noting that to date only few other antibodies against CA XII and CA IX have been shown to inhibit enzyme catalytic activity [54–57], and that the inhibition mechanism has not been clarified in any of these.

Altogether, these data highlight that Fab6A10 retains all the features of the parental 6A10 antibody, turning it into an attractive candidate for various clinical applications. It is also important to highlight that the knowledge of the CA XII/Fab6A10 structure allows for the design of small molecules, which could act as CA XII selective inhibitors. Further studies are currently underway in our lab to investigate this topic.

Material and Methods

Preparation of the Fab6A10

Variable sequences of heavy (H) and light (L) chain of the 6A10 antibody were obtained by rapid amplification of cDNA ends (RACE; Kit by Invitrogen). For this, original rat-hybridoma cell lines were harvested (10^7) and the total RNA was isolated by phenol/chloroform and isopropanol precipitation. Subsequently, the RACE protocol was processed, beginning with cDNA synthesis, using isotype-specific primers (rat IgG2: ACAAG-GATTGCAATCCCTTGCG, Rat kappa: CTCATTCCCTGTTGAGCTCCTTGACGAC, as LC reverse primer, respectively). The products were sequenced to elucidate the sequence of the variable parts of the 6A10 antibody. Then, a fused construct of the variable sequences, a secretion signal (IL-2), and the hlgG1-kappa constant part from the Fab site was synthesized (GenScript).

The sequences were subcloned into a pHIT basis vector (established by Fraunhofer Institute for Toxicology and Experimental Medicine) and transfected into CHO Hit cells (established by Fraunhofer Institute for Toxicology and Experimental Medicine). The Fab6A10 was purified on a CaptureSelect kappaXL column (GE Life Science). Elution of bound Fab was performed at pH = 5.0 as described in the manufacturer’s instructions. Fab was eventually dia lyzed against PBS and concentrated by Amicon-Ultra centrifugation filter units (Millipore).

Flow cytometry

Cells were harvested and incubated with Fab6A10 followed by incubation with a donkey anti-human IgG secondary antibody labeled with Alexa467. Cells were analyzed on a BD FACS Canto™ cytometer and analyzed with FlowJo 9 (FlowJo, LLC). Further data analysis and Kd determination was performed by GraphPad Prism 7.

Production of recombinant CA XII catalytic domain

DNA-sequence of CA XII catalytic domain was synthesized by GenScript including a C-terminal His-tag and delivered in a pUC57 vector plasmid. The sequence was recloned into a pETM13 vector by using the restriction sites as in (A), (D) Detail of the CA XII/Fab6A10 complex. Fab6A10 is reported in blue (heavy chain) and cyan (light chain), whereas CA XII is reported in yellow. Fab6A10 residues buried at complex interface and CA XII zinc ion with the three coordinating histidines are shown in stick representation. H3 is colored in red. The close position of Tyr98 to the Zn^{2+} ion is highlighted.
XbaI (5’) and HindIII (3’). The plasmid was transformed into *Escherichia coli* Rosetta (DE3). Bacteria were grown in an autoinduction medium [58] to an OD600 of 1.0 as a preculture, and subsequently diluted to an OD600 of 0.1 for final culture. Bacteria were grown to an OD600 of 2–5, harvested by centrifugation, resuspended in lysis buffer (50 mM sodium phosphate, 300 mM sodium chloride, 20 mM imidazole, 1 mg/mL lysozyme, pH 8.0) and incubated on Agarose-NTA-beads overnight. The beads were washed three times with wash buffers containing increasing imidazole content (50 mM sodium phosphate pH 8.0, 300 mM sodium chloride, 20/50/100 mM imidazole). Finally, CA XII was eluted with buffer containing 500 mM imidazole and dialyzed against 1× PBS.

### Surface plasmon resonance

The ligand, here represented by Fab6A10 or mAb 6A10, was immobilized on a CM5 sensor chip (GE LifeSciences) by amine-coupling using a Biacore 3000™ according to the manufacturer’s instructions. Conditions: 10 µg/mL ligand, solved in 10 mM sodium acetate, pH 4.5; 0.05 M N-hydroxysuccinimide (NHS), 0.02 M 1-ethyl-3-(3-dimethyl-aminopropyl) carbodiimide (EDC); washing solution 1 M ethanolamine. Subsequently, the analyte, represented by recombinant CA XII, was applied to the chip using different concentrations such as 5, 10, 50, 100 nM (each in duplicates) to perform a kinetic analysis. 4 M MgCl₂ served as an indicator, working at an absorbance maximum of 557 nm, with 10 mM Hepes (pH 7.4) as a buffer, 0.1 M Na₂SO₄ or NaClO₄ (for maintaining constant the ionic strength; these anions are not inhibitory in the used concentration), following the CA-catalyzed CO₂ hydration reaction for a period of 5–10 s. Saturated CO₂ solutions in water at 25 °C were used as substrate. Stock solution of Fab was diluted up to 0.01 nM with the assay buffer. At least seven different inhibitor concentrations were used for measuring the inhibition constant. Inhibitor (I) and enzyme (E) solutions were preincubated together for 1 h at 4 °C before assay, to allow for the formation of the E-I complex. Triplicate experiments were done for each inhibitor concentration, and the values reported in this paper are the mean of such results. The inhibition constants were obtained by nonlinear least squares methods using the Cheng-Prusoff equation, as reported earlier [60] and represent the mean from at least three different determinations. All CA isozymes used here were recombinant proteins obtained as reported earlier by our groups [60–63].

### Immunohistochemistry

For CA XII immunohistochemistry, 8-µm-thick frozen sections were cut and fixed in Delaunay for 1 min. After that, pretreatment with H₂O₂ and blocking with normal horse serum at room temperature for 30 min followed. mAb 6A10 was incubated overnight at 4 °C in a dilution of 1:100, followed by incubation of biotinylated secondary anti-rabbit IgG antibody (Vector Laboratories, USA) in a dilution of 1:400 for 30 min. ABC-reagent (Vector Laboratories, USA) was applied for 30 min, followed by diaminobenzidine peroxidase (a kind gift of R. Feederle, Munich), which is a 6A10-specific antibody raised in mice, was used as a secondary antibody to specifically detect the Fab fragment. 1:1 M ratio for 18 h at 20 °C. The complex was finally purified by Superdex 200 in 30 mM Tris, 150 mM NaCl, pH 8.0. CA XII/Fab6A10 complex was obtained incubating CA XII/Fab6A10 overnight at 4 °C in a dilution of 1:20.31A4/ peroxidase (a kind gift of R. Feederle, Munich), which is a 6A10-specific antibody raised in mice, was used as a secondary antibody to specifically detect the Fab fragment. 1:1 M ratio for 18 h at 20 °C. The complex was finally purified by Superdex 200 in 30 mM Tris, 150 mM NaCl, pH 8.0.

Quaternary structure investigations on CA XII, Fab6A10, and CA XII/Fab6A10 complex were performed by SEC-MALS-QELS (size exclusion chromatography—multiscattering analysis—quasi elastic light scattering) as previously reported [64,65]. In particular, analyses were carried out loading 50 µL of 2.0 mg/mL Fab6A10, 50 µL of 3.9 mg/mL CA XII, and 50 µL of 2.0 mg/mL CA XII/Fab6A10 complex on a Superdex 75 or 200 column (GE Healthcare), equilibrated in 30 mM Tris, 150 mM NaCl, pH 8.0 and connected to a FPLC AKTA, coupled to a light scattering detector (mini-DAWN TROOS, Wyatt Technology) and a refractive index detector (Shodex RI-101). Data were analyzed using the program ASTRA 5.3.4.14 (Wyatt Technology Corporation).
Cocrystallization of CA XII/Fab6A10 complex and structure determination

CA XII/Fab6A10 complex was crystallized at 293 K using the hanging drop vapor diffusion technique. Drops were prepared by mixing 1 μL of complex solution (5 mg/mL in 30 mM Tris-HCl, pH 8.0, 150 mM NaCl) with 1 μL of precipitant solution (1.5 M ammonium sulfate, 0.1 M sodium acetate, pH 5.5, 0.02 M CdCl₂), and equilibrated over a well containing 1 mL of precipitant solution. Diffraction data were collected to 2.8 Å resolution, in-house at 100 K, using a Rigaku MicroMax-007 HF generator producing Cu Kα radiation and equipped with a Saturn 944 CCD detector. Before cryogenic freezing, crystals were transferred to the precipitant solution with the addition of 20% (w/v) glycerol. Data were processed using the HKL2000 software package [66]. The crystals belonged to the space group I222 with unit cell dimensions of a = 77.3 Å, b = 222.2 Å, c = 287.0 Å. The Matthews coefficient (Vm = 3.6 Å³/Da) indicated that the crystallographic asymmetric unit contained two CA XII/Fab6A10 complexes according to a solvent content of 66%. Data collection statistics are reported in Table 2.

The structure of CA XII/Fab6A10 complex was solved by molecular replacement technique using the program AMoRe [67], and the crystallographic structure of hCA XII (PDB code 4WW8) [34] and the Fab portion of the antibody 13G5 (PDB code 3FOO) [35] as model templates. The first cycles of structure refinement were carried out with CNS [68,69], using twofold NCS-restraints and an energy barrier of 300 kcal mol⁻¹ Å². Further cycles of restrained refinement were carried out using Refmac5 [70] to produce the final model with crystallographic Rwork and Rfree values (in the 49.6–2.8 Å resolution range) of 20.3% and 23.2%, respectively, the noncrystallographic symmetry (NCS) restraints were reduced to 150 kcal mol⁻¹ Å². Further cycles of restrained refinement were carried out to produce the final model with crystallographic Rwork and Rfree values (in the 49.6–2.8 Å resolution range) of 20.3% and 23.0%, respectively. Local NCS restraints, automatically generated by Refmac, were used in the refinement. Test set reflections were selected in thin shells with the program DATAMAN [71]. Model building was performed with the program O [72]. Data refinement statistics are summarized in Table 2. For the analysis of CA XII/Fab6A10 complex interactions, h-bonds, salt bridges, and buried residues have been determined by PISA [73], considering a distance between the heavy atoms (donor and acceptor) less than 3.5 Å and 4 Å for h-bonds and salt bridges, respectively.

Accession numbers

Coordinates and structure factors of the CA XII/Fab6A10 complex have been deposited in the Protein Data Bank (accession code 6RPS). Authors will release the atomic coordinates and experimental data upon article publication.

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Disclosure of interest

The authors declare that they have no conflict of interest.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.jmb.2019.10.022.

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- carbonic anhydrase XII complex;
- crystal structure;
- monoclonal antibody

These authors contributed equally to the work.

Abbreviations used:
- hCAs, human carbonic anhydrases;
- P-GP, P-glycoprotein;
- Fab, antigen-binding fragment;
- AAZ, acetazolamide;
- mAb, monoclonal antibody.

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