Conformational fingerprint of blood and tissue ACEs: Personalized approach

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Abstract

Background

The pattern of binding of monoclonal antibodies (mAbs) to 18 epitopes on human angiotensin I-converting enzyme (ACE)—“conformational fingerprint of ACE”—is a sensitive marker of subtle conformational changes of ACE due to mutations, different glycosylation in various cells, the presence of ACE inhibitors and specific effectors, etc.

Methodology/Principal findings

We described in detail the methodology of the conformational fingerprinting of human blood and tissue ACEs that allows detecting differences in surface topography of ACE from different tissues, as well detecting inter-individual differences. Besides, we compared the sensitivity of the detection of ACE inhibitors in the patient’s plasma using conformational fingerprinting of ACE (with only 2 mAbs to ACE, 1G12 and 9B9) and already accepted kinetic assay and demonstrated that the mAbs-based assay is an order of magnitude more sensitive. This approach is also very effective in detection of known (like bilirubin and lysozyme) and still unknown ACE effectors/inhibitors which nature and set could vary in different tissues or different patients.

Conclusions/Significance

Phenotyping of ACE (and conformational fingerprinting of ACE as a part of this novel approach for characterization of ACE) in individuals really became informative and clinically relevant. Appreciation (and counting on) of inter-individual differences in ACE conformation and accompanying effectors make the application of this approach for future personalized medicine with ACE inhibitors more accurate. This (or similar) methodology can be applied to any enzyme/protein for which there is a number of mAbs to its different epitopes.
Introduction

Angiotensin I-converting enzyme (ACE, CD143, EC 3.4.15.1), a zinc-metallopeptidase, is a key regulator of blood pressure participating in the development of vascular pathology and remodeling. The somatic isoform of ACE is highly expressed as a type-I transmembrane glycoprotein in endothelial, some epithelial, as well as macrophage and dendritic cells. Somatic ACE also presents as a soluble form, e.g., in plasma, cerebrospinal and seminal fluids, which lacks the transmembrane domain responsible for membrane attachment (for review see [1–2]).

Two homologous domains (N and C domains) within a single polypeptide chain comprise the majority of the structure of ACE, each containing a functional active center [3]. The three-dimensional crystal structure of ACE is still unknown. However, the model of the two-domain ACE has been recently suggested, based on the solved crystal structures of the individual C and N domains, epitope mapping of monoclonal antibodies (mAbs) to ACE, and on the electron microscopy picture of pig somatic ACE [4].

Recent ACE studies with mAbs recognizing different conformational epitopes on the surface of the catalytically active N domain (eight mAbs) and the C domain (eight mAbs) of human ACE molecule revealed that the pattern of mAb binding to ACE is potentially a very sensitive marker of the local conformation of ACE globule [5]. The changes of this pattern could be definitely attributed to the changes of the topography of the epitopes for the distinct mAbs due to denaturation of ACE globule, chemical modification, mutations, or the binding of inhibitors, as well as protein and low molecular weight (LMW) effectors [5–9]. It is noteworthy that ACE contains 17 potential N-glycosylation sites and the epitopes of all mAbs contain at least one site. As ACE glycosylation could be both cell- and tissue-specific due to different post-translational modification of ACE globule in different cells, the local topography of ACE surface produced by different cells could be also unique. We demonstrated previously that the pattern of ACE binding by a set of mAbs to 16 epitopes of human ACE—“conformational fingerprint of ACE”—is indeed cell- and tissue-specific [5, 9–10] and, moreover, confirmed that tissue-specific glycosylation of ACE is an important structural requirement for this specific “conformational fingerprint” [10–12].

Here, we described in detail a methodology of “conformational fingerprinting” of ACE using a panel of monoclonal antibodies to this enzyme. This approach allows one to detect the presence of ACE inhibitors in the patient’s blood and provides valuable information on the presence of low-molecular weight (LMW) effectors and ACE-binding proteins in the plasma [8] and this study and tissues (this study). Moreover, we demonstrated that conformational fingerprinting of ACE is a sensitive potential tool for detection of even inter-individual differences in ACE conformation.

We believe that similar approach could be applied to any enzyme/protein for which there is a number of mAbs to its different epitopes. This approach could provide unique information about an enzyme in question and its effectors, not available by any other method.

Materials and methods

ACEs from different sources

The work was carried out in accordance with The Code of Ethics of World Medical Association (Declaration of Helsinki) and was approved by the Institutional Review Boards of the Bakulev Center of Cardiovascular Surgery, Moscow University and the University of Illinois at Chicago. None of the donors were from the vulnerable population and all donors or next of kin provided written informed consent that was freely given. Human citrated plasma and...
serum, seminal fluid, lung and heart tissue homogenates were used as sources of somatic, two-domain ACE. Human plasma was used as individual plasma samples from different donors as well as a pool from 80 donors. Seminal fluid was pooled from ejaculates of more than 30 individuals. Heart and lung homogenates were prepared from tissues of individual donors as described [9] using 1:9 (weight: volume of PBS) ratio. Previously we found that ACE activity in tissue homogenates (or cell lysates) prepared with 0.25% Triton X-100 do not tolerate long storage as frozen samples (unpublished). In this study, we used freshly prepared homogenate, no proteinase or sialidase inhibitors were added.

ACEs were purified from corresponding sources using anion-exchange chromatography on DEAE-Toyopearl 650M and lisinopril affinity chromatography as in [13], then further washed and concentrated on Vivaspin filtration membranes (GE Healthcare, Sartorius Corp., Bohemia, NY) with 100 kDa limit as in [9]. Recombinant soluble ACE was obtained from culture fluid of CHO cells transfected with ACE lacking transmembrane anchor [14], kindly provided by F. Alhenc-Gelas (then INSERM Unit 352, Paris, France).

**ACE activity assay**

ACE activity in blood serum/plasma, seminal fluid or homogenates of human organs was measured using a fluorimetric assay with two ACE substrates, 2 mM Z-Phe-His-Leu (ZPHL) and 5 mM Hip-His-Leu (HHL), at pH 8.3 [15]. Briefly, 20–40 μl aliquots of samples diluted in PBS-BSA (0.1 mg/ml) were added to 200 μl of ACE substrate and incubated for an appropriate time at 37˚C. The His-Leu product was quantified fluorimetrically via complexing with o-phtaldialdehyde.

**Immunological characterization of ACE (Plate immunoprecipitation assay)**

Ninety six-well plates (high binding, Corning Inc., Corning, NY, USA) were coated with anti-ACE mAbs via goat anti-mouse IgG (Pierce, Rockford, IL, USA or IMTEK, Moscow, Russia) bridge [16] and incubated with different sources of ACE, which were equilibrated for ACE activity with ZPHL as a substrate. After washing off unbound ACE, plate-bound ACE activity was measured by adding a substrate for ACE (ZPHL) directly into the wells [16]. Sixteen mAbs to human ACE were generated in our lab [5], while mAb BB9 [17] to the N domain of ACE was kindly provided by Paul J Simmons (then Brown Foundation of Molecular Medicine, University of Texas Health Science Center, Houston, TX, USA). Additional mAb to ACE (clone 2H4 to the N domain of ACE) was generated in collaboration with Ilya N. Trakht and Gavreel F. Kalantarov (Columbia University, New York, NY, USA) (unpublished). For most experiments, we used a panel of 16 mAbs that were generated in our lab, while in some we tested also two new mAbs.

**Sequencing and genotyping**

Genomic DNA was obtained from heart tissue of patients #11 and # 27 by QIAamp DNA Mini Kit (Qiagen, Hilden, Germany), and 9 exons of ACE gene (14th–20th, 23th and 25th) were sequenced, using primers designed by Dufour et al. [18] as well from other sources (listed in Table 1). Briefly, PCR was made under the following conditions: 95˚C for 5 min and then 40 cycles including melting at 95˚C for 10 s, annealing at T_{annealing} (listed in Table 1) for 10 s and elongation at 72˚C for 20 s. PCR products were separated in 1% agarose gel), isolated by QIAquick Gel Extraction Kit (Qiagen) and sequenced using the same primers, BigDye Terminator v3.1 Cycle Sequencing Kit (Thermo Fischer Scientific, Waltham, MA, USA) and 3730xl DNA Analyzer (Applied Biosystems, Waltham, MA, USA).
Statistical analysis

All data are means ± SEM. Significance was analyzed using the Mann-Whitney test with STATISTICA 6 (StatSoft, Inc., OK).

Results and discussion

Enzyme (ACE) immune-capture assay

In order to quantify the amount of immunoreactive ACE protein and to analyze the changes in the local ACE surface conformation we used enzyme (ACE) immune-capture assay with panel of mAbs to different epitopes on the surface of N- and C domains of ACE [16, 19]. The scheme illustrating the method is presented on the Fig 1A. High-binding 96-well plate was coated with different mAbs to ACE via goat-anti-mouse IgG bridge (10 μg/ml, 50 μl in each well) in PBS. After washing of unbound mAbs the source of ACE was added (usually 2–20 mU/ml) and then, after overnight incubation and washing of unbound ACE, substrate for ACE (usually 1 mM ZPHL in 100 mM phosphate buffer, pH 8.3, containing 300 mM NaCl,) was added and precipitated ACE activity was estimated directly in the wells [16]. Generally, the linear relation of added ACE and precipitated ACE activity in wells (the more loaded ACE activity—the more precipitated ACE activity) was observed with not more than 20 mU /ml of loaded ACE activity especially when high-binding mAb, e.g., mAb 9B9, was used (Fig 1B–1D).

In order to save valuable mAbs, we used not more than 3 μg/ml of pure mAbs (or 1/20 dilution of hybridoma cell culture medium) which was found to be sufficient for coating (Fig 1C and 1D). Usually the amount of ACE immunoreactive protein was estimated using the strongest mAb to ACE, clone 9B9 [16, 20], while a pattern of ACE activity precipitation by a panel of mAbs to different epitopes on ACE—conformational fingerprint of ACE—was used for the estimation of local conformational differences in ACE surface topography due to disease [5, 7, 21] or due to tissue origin of ACE [5, 9, 10].

A great advantage of this approach is a possibility to measure ACE activity in plasma taken with EDTA or in plasma containing ACE inhibitors, because EDTA or ACE inhibitors are washed out during washing step with distilled water with Tween-20 while ACE is still bound to mAbs. We determined the number of washings necessary to eliminate strong specific ACE inhibitor, enalaprilat, from the complex with ACE. It is worth noting that, while a compound

Table 1. Primers used for sequencing.

| Exon | Sequence | Source | Tannealing |
|------|----------|--------|------------|
| 14   | GCAGAGTTTGTCTGTTTCC TGTGATGACATGCTCAGG | [18]    | 56°C       |
| 15&16 | GCCCTCAGTCCCACCTTG CTTGGGCTCAGCCACG | [18]    | 56°C       |
| 17   | TCCTGCTTCTCCATTTTC AGTACGGCAATTTTGGG | SeqWright, Inc. (Houston, TX) | 56°C |
| 18   | AAATCAGTGGCAGCTTGGGT AGATCTGCGAGTGAATGGC | In-house designed | 56°C |
| 19   | GCCCTTGGATCTGGAGGSC TCTGAATCGAGGGGCCTCC | In-house designed | 64°C |
| 20   | TGCTTTTCTCTCTCTGCGTC CACGAGTCCCAAACCTCGG | SeqWright, Inc. (Houston, TX) | 56°C |
| 23   | CACCTTGTAGGTTCTCCAAAATC AAACAAAGCCTGTTGACTCTG | In-house designed | 65°C |
| 25   | CCAGAAGTCGACGGCGATC | [18]    | 56°C       |

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can be a strong ACE inhibitor in physiological conditions, its binding constant to ACE significantly decreases in the absence of chloride [22], in this case, in distilled water. We measured ACE activity precipitated by 4 different mAbs using two substrates (ZPHL and HHL) and used the ratio of their rates of hydrolysis, ZPHL/HHL ratio, as a sensitive read-out of the presence of an inhibitor in complex with ACE [15, 23]—Fig 2. We found that the residual levels of ACE inhibitor in ACE precipitated by different mAbs are different, which is reflected in different ZPHL/HHL ratios at 1–3 washings, but five washings were enough for the complete dissociation of enalaprilat from the complex with ACE precipitated by any mAb (Fig 2).

Previously we found that the pattern of ACE activity precipitation by a set of mAbs—conformational fingerprint of ACE—could be sensitive to the presence of detergents [5]. Because we actively used this approach for the study of the fine conformation of ACE from different human tissues—tissue ACE specificity [9–10, 21]—we studied the effect of Triton X-100, used for ACE solubilization, on ACE conformational fingerprint. Fig 3 demonstrates the effect of different concentrations of Triton X-100 on ACE purified from human lung homogenate. It
appeared that the binding of 4 mAbs to purified ACE was sensitive to the presence of the detergent—two mAbs, i1A8 and 5F1, to different epitopes on the N domain and two mAbs, 1B8 and 3F11, to the epitopes on the C domain (Fig 3). Intuitively, an effect of detergent on mAbs binding to ACE should be bigger with membrane form of ACE, i.e., with the enzyme containing transmembrane anchor, than with soluble ACE. We have shown previously [24] that even purified membrane form of ACE from bovine lung carried significant amount of Triton X-100. So, we tested an effect of Triton on mAbs binding with different ACEs and found that, indeed, while the effect of Triton was clearly seen with lung ACE, in crude homogenate or purified enzyme (Fig 4A and 4B), the presence of Triton did not influence on mAbs binding to soluble ACEs, recombinant ACE (Fig 4C) or ACE from seminal fluid (Fig 4D).

Storage of human tissue homogenates can also alter ACE properties. We found that while ACE activity (with ZPHL as a substrate) in human lung homogenate was quite stable during up to 4 weeks of storage in a refrigerator, storage in a freezer at -20°C longer than one month resulted in a complete loss of ACE activity in human tissue homogenates or lysates of ACE...
expressing cells (not shown). The binding of some mAbs to ACE in human lung homogenate, however, noticeably changed already after one week of storage in a refrigerator and also changed even after short freezing (Fig 5). These data indicate on the changes of the topography of the surface of ACE protein globule during storage while active centers located deeply inside the protein globule [25] maintain their enzymatic functions. This observation is in accordance with our previous finding [26] by second derivative UV spectra of ACE solutions.

Fig 3. Effect of TritonX-100 on the mAbs binding to purified lung ACE. A-C. The effect of different concentrations of detergent Triton X-100 on the precipitation of ACE activity from ACE purified from human lung homogenate. Data are expressed as a % of control (i.e. without Triton X-100) and presented as a mean of at least 3 independent experiments. N/T–not tested mAbs. The coloring is as in the legend to Fig 2, and, in addition, values decreased more than by 50% are highlighted in deep blue.

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that γ-irradiation resulted in a decomposition of tyrosine and tryptophan residues on the surface of bovine ACE occurring even at low doses, while the enzyme retained its full enzymatic activity.

Thus, data presented in Figs 3–5 demonstrate that the fine changes in the surface topography of ACE from different sources should be measured in identical (or similar) conditions.
Fig 5. Effect of storage of lung homogenate on the recognition of ACE by mAbs. Freshly prepared human lung homogenate was prepared and then aliquoted into individual volumes for storage. Precipitation of ACE activity was compared for freshly prepared homogenate and the same homogenate but stored for different time and at different temperatures. A-C. Storage of homogenate at 4°C for different periods of time. D. Short freezing at -20°C. Data are expressed as a % of control (i.e., initial precipitation of ACE from freshly prepared lung homogenate) and presented as a mean of at least 3 independent experiments. N/T–not tested mAbs. The coloring is as in the legend to Fig 3.

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bearing in mind the presence of membrane anchor in some ACE molecules, as well as the conditions of ACE solubilization and storage.

**Detection of ACE inhibitors in patient’s blood**

Previously we demonstrated the possibility of the detection of commercial ACE inhibitors in the patient’s blood using two different approaches—calculation of the ratio of the rates of the hydrolysis of two substrates, ZPHL/HHL ratio [15], and calculation of the ratio of the binding of two mAbs, 1G12 (or 6A12) and 9B9, to ACE [6], both ratios increasing in the presence of ACE inhibitors. The detection and quantification of ACE inhibitors in the patient’s blood (i.e. anti-hypertensive adherence) definitively has clinical significance, because patients with effective suppression of ACE has better blood pressure control than those with weak response to inhibitors [27]. Bearing this in mind, we accurately compared the sensitivity of both methods (Fig 6) and demonstrated that one of common ACE inhibitors, enalaprilat, could be detected in the patient’s blood at 1 nM using ZPHL/HHL ratio (Fig 6B and 6D) and even at less than 0.1 nM (Fig 6C and 6D) using mAb-based approach. For reference, the concentration of enalaprilat in the blood at its peak (4 hours after oral administration of 10 mg of enalapril) was reported to be about 50 nM [28]. Thus, even “restricted” conformational fingerprinting of ACE with only two mAbs allows obtaining valuable information.

**Conformational fingerprint of blood ACE from different donors**

We quantified the precipitation of ACE activity from the plasma samples from different donors using wells coated (via goat-anti-mouse bridge) with 17 different mouse anti-ACE mAbs (Fig 7). Relative precipitation of ACE by strong mAbs (9B9, 2B11, 3A5, 2H9) dramatically differed compared to precipitation by weak mAbs (i1A8, 3G8, 1E10, and 3F11), likely reflecting difference in their affinity to ACE (Fig 7A). We should mention that some mAbs demonstrated anti-catalytic effect on ACE in solution at μg/ml concentrations [6, 15], whereas in the enzyme immunocapture assay (at an order lower concentrations the anti-catalytic effect did not exceed 20% (S1 Fig in [7]) in comparison with 10-fold differences in fluorescence signals demonstrated by ACE precipitated by different mAbs, weak and strong (Fig 7A). Therefore, we can consider anti-catalytic effect of some mAbs in this format as negligible.

We performed such immuno-capture assay with 8 randomly chosen plasma samples from healthy donors in two separate experiments with 5 and 3 donors. In order to see the putative inter-individual differences in the pattern of anti-ACE mAbs binding we presented the results as a ratio of ACE precipitation by any mAb (normalized for binding of mAb 9B9 to ACE corresponding to the level of the enzyme in plasma) for a certain donor to the mean value of ACE precipitation by this mAb for a number of donors (Fig 7B, 7D and 7E). The conformational fingerprint of blood ACE in general is very stable characteristics of the enzyme, as the pattern of mAbs binding to ACE from the pool from 3 plasma samples fairly well coincided with the pattern obtained for pool from 80 other plasma samples (Fig 7F). This analysis did not also reveal any peculiar ACE from blood of almost all patients, e.g., patients #5 and #L (Fig 7B and 7D), but showed that conformational fingerprint of ACE from the blood of patient #1 differed from that for other donors (Fig 7E). Moreover, if to compare conformational fingerprint in pairs with more samples, it is possible to catch some differences in ACE conformation between individuals as well. Thus, we revealed such difference for ACE from patient #5 and patient #14 (Fig 7C), likely reflecting small inter-individual differences in ACEs from the blood of different donors.

It should be emphasized, that the effect of ACE inhibitors on mAbs binding (i.e., conformational changes in ACE) was much more prominent with up to 4-fold increase of the binding of
mAbs 1G12/6A12 (Fig 6C and 6D), than inter-individual differences in ACE conformation which, in our experiments, did not exceed 30–40%. Nevertheless, the inter-individual differences observed were statistically significant (p < 0.05) and reliable. While the clinical relevance of such inter-individual differences awaits further investigation, however, it could lead to the different behavior of the enzyme in some physiological processes, e.g. those including ACE interplay with effectors [7, 8] in blood and tissues.

Conformational fingerprint of tissue ACE from different donors

Inter-individual differences in ACE conformation are more prominent for tissue ACEs. As an example, we show conformational fingerprints of heart and lung ACEs from different donors (Fig 8). Previously, we demonstrated that conformational fingerprint of ACE is very tissue-
specific [9–10, 12] and the pattern of the binding of mAbs to ACE from heart (9 samples from different donors) remarkably differed from the pattern obtained for ACE from lung samples from the same donors. However, if to compare conformational fingerprint of ACE using more samples, it is possible to find some differences in ACE surface topography between individuals, as, for example, for donor #13 (Fig 8B and 8F) and other donors. Moreover, these differences could be tissue-specific, as seen for heart and lung ACEs (Fig 8).

Conformational fingerprint of ACE from tissues of another donor, donor #27, revealed significant decrease in binding of two mAbs, 1B8 and 3F10 (Fig 9), which was observed when conformational fingerprints of ACE were compared in pairs (Fig 9A–9C), as well when conformational fingerprint of donor #27 was compared with that for mean from two other donors (Fig 9D). As the differences in 1B8/3F10 binding were found for ACEs from different tissues, heart and lung, we suspected a mutation in ACE in the overlapping region of the epitopes for...
However, when we sequenced 9 exons of the C domain which coded the overlapping region of the epitopes for mAbs 1B8 and 3F10 [29], we did not find any ACE mutation that could be responsible for this phenotype. Moreover, we did not identify any known polymorphic variants of ACE gene in patient #27, but, at the same time, we identified silent single nucleotide polymorphism rs4331 in heterozygous state (AG) in patient #11, which does not lead to amino-acid substitution but may be associated with higher activity of ACE [30].

It is worth noting that tissue- and cell-specific post-translational modifications (PTM) are common for various proteins [31–32], which can subtly or dramatically alter the protein surface topography. The most common PTM is glycosylation. The inter-individual heterogeneity of the N-glycome is already reported, at least for plasma [33–34]. Therefore, it is possible that the mutation in one of the genes (out of more than 100 genes) responsible for glycosylation of the proteins in human [35] is a reason for the increase in 1B8/3F10 to ACE from patient #27.

mAbs 1B8/3F10. However, when we sequenced 9 exons of the C domain which coded the overlapping region of the epitopes for mAbs 1B8 and 3F10 [29], we did not find any ACE mutation that could be responsible for this phenotype. Moreover, we did not identify any known polymorphic variants of ACE gene in patient #27, but, at the same time, we identified silent single nucleotide polymorphism rs4331 in heterozygous state (AG) in patient #11, which does not lead to amino-acid substitution but may be associated with higher activity of ACE [30].

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Somatic ACE is a N-type glycoprotein characterized by a considerable content of its sugar moiety, e.g., human kidney ACE was reported to have about 18% of sugars [36]. The sequence of human somatic ACE contains 17 potential sites for N-glycosylation [3], however, the information about the structure and exact positions of glycan chains in ACE from different tissues

**Fig 9.** Inter-individual differences of the conformation of ACE from heart and lung. The ratios of precipitated ACE activity from any tissue homogenate to precipitated ACE activity from another homogenate is shown for clarity. A. The ratio of precipitated ACE activity from lung homogenate of 27th donor to that of 17th donor. B. The ratio of precipitated ACE activity from heart homogenate of 27th donor to that of 17th donor. C. The ratio of precipitated ACE activity from heart homogenate of 27th donor to that of 11th donor. D. The ratio of precipitated ACE activity from heart homogenate of 27th donor to that for mean from 17th and 11th donor. Data are expressed in % and presented as a mean of at least 3 independent experiments. N/T–not tested mAbs. The coloring is as in the legend to Fig 3.

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Somatic ACE is a N-type glycoprotein characterized by a considerable content of its sugar moiety, e.g., human kidney ACE was reported to have about 18% of sugars [36]. The sequence of human somatic ACE contains 17 potential sites for N-glycosylation [3], however, the information about the structure and exact positions of glycan chains in ACE from different tissues
is very limited [10, 12, 37–38]. The main part of glycans in somatic ACE belongs to the biantennary complex type oligosaccharides as well as high-mannose and hybrid type oligosaccharides [39]. These types of oligosaccharides possess the largest structural variation due to different possible amounts of the outer chains linked to the trimannosyl core and different structures of these chains. Besides, the outer chains can bear different amounts of neuraminic acid residues on their ends.

Almost all conformational epitopes for mAbs that we generated to the catalytically active human lung ACE contain potential glycosylation sites [5]. In particular, epitopes for mAbs 1B8 and 3F10 contain glycosylation site Asn731 [29]. Therefore, we can expect that it is glycan in glycosylation site Asn731 which is different in ACE from donor #27, whereas we could not also exclude another PTM for this donor at the moment.

Further, an analysis of the pattern of a set of mAbs binding to ACE from different tissues and cells can provide some information about differences in glycosylation of ACE in these tissues [5, 9–10, 12], in particular, in sialylation of definite glycans. Fig 10A demonstrates the differences in the patterns of binding of mAbs to ACEs purified from the lung homogenate and from the blood plasma, i.e., the differences in the local surface topography of ACEs from different sources. Plasma ACE originates mainly from the lung capillaries (by 75%, according to our estimation [unpublished]) based on heterogeneous ACE expression in the capillaries from different organs [40]. It is known that sialic acid-deficient glycoproteins can be selectively removed from serum by macrophages and hepatocytes in the liver via lectins specifically recognizing penultimate galactose residues in oligosaccharide chains of a glycoprotein [41]. As an example, ACE from rabbit serum contains 3-fold higher molar ratio of sialyl- to galactosyl- residues than lung ACE [42]. Therefore, higher binding of mAbs i1A8 and 5F1 and much lower binding of mAbs 1B8 and 3F10 to lung ACE compared to plasma ACE (Fig 10A) could be attributed to the different sialylation of the oligosaccharide chains present in the epitopes of the corresponding mAbs. In particular, we can conclude that mAbs 1B8 and 3F10 to the C domain of ACE bind much better to the enzyme when glycan in the potential glycosylation site Asn731 is sialylated. The less effective binding of mAbs 1B8 and 3F10 to ACE from donor #27, therefore, indicates on less extent of sialylation (or even the absence of sialic acid residues) of this particular glycan in glycosylation site Asn731 in ACE from this donor compared to ACEs from other donors. Thus, conformational fingerprint can help to reveal some fine characteristics of the enzyme from a definite individual.

Besides, from the same data (Fig 10A) it is seen that mAb 1E10 to the C domain of ACE and mAbs 5F1 and i1A8 to the N domain bind better to the enzyme when oligosaccharide chains in the corresponding glycosylation sites, Asn666, Asn117 and Asn82, respectively, are not or, at least, less sialylated. This finding could be further extrapolated to the analysis of conformational fingerprint of ACE from other donors.

**Detection of ACE effectors in blood and tissue homogenates**

Conformational fingerprinting of ACE can also provide valuable information on putative ACE effectors and ACE-binding proteins in the blood or different tissues. The influence of these effectors on the access of epitopes on the surface of ACE for mAbs is seen from the comparison of the conformational fingerprints of ACEs purified from lung and heart homogenate and from human plasma (Fig 10). Purification of ACE from human plasma resulted in an increase of binding of four mAbs (and decrease of one mAb) the N domain and one mAb to the C domain (Fig 10A). Purification of ACE from human heart decrease binding of five mAbs to the N domain, while one mAb to the N domain and five mAbs to the C domain increased
their binding (Fig 10B). Purification of ACE from lung decreased binding of 3 mAbs to the N domain and increased binding of 5 mAbs to the C domain (Fig 10C).

From previous studies, we know that common ACE inhibitors, when bind to ACE, significantly increase the binding of two mAbs, 1G12 and 6A12, to the N-domain and decrease the
binding of two anticatalytic mAbs, 1E10 and 4E3, to the C domain of ACE [6–8, 29]. Thus, the decrease in the binding of 1G12/6A12 mAbs and the increase in the binding of 1E10/4E3 mAbs with heart and lung ACEs could be attributed, at least partly, to the removal of some endogenous ACE inhibitors, accompanying ACE in the heart and lung, as a result of ACE purification. However, the significant increase in 1B3, 1B8 and 3F10 mAbs binding cannot be explained just by the removal of ACE inhibitors but rather by the removal of some other ACE effectors/ACE binding proteins accompanying ACE in the heart and lung and shielding (masking) epitopes for mAbs 1B3, 1B8/3F10, 1E10/4E3.

The strikingly different effect of the enzyme purification on its conformational fingerprint was obtained for plasma ACE. Namely, purification of ACE from human plasma resulted in the dramatic increase in binding of 4 mAbs to the overlapping region on the N domain, 1G12/6A12/i2H5 [6], and 3G8 [20], which we should explain mainly by the dissociation of bilirubin from its complex with plasma ACE as a result of the enzyme purification, as we have shown that bilirubin dramatically decreases the binding of these very mAbs [8]. The increase of mAb 1E10 binding to the C domain of plasma ACE after purification can be explained either by dissociation of endogenous inhibitors or/and by dissociation of lysozyme which forms complexes with ACE via binding to the cleft between ACE domains and to the region of 1E10 epitope [8].

Therefore, the conformational fingerprinting of ACE in different human tissues (using a set of mAbs to different epitopes of human ACE) provides a novel structural information about characteristics of ACE in different individuals (and in different tissues), as well as its effectors, that could be clinically important.

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