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Section: Molecular biophysics

Prothrombin is a binding partner of the human Receptor of Advanced Glycation End products

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Abstract

The receptor for advanced glycation end products (RAGE) plays a key role in mammal physiology and in the etiology and progression of inflammatory and oxidative stress-based diseases. In adults, RAGE expression is normally high only in the lung where the protein concentrates in the basal membrane of alveolar Type I epithelial cells. In diseases, RAGE levels increase in the affected tissues and sustain chronic inflammation. RAGE exists as a membrane glycoprotein with an ectodomain, a transmembrane helix and a short carboxyl-terminal tail, or as a soluble ectodomain that acts as a decoy receptor (sRAGE). VC1 domain is responsible for binding to the majority of RAGE ligands including advanced glycation end products (AGEs), S100 proteins and HMGB1. To ascertain whether other ligands exist, we analyzed by MS the material pulled-down by VC1 from human plasma. Twenty out of 295 identified proteins were selected and associated to coagulation and complement processes and to extracellular matrix. Four of them contained a ycarboxyl glutamic acid (Gla)-domain, a calciumbinding module, and prothrombin (PT) was the abundant. Using MicroScale most Thermophoresis, we quantified the interaction of PT with VC1 and sRAGE in the absence or presence of calcium that acted as a competitor. PT devoid of the Gla-domain (PT des-Gla) did not bind to sRAGE, providing further evidence that the Gla-domain is critical for the interaction. Finally, the presence of VC1 delayed plasma clotting in a dose-dependent manner. We propose that RAGE is involved in modulating blood coagulation presumably in conditions of lung injury.

Introduction

RAGE is a multi-ligand receptor of the superfamily of immunoglobulins and is related to the category of cell adhesion proteins (1). RAGE was initially identified for its capability to bind advanced glycation end products (AGEs), a highly heterogeneous class of molecules that derive from non-enzymatic reaction products of sugars with proteins (Maillard reaction) (2). Beside AGEs,

RAGE binds to widely different pro-inflammatory among those members of ligands the S100/calgranulin protein family, High mobility group protein box-1 (HMGB1 or amphoterin), the lipopolysaccharide of bacterial cell walls (LPS). amyloid β -peptides, and the complement C1q and C3a proteins. Further ligands identified are extracellular matrix proteins such collagen I/IV, Mac-1 (member of the β 2-integrin family) and complex polymers such as heparan sulfates [reviewed in (3)]. In view of the large variety of ligands, RAGE was assigned to the pattern recognition receptor (PRR) family (4-7).

A wide spectrum of RAGE functions have been reported, reflecting the variety of the different ligands. RAGE is involved in the innate and adaptive immune response, in the onset and establishment of inflammation, as well as in tissue homeostasis and repair (3,8). RAGE is encoded by AGER, a single gene unique to mammals, located in proximity of the MHC locus on Chr 6 where the MHC genes and some genes of the complement system (C4-A and C2) are located [(9) and human genome at the UCSC Genome Browser (10)]. AGER is highly expressed in many tissues during embryo development, but in healthy adults its expression decreases in all tissues except in the lung where it is abundant but restricted to alveoli (11-14). Moreover, very low RAGE levels were detected in the heart, skeletal and smooth muscle, thyroid gland and adrenal cortex, in specific areas of the brain and in immune cells (dendritic, CD4+ and CD8+ T cells and monocytes) (12,15). Notably, RAGE expression increases during the inflammatory response due to a positive feed-back loop triggered by the RAGE-dependent activation of the transcription factor NF-KB and to proinflammatory stimuli (16).

The various functions of RAGE are ascribable to several intracellular pathways that are activated depending on the specific ligand, context and cell type (6). Activation of the signaling pathways occurs via the short cytoplasmic domain that is devoid of any enzymatic activity and oligomerization of the receptor facilitates the recruitment of adapter proteins, such as the cytoskeleton Diaphanous-1 protein (DIAPH1), TIRAP and MyD88, essential for signaling (3,5). Primary signaling pathways activated by RAGE are the p21ras/mitogen-activated cascades of protein kinases (MAPKs), such as ERK1/2, JNK and p38. NF-KB and JAK/STAT signaling pathways resulting in the activation of transcription of Type I/V collagen that triggers cell adhesion and cell spreading, taking also into account that collagen activates RAGE on the cell surface (6,8). In addition, the binding of RAGE to Mac-1 integrin on leukocytes triggers expression ICAM1/VCAM1 and promotes leukocyte adhesion. Interestingly, heparin was reported as a ligand that inhibits RAGE signaling (17).

Besides the full-length membrane bound receptor, a primary soluble form of RAGE is generated by alternative splicing from the *AGER_v1* mRNA, named endogenous soluble (esRAGE) (18), or by ADM10-mediated protease digestion of full-length RAGE at the cell surface (19,20). Extensive work has revealed that soluble RAGE (sRAGE) acts as a decoy for RAGE-ligands and, therefore, blocks the establishment and progression of a wide-range of pathological states in animal models such as cardiovascular diseases, diabetes, cancer and neuronal dysfunctions (21-24).

RAGE is involved in the etiology and progression of several human diseases and in ageing. In particular, RAGE has emerged as a harmful mediator of tissue damage caused by inflammation and oxidative stress, conditions that accelerate the reaction of proteins with sugars and the formation of AGEs by sustaining the activation of NF- κ B causing chronic inflammatory disorders (25).

In this work, we report the results of a study aimed to identify new RAGE interactors. RAGE consists of the extracellular IgG-like domains V, C1 and C2, a single transmembrane α -helix, and a short C-terminal cytosolic domain. V and C1 form a structural integrated domain, named VC1, which is connected to the C2 domain through a hinge region. The vast majority of RAGE ligands bind to VC1. Therefore, we exploited VC1 to pull-down new RAGE ligands from human plasma. We identified proteins associated to the blood coagulation and complement processes and to the extracellular matrix (ECM). Prothrombin (PT) was the major interactor. Moreover, we quantified a moderate-affinity binding of PT to VC1 and a higher affinity to sRAGE. These results, along with the inhibition of blood coagulation by VC1, suggest a potential new role of sRAGE in modulating blood coagulation likely in the lung where the protein is abundantly expressed.

Results

Multiple band analysis identifies new plasma interactors of VC1

In the attempt to obtain a more comprehensive picture of the network of molecules interacting with RAGE, we used a previously described in vitro pull-down assay to identify potential new binding partners present in human plasma. Briefly, the ligand-binding domain of human RAGE (VC1) was immobilized through a C-terminal Strep-tag on streptavidin (STV)-coated magnetic beads (26). The Strep-tag has high affinity for STV allowing for stringent washing and eliminating weak or unspecific binders. VC1 domain was expressed and secreted from the yeast Pichia pastoris (Fig. S2). VC1 was N-glycosylated and showed a remarkably high thermal stability and solubility, two important features to set up a robust assay (27). The use of VC1 in pull-down assays was validated by demonstrating its capability to capture in vitro modified human serum albumin (AGE-HSA) whereas lacking any interaction with unmodified HSA (26-28).

Aliquots of plasma from healthy individuals were diluted to 5 mg protein/ml and 800 µg were used in each pull-down assay. After 2 h of incubation with plasma at 4 °C, the VC1-STV-coated beads were extensively washed and the bound material was solubilized in SDS sample buffer and analyzed by SDS-PAGE. As a control, the same procedure was applied using the STV-coated beads alone (STV-beads). A major band of about 75 kDa was reproducibly detected by SDS-PAGE and Coomassie staining in the material solubilized from VC1-STV beads, but not in the material solubilized from STV-beads alone (Fig. 1A). To identify the ligands bound to VC1, we analyzed only the region of the gel above the 37 kDamarker. The abundance of STV (~14 kDa) and VC1 (32-34 kDa) proteins, solubilized as well

from the beads, did not allow the identification of ligands migrating in the range of MW lower than 37 kDa (see Experimental procedures).

Gel pieces were excised and processed for peptide LC-MS fingerprinting (*multiple band analysis*). To increase the specificity in the identification without losing sensibility, different pull-down assays were set up as follows: (*i*) 5 independent pull-down assays were performed, each with plasma from a single individual, (*ii*) a pull-down assay from a pool of three plasma samples from other individuals and (*iii*) 2 pull-down assays from the plasma of other two single subjects. The set of proteins in common to both VC1-STV and STV-beads was discarded during data elaboration. The experimental design is summarized in **Fig. S1**.

Dataset S1 lists the human plasma proteins identified by MS (for details see Experimental procedures). Twenty out of 295 identified proteins were selected by applying a cut-off of positive identifications in \geq 3 pull-down assays over a total of 8. The subset of twenty proteins, shown in **Table 1**, was analyzed by the software STRING that reveals functionally associated protein networks (29). The identified proteins converged on the complement and coagulation cascades in the KEGG classification (pathway hsa04610) with a false discovery rate of 1.05 e⁻¹⁸ (**Fig. S3**).

As indicated in Table 1, prothrombin (PT), the coagulation factors IX, the anticoagulant proteins S and Z were reproducibly detected. These proteins share the Gla-domain, a domain rich in glutamic acid residues that are modified by γ carboxylation to γ -carboxylated residues (Gla). Among the other interactors, the α -chain of the complement C4b-binding protein (C4BP) and the complement C4A are notable. C4BP, an inhibitor of C4 in the complement system, is a multimeric and abundant plasma protein (25 µg/ml), containing 7 identical α -chains and a unique β chain. Protein S (PROS1) binds C4BP through the β-chain. Approximately 60% of total PROS1 (~15 µg/ml) circulates in complex with C4BP, whereas approximately 40% (~10 µg/ml) is in free form (30). C4A occurred in all the analyzed samples (Table 1). Prior to secretion, C4A cleavage generates three chains, the largest of which is C4b (192.7 kDa), that interacts with α -chains of C4BP.

Thus, C4BP and C4A could be direct interactor or captured in complex with Protein S. Finally, C4b undergoes further processing that leads to the assembly of a complex that activates C5 and C3 convertases. Notably, C3a, a proteolytic fragment of C3 that was previously identified as a RAGE-ligand, was also identified in our study (31). In addition, α -2-macroglobulin is another interesting interactor since it is highly expressed in the lung.

Finally, another class of interactors is related to the extracellular matrix (ECM): vitronectin, thrombospondin-1, and the cartilage oligomeric matrix protein, (also known as thrombospondin-5) and support a role in cell adhesion, a process in which RAGE is known to be involved into.

Finally, to verify whether AGE modifications contributed to the binding of the interactors to VC1, we searched for the presence of AGE/ALE (advanced lipoxidation end products) modifications in the identified proteins. No AGE or ALE modifications were detected by MS using a previously described search method (26,32) (see Discussion).

Single band analysis identified the 75-kDa polypeptide as prothrombin

The major polypeptide present in pull-down assays, the 75-kDa band of **Fig. 1A**, was excised from Coomassie-stained gels and MS analyses identified this band as prothrombin (PT) (**Table 2**). Single band analysis was repeated and PT was invariably detected as the most abundant interactor of VC1.

In order to confirm that the 75-kDa band indeed represented PT, we performed Western blot analyses of the pull-down assays. A monoclonal anti-PT antibody detected clearly the 75-kDa polypeptide from VC1-STV beads which was absent in samples prepared with STV beads alone (**Fig. 1B**). Thus, PT was identified as the most prominent RAGE interaction partner from plasma. Furthermore, we performed Western blot analysis of a VC1 pull-down assay using anti-Gla mAb. As shown in **Fig. 1C**, the 75 kDa band was present, showing that PT is γ -carboxylated, a posttranslational modification that could not be detected in the conditions used in our LC-MS analysis. Accordingly, PT is the most abundant Gla-domain-containing protein captured by VC1 from plasma. This is likely related to the level of PT in the plasma that is higher than that of the other circulating Gla-domain containing proteins (see also Discussion). Accordingly, Western blot analysis failed to reveal other Gla-domaincontaining polypeptides. As expected, no band was detected in the material from the control STVbeads.

Hence, the 75-kDa polypeptide present in the VC1 pull-down assays is PT and is also the only Gladomain-containing protein detected in the pulldown from plasma using Western blot analysis, taken into consideration that its level is about one order of magnitude higher than the level of the other detected Gla-containing proteins.

PT is a direct interactor of VC1 and binding is mediated by the Gla-domain

To assess whether PT is a direct interactor of VC1, we repeated the pull-down using purified PT. As shown in the Coomassie-stained gel of **Fig. 1D**, VC1-STV beads captured a 75-kDa polypeptide that was absent in the material from the STV-beads alone. Thus, PT is a direct interactor of VC1.

The interaction of PT with VC1 is competed by Ca^{2+}

The occurrence of proteins sharing the Gla domain in the VC1 plasma interactome raised the hypothesis that the binding of VC1 to PT was mediated by this domain. The Gla domain is a calcium-binding module (K_d for calcium 0.3-0.5 mM) that in the calcium-bound form defines the binding of Gla-containing proteins to membrane phospholipids (33). In human PT, the Gla domain is an N-terminal region of 46 amino acids with 10 y-carboxyl glutamic acid residues (Gla) that are responsible for the binding to calcium ions. Upon binding calcium ions, the negative charges of the Gla residues are neutralized and the Gla domain undergoes conformational changes that sequester calcium ions in the interior of the structure. We reasoned that if the PT-VC1 interaction was mediated by electrostatic interactions between the positively charged V domain and the highly

negatively charged Gla-domain, calcium ions would affect the binding. Since the human plasma was devoid of free calcium, due to the use of EDTA to inhibit coagulation, we repeated the pulldown assays after re-addition of CaCl₂. As shown in Fig. 2A, in the presence of different concentrations of Ca^{2+} , the level of PT bound to VC1 decreased in a dose-dependent manner. Both Western blot analysis with anti-Gla and anti-PT antibodies confirmed that the 75-kDa polypeptide, whose intensity decreased, was PT (Fig. 2B and C). To rule out that the displacement of the interacting partner by calcium was simply due to the increase of the ionic strength of the environment, we compared the effect of the addition of calcium chloride to the PT-VC1 binding reaction with those obtained by addition of NaCl in amounts that yielded the same ionic strength (i.e. 10 mM of CalCl₂ is equivalent to 30 mM of NaCl) and no relevant effects on the VC1 pull-down were detected (data not shown).

Next, we tested whether addition of calcium could displace PT already bound to VC1-STV beads. Fifty mM CaCl₂ was capable to elute PT from the beads (Fig. 3A). Any residual material still bound to the resin was recovered by boiling the beads in SDS sample buffer. The majority of PT was released in the calcium-elution fraction (B1). To rule out any effect of 50 mM CaCl₂ on VC1 binding to the STV-beads, we stained a parallel gel with Coomassie (Fig. 3B). In the fraction eluted with calcium, where the majority of PT was released, only a tiny amount of VC1 was detected. The amount was negligible compared to that obtained by boiling the resin in SDS (Fig. 3B, lane B2 compared to B1). Thus, high concentrations of Ca²⁺ displace the Gla-domain of PT from VC1 without significantly affecting VC1 binding to the resin.

Finally, to further prove that the Gla domain was the site of interaction of PT with VC1, we repeated the pull-down assay using purified full-length PT and PT devoid of the Gla-domain, (PT des-Gla), obtained by treating PT with α -chymotrypsin (34) (**Fig. 4**). In contrast to the strong interaction of full-length PT, PT des-Gla showed no binding to VC1, corroborating the finding that the interaction of PT with RAGE occurs via the Gla-domain.

MST analysis of the molecular interaction of PT with VC1 or sRAGE

To further characterize the PT-RAGE interaction, we analyzed the binding of PT to VC1 by MicroScale Thermophoresis (MST), a powerful technique to analyze and quantify interactions between proteins in solution (35,36). In preliminary studies, the well-known RAGE ligands, S100B and S100A12 proteins, were used as positive control in MST and affinity values were similar to those obtained by Biacore/surface plasmon resonance assays, where one of the reactant is immobilized on a matrix (37) (data not shown).

Fig. 5A shows the binding curve of fluorescently labeled PT to VC1 in the absence of calcium. MST analysis revealed the formation of a 1:1 stoichiometry complex between the two proteins with an apparent dissociation constant (K_d) of 46.4 \pm 13.3 μ M, indicating a moderate affinity of PT for VC1.

In addition, we analyzed the binding of PT to human recombinant sRAGE, spanning from residue 23 to 322 and corresponding to the VC1-C2 domains. sRAGE was produced as a secreted glycoprotein in P. pastoris, as described under procedures. Experimental Purified **sRAGE** migrated as a smeared polypeptide of about 45-60 kDa in agreement with its glycosylated nature (Fig. S2). The binding reaction of PT to sRAGE showed a K_d of 4.4 μ M \pm 1.6 μ M, indicating that the affinity of PT for the ectodomain of RAGE is significantly higher than that for VC1 (Fig. 5A). As a next step, we tested the effect of calcium on PT binding to VC1 in MST assays. Increasing concentrations of calcium decreased the affinity of PT for VC1 and at the highest concentration, 20 mM CaCl₂, the binding was almost abolished (Fig. **5B**). Fig. S4 shows that also the binding of PT to sRAGE high was abolished at calcium concentration.

We also analyzed by MST the interaction of PT des-Gla with VC1 and sRAGE. PT des-Gla did neither bind to VC1 (**Fig. 5B**) nor to sRAGE (**Fig. S4**) providing further evidence that the Gla domain is crucial for the interaction.

VC1 inhibits plasma clotting

Since PT has a fundamental function in blood coagulation, we explored whether the interaction of PT with RAGE might modulate this process. To assess a potential role of VC1 in blood coagulation we measured plasma clotting by a conventional assay based on the use of activated partial thromboplastin time (aPTT) reagent to activate the intrinsic pathway. Different amounts of VC1 were added to the plasma prior the coagulation reaction was started. The clotting of the control initiated after 50 sec, which is typical in these assays. Remarkably, the presence of VC1 drastically prolonged the onset of coagulation in a dose dependent manner up to fourfold with an onset of the clotting after approximately 200 sec (**Fig. 6**).

To rule out any effect of VC1 on thrombin activity, VC1 was added to a thrombin activity assay using a substrate with a thrombin cleavage site and activity was unaffected by VC1 (data not shown).

Discussion

In this work, we demonstrated that the RAGE ligand-binding domain, VC1, and sRAGE bind PT via the Gla-domain. The Gla-domain is highly conserved among species and is present at the Nterminal end in proteins that undergo vitamin Kdependent γ -carboxylation of glutamate residues. This post-translational modification occurs in the ER and is essential for life (38). Among the Glacontaining proteins identified in this study, four (PT, protein S, protein Z, Factor IX) were reproducibly enriched as detected by MS analysis of VC1-interacting material from human plasma. The Gla-domain-containing factor VII escaped detection in the VC1 pull-down experiments, most likely due to the tiny circulating amounts in the plasma [levels are from (39) and from Enzyme Laboratories Research Inc. https://enzymeresearch.com/]. Indeed. the concentration of Factor VII is only 0.5 $\mu\text{g/ml}$ compared to the levels of circulating PT (90 μ g/ml), protein S (free: 10 μ g/ml, total: 25 μ g/ml), factor IX (5 µg/ml) and protein Z (2.2 µg/ml). Another Gla-domain-containing factor, Factor X (8 µg/ml) is nevertheless not reported in **Dataset** S1, since it was present both in the material bound to VC1- and control-beads, and according to the adopted criteria, was removed from the dataset of proteins specifically interacting with VC1. Regarding the interaction of VC1 with C4BP, it is remarkable to note that C4BP binds apoptotic and necrotic cells as well as DNA, to clean up after injury [30]. This interaction is mediated by the Gla domain of protein S and does not affect the ability of C4BP to inhibit the complement. Notably, the known RAGE ligand C3a [31] was also identified in our analysis.

No AGE or ALE modifications were detected in the identified proteins. This is not surprising since AGEs and ALEs require long time to form and therefore in physiological conditions accumulate in proteins with very low turnover rates such as collagen. The turnover of plasma proteins is much higher preventing an accumulation of AGEs in amounts compatible with the affinity of a pulldown assay (39). The common RAGE ligands, HMGB1 and S100 proteins, could not be identified since members of the S100 protein family and HMGB1, are small proteins of ~10-14 kDa (monomer) and 29 kDa, respectively, and in our study we restricted the analyses to the region of the gels containing polypeptide bands with molecular mass higher than 37 kDa. The reason of this choice was that in the elution step, STV (~14 kDa) and VC1 (~32-34 kDa) were released from the beads in high abundance and the identification of specific low-abundance VC1 ligands by MS, was not feasible (see Experimental procedures).

According to previous reports, proteins other than RAGE specifically interact with Gla-domaincontaining proteins through the Gla-domain. Factor X zymogen binds to the adenovirus serotype 5 hexon protein (Ad5 hexon) via the FX Gla domain and this interaction is required for the entry of the virus into hepatocytes (40). Another example is the superantigen-like protein 10 (SSL10) of Staphylococcus aureus, a bacterial exoprotein which directly binds to the Gla-domain of PT and of factor Xa, and binding is competed by Ca^{2+} (41). SSL10 has two short and highly basic segments (pI ~11 and pI ~9.8) that are responsible for the binding of SSL10 to PT (42). The interaction likely occurs between the highly positively charged segments and the highly negatively charged Gla-domain of PT. Thus, electrostatic attractions could dominate the bimolecular PT-SSL10 interaction.

The nature of interaction of VC1 with Gla-domain containing proteins seems to be similar to that reported for SSL10 and has a marked electrostatic nature. In particular, the V domain, a so called "basic trap" of RAGE, is rich in lysine and arginine residues that are positively charged at physiological pH. In addition, the structure of RAGE reveals the presence a shallow binding region that could provide a surface of interaction with the Gla-domain. The binding of PT via its Gla-domain to sRAGE was confirmed by pulldown and MST experiments corroborating the interaction of PT with VC1 domain in the context of the entire ectodomain. Interestingly, sRAGE displayed a higher affinity towards PT than VC1 alone (Fig. 5A) raising the question whether C2 domain is involved in the interaction. C2 domain of RAGE displays a net negative charge and interaction with the negatively charged Gladomain of PT is rather unlikely. Nevertheless, C2 domain might interact with other regions of PT and contribute to the stability of the complex. Furthermore, it was reported that the linker between C1 and C2 domain as well as the C2 domain contribute to dimerization of sRAGE (43,44) although in our MST set up and using the glycosylated sRAGE produced in *P.pastoris* we did not have evidence of a partial sRAGE dimerization in solution.

Ca²⁺ could effectively compete with RAGE-PT binding as observed for the interaction of SSL10 with PT. The competition effect of Ca²⁺ was clearly stronger than a simple increase of ionic strength. Gla-domain has an affinity for Ca²⁺ ions that mask the highly negative charges of the γ carboxylated-Glutamate residues and induce a transition from a disordered conformation to a structure in which the acidic residues complexed with calcium are buried inside. This contributes to confer PT the ability to bind membrane phospholipids (PS). We propose that these effects mask the Gla-domain from interacting with basic proteins, such as SSL10 or the "basic trap" V domain of RAGE.

The physiological importance of the binding of Gla-domain to calcium ions is well established contributing to the assembly of active coagulation factors complexes: Factor VIIIa-IXa that activates Factor X to factor Xa and factor Va-factor Xa complex (prothrombinase complex) on the surface of membranes and leads to efficient proteolytic conversion of PT to thrombin. For this reason, the interaction of the Gla-domain with VC1 impairs the binding to calcium and interferes with the activation of the coagulation factors causing a delay in plasma clotting. In this respect, the Gladomain could function as a sticky segment that interacts with positively charged domains. To address the exact physiological role of RAGE-PT interaction, further in vivo studies are required. It should be noted that the amount of circulating sRAGE in healthy subjects is too low (1.1-1.7 ng/ml) to affect blood coagulation at the ~1.2 mM free calcium concentration typical for blood. A different scenario could be envisaged in the lung. The basal membrane of alveolar Type I (ATI) cells of the lung, rich in membrane-bound RAGE, is in direct contact with the endothelium of capillary vases and with the interstitial space, and together constitute the capillary-alveolar membrane whose integrity is fundamental for gas-exchanges. In the adult, RAGE is mainly expressed in the ATI cells and it may be in contact with blood during the gas exchanges and also be present as sRAGE in the interstitial space of the capillary-alveolar membrane (13,45). In conditions of severe lung injury as acute lung injury (ALI) and/or acute respiratory distress syndrome (ARDS), losses of integrity of the alveolar-capillary membrane or changes in capillary permeability result in an extensive efflux of protein-rich oedema fluid into the alveolar air-space compartment. Herein, release cytokines, macrophages TNF- α , interleukins and a cascade is triggered that leads to induction of coagulation in the alveolar compartment and generates а pulmonary coagulopathy (46-48). In severe ALI the plasma level of sRAGE increases and reaches a concentration of 6-12.3 ng/ml. In oedema fluid of ALI/ARDS patients, sRAGE concentration is in the range of 0.9-3.2 μ g/ml (49). Therefore, in these diseases, sRAGE in the alveolar space may contribute in modulating coagulation bv sequestering PT from activation to thrombin. Alternatively, since thrombin plays not only a role in hemostasis but also in promoting fibrosis via thrombin receptors (50), the binding of PT to sRAGE could be beneficial in reducing/delaying

thrombin-signaling to fibrosis. Indeed, it is known that sRAGE exerts anti-fibrotic effects in a model of idiopathic pulmonary fibrosis (51). However, studies using RAGE constitutive KO homozygous mice are controversial. Lack of RAGE seems to either protect mice from lung fibrosis (50,52) or to enhance fibrosis in some models of induced pulmonary fibrosis (51). In addition, RAGE^{-/-} mice spontaneously develop pulmonary fibrosis with age (53). The model of disease used appears to affect the conclusions of these studies and the capability of the lung to respond to injuries.

In conclusion, the new interactors identified in this work may contribute in the future to deepen the understanding of the physiological functions of RAGE and its involvement in lung pathologies.

Experimental procedures

Plasma collection

At the recruitment, blood samples from six healthy volunteers were collected between 8 and 9 a.m., following at least 6 h of fasting. To inhibit coagulation, VACUETTE[®] EDTA 7 ml tubes containing K3EDTA or VACUETTE[®] 6ml- tubes containing a buffered sodium citrate solution at the concentration of 0.109 mol/1 (3.2%) mixed at a ratio of 1 part citrate solution to 9 parts blood, were used. Plasma-EDTA or plasma-citrate samples were obtained after centrifugation at 1200 g at room temperature for 15 minutes. The protein concentration of the supernatant was determined by BCA protein assay.

The studies were approved by the Ethical Committee of Ca' Granda Ospedale Maggiore Policlinico and University of Milan. All subjects gave their informed consent to participate in the studies. The studies abide by the Declaration of Helsinki principles.

VC1 pull-down assay

The VC1 portion of human RAGE was expressed and purified with a C-terminal tandem tag (His and Strep tag) from *Pichia pastoris* culture supernatant as previously described (26). The recombinant protein was immobilized on Streptavidin (STV)-coated magnetic beads (Streptavidin Mag SepharoseTM, GE Healthcare) by exploiting the affinity of the Strep tag for STV. In order to obtain the VC1-STV-beads, 50 μ g of purified VC1-His-Strep in 170 μ l of Buffer A (20 mM HEPES pH 7.1, 100 mM NaCl) were added to 5 μ l of packed STV coated-beads, previously equilibrated with the same buffer. 170 μ l of Buffer A were added to the same amount of beads in order to obtain the STV-beads. After 1 hour of incubation at 4 °C on a rotary mixer, the unbound material was carefully removed and the magnetic beads were washed twice with 500 μ l of Buffer A.

For pull-down assays, plasma samples were centrifuged at 4 °C for 10 min at 16000 rcf and then diluted to 5 mg/ml in buffer B (20 mM HEPES, 100 mM NaCl, 0.5 % Triton X-100 pH 7.1). The VC1-STV- and the STV-beads were incubated for 2 h at 4 °C with 160 µl of diluted plasma (called input). The unbound material was carefully removed and the beads were washed three times with 500 µl of Buffer B. Protein bound to VC1-STV and STV were eluted boiling the beads for 5 min in 15 µl of Laemmli Sample Buffer 4X [0.25 M Tris-HCl pH 6.8, 8% (w/v) SDS, 400 mM DTT, 40 % (w/v) glycerol and 0.04% Bromophenol blue]. The beads were washed with 15 μ l of Buffer A and the two eluates were pooled. For the pull-down assay performed in the presence of calcium, the same protocol was used, adding to buffer B different amount of CaCl₂ (1, 5 or 10 mM).

sRAGE expression and purification

Human sRAGE (residues 23-322) was produced, as an untagged protein, in P. pastoris from strain KM71H transformed with pPICZaA-sRAGE (54). Cells grown in MGY (1% yeast extract, 2% peptone, 1% glycerol, 1.34% YNB, 4x10-5 % biotin) were diluted to 1 OD_{600} and induced to produce sRAGE in MEMY (1% yeast extract, 2% peptone, 0.5% methanol, 0.7% Ethanol, 1.34% YNB, 4x10-5 % biotin) in flask at 30 °C and 0.5 % (v/v) MetOH was added daily. The culture supernatant from 400 ml-culture was collected after 72 h from induction and was concentrated to ~ 30 ml by ultrafiltration using a 10,000 cut-off membrane and the retentate was dialyzed overnight against 20 mM MES, pH 6. The solution was applied to a cation-exchange HITRAP SP HP

column (GE-Healthcare) equilibrated with the same buffer and connected to an AEKTA PRIME PLUS system (GE-Healthcare). After column washing, elution was performed with a 0 -1 M NaCl gradient in the same buffer. Fractions were analyzed by SDS-PAGE and fractions containing sRAGE were pooled and applied to a size exclusion chromatography Superdex 75 column (GE-Healthcare) equilibrated with 20 mM MES, 300 mM NaCl, pH 6, and connected to an AEKTA PRIME PLUS system (GE-Healthcare). Fractions were analyzed by SDS-PAGE and fractions containing sRAGE were pooled. Amicon centrifugal concentrators equipped with 3 kDacutoff membranes were used to exchange the buffer with 20 mM HEPES, 100 mM NaCl, pH 7.1.

Electrophoresis and immunoblotting procedures

Input, unbound and bound fractions obtained from pull-down experiments were analyzed by SDS PAGE. To 3 µl of input and unbound fractions, 12 µl of MilliQ water and 5 µl of Laemmli Sample Buffer 4X were added. The samples were denatured incubating for 5 min at 95 °C. SDS-PAGE was performed on Any KD TM Mini Protean® TGX TM precast gels. Protein bands were stained with Bio-Safe Coomassie blue (Bio-Rad). Images were acquired using the calibrated densitometer GS-800 and analyzed by the software Quantity one (Bio-Rad). For detection by Western blotting, proteins were transferred to nitrocellulose membranes using a Trans-Blot® Turbo[™] Transfer System (Bio-Rad). Proteins were detected using a mouse monoclonal antibody against thrombin (F-1) (Santa Cruz Biotechnology, dilution 1:500) or mouse monoclonal antibody against human ycarboxyglutamate (Gla) residues (Sekisui diagnostics, Cat# 3570, dilution 1:200) in combination with the peroxidase-conjugated antisecondary antibody (Santa mouse Cruz Biotechnology). Bound antibodies were revealed with the LiteAblot®PLUS Western blotting detection reagent (Euroclone, Italy).

Preparation of human Prothrombin des-Gla

Plasma derived prothrombin was purchased by Sekisui Diagnostics GmbH, Pfungstadt, Germany (Cat# ADG417 > 95% pure by SDS-PAGE and MS analysis). Gla-domain less PT (PT des-Gla) was obtained by proteolytic treatment essentially as described in (34). PT was diluted at 1 mg/ml in 50 mM HEPES, pH 8 and chymotrypsin (Sequencing grade, Roche) was resuspended at 1 mg/ml in 1 mM HCl (Pierce) and then diluted to 0.025 mg/ml in the same buffer. The reaction mix containing chymotrypsin and PT at a ratio of 1/1000 and a control reaction mix were incubated for 20 min at 37 °C. Proteolytic digestion was stopped by addition of N-tosyl-L-phenylalanine chloromethyl ketone (TPCK, 400 µM) and phenylmethylsulfonyl fluoride (PMSF, 2 mM). Aliquots of the reaction mixes were denatured at 100 °C for 5 min and analyzed by SDS-PAGE on 12% polyacrylamide minigels prepared using a solution of acrylamide/bis-acrylamide with ratio of 35/1. PT migrated as a single polypeptide with a Mr of 69,000 and after digestion totally converted to a slightly faster migrating band, that still reacted with anti-PT antibody but not with anti-Gladomain mAb. Thus, the whole Gla-domain was removed by this treatment in agreement with previous reports (34).

In gel digestion of polypeptides

After SDS-PAGE analysis of pull-down assays, areas of the gel above the 37 kDa marker were excised using a scalpel, finely chopped, transferred to a new Eppendorf and washed with 200 μ l of digestion buffer (50 mM ammonium bicarbonate). The region below the 37 kDa- marker was excluded since the detection of anything other than the highly abundant streptavidin and VC1 was unfeasible, despite the high sensitivity of the high-resolution spectrometer used. The reason of this was the dynamic protein concentration range in turn ascribable to the experimental needs.

An aliquot of 200 μ l of destaining solution (50% Ammonium bicarbonate 50 mM – 50% Acetonitrile) was added to each gel portion and heated at 37 °C for 10 min in the thermomixer (1400 rpm); the destaining solution was then discarded and this step was repeated until destaining was completed. Afterwards, gel pieces were incubated with 150 μ l of reducing solution (10 mM dithiothreitol in 50 mM ammonium bicarbonate) at 56 °C for 1 hour and then with 150 μ l of alkylating solution (55 mM iodoacetamide in

50 mM ammonium bicarbonate) at room temperature for 45 min in the dark. In-gel protein digestion was performed by two sequential digestions, one with 1 μ g of sequencing-grade trypsin (Roche) dissolved in digestion buffer and incubated for 16 hours, followed by a second digestion with a sequencing-grade chymotrypsin $(1 \mu g)$ for 7 hours at 25 °C in the presence of calcium chloride (10 mM). Peptide mixtures were extracted by 10 min-incubation with extraction solution (acetonitrile/trifluoroacetic acid/water; 30/3/67; v/v/v) and by additional 10 min incubation with pure acetonitrile. The two extracts were combined and dried in a vacuum concentrator (Martin Christ.). Digested peptide mixtures were then dissolved in an appropriate volume (20 µl) of 0.1% formic acid for mass spectrometry (MS) analysis.

Mass spectrometry analyses

The flowchart of Fig. S1 outlines the entire experimental design of the MS analyses of pulldown assays. Peptide mixtures from the in-gel digestion were separated by reversed-phase (RP) capillary liquid chromatography nanoscale (nanoLC) and analyzed by electrospray tandem mass spectrometry (ESI-MS/MS). For each analysis 5 µl of solubilized peptides were injected onto an Acclaim PepMap[™] C18 column (75 µm x 15 cm, pores 100 Å, Thermo Scientific, Waltham, Massachusetts, United States), protected by a precolumn, namely the Acclaim PepMapTM (100 µm x 2 cm, pores 100 Å, Thermo Scientific, Waltham, Massachusetts, United States). Samples were loaded onto the pre-column by means of the loading pump at 5 µl/min of mobile phase consisting of 99% of buffer A_LP (0.1% TFA) and 1% of buffer B_LP (0.1% HCOOH in CH₃CN) for 3 minutes. After the loading valve switching, peptide separation was performed by the NC_pump with a 91 minutes linear gradient of buffer B_NP (0.1% HCOOH in CH₃CN) from 1% to 40%, and further 12 minutes of linear gradient from 40% to 95% (Buffer B_NP). 9 minutes at 95% of buffer B_NP to rinse the column following the separative gradient, and the last 5 minutes served to re-equilibrate the column to initial conditions. The nano-chromatographic system, an UltiMate 3000 RSLCnano System (Dionex), was connected to an LTQ-Orbitrap XL mass

spectrometer (Thermo Scientific Inc., Milan, Italy) equipped with a Thermo Scientific dynamic Nanospray ion source set as follows: positive ion mode, spray voltage 1.7 Kv; capillary temperature 220 °C, capillary voltage 35 V; tube lens offset 120 V. The LTQ-Orbitrap XL mass spectrometer was operated in data-dependent acquisition mode (DDA) to acquire both the full MS spectra and the MS/MS spectra. Full MS spectra were acquired in "profile" mode, by the Orbitrap (FT) analyzer, in a scanning range between 300-1800 m/z, using a capillary temperature of 220 °C, AGC target = 5x105 and resolving power 60,000 (FWHM at 400 m/z). Tandem mass spectra MS/MS were acquired by the Linear Ion Trap (LTQ) in CID mode, automatically set to fragment the nine most intense ions in each full MS spectrum (exceeding 1×10^4 counts) under the following conditions: centroid mode, normal mode, isolation width of the precursor ion of 2.5 m/z, AGC target 1×10^4 and normalized collision energy of 35 eV. Dynamic exclusion was enabled (exclusion dynamics for 45 seconds for those ions observed 3 times in 30 seconds). Charge state screening and monoisotopic precursor selection were enabled, singly and unassigned charged ions were not fragmented. Xcalibur software (version 2.0.7, Thermo Scientific Inc., Milan, Italy) was used to control the mass spectrometer.

Protein Identification

The software Proteome Discoverer (version 2.2.0.338, Thermo Scientific, USA), implemented with the algorithm SEQUEST, was used to compare the experimental full and tandem mass spectra with the theoretical ones obtained by the *in silico* digestion of all the proteins within the UniProt *Homo sapiens* proteome database (https://www.uniprot.org), (UniProt release 2017_03 - March 15, 2017, Reviewed / Swiss-Prot 26,546 entries).

Trypsin and chymotrypsin were selected as the cleaving proteases, allowing a maximum of 2 missed cleavages. Peptide and fragment ion tolerances were set to 5 ppm and 10 mmu, respectively. Cysteine carbamidomethylation was set as fix modification (+57.02147), while methionine oxidation was allowed as a variable modification. As a quality filter, only peptide with an XCorr value grater then 2.2 for doubly-charged

peptides, 2.5 for triply-charged, 2.75 for quadruply charged peptide ions, and 3 for charge states quintuple or higher were considered as genuine peptide identifications. To ensure the lowest number of false positives, the mass values experimentally recorded were further processed through a combined search with the Database Decoy, where the protein sequences are inverted and randomized. This operation allows the calculation of the false discovery rate (FDR) for each match, so that all the proteins out of range of FDR between to 0.01 (strict) and 0.05 (relaxed) were rejected.

MicroScale Thermophoresis

The technique is based on thermophoresis, the directed motion of molecules in temperature gradients (36). Purified PT (Enzyme Research Laboratories) was labeled using the RED-NHS (Amine Reactive) NT-647 fluorescent dye (kit #MO-L001, NanoTemper Technologies) according to the manufacturer's instructions. Unreacted dye was removed with the supplied dye removal column equilibrated with 20 mM HEPES pH 7.1, 100 mM NaCl, 0.05% Tween 20. The labeled PT was adjusted to 200 nM with 20 mM HEPES pH 7.1, 100 mM NaCl, 0.05% Tween 20. Unlabeled recombinant VC1-His-Strep was subjected to 16 serial dilutions in 20 mM HEPES pH 7.1, 100 mM NaCl, 0.05% Tween 20 ranging from 666 µM to 20 nM. Unlabeled recombinant sRAGE was subjected to 16 serial dilutions in 20 mM HEPES pH 7.1, 100 mM NaCl, 0.05% Tween 20 ranging from 528 µM to 16 nM. Labeled prothrombin and VC1 or sRAGE were mixed at a volume ratio of 1:1, incubated at 24°C for 30 min and then loaded into standard capillaries (Monolith NT.115 Capillaries, NanoTemper Technologies). During a MST experiment, a temperature gradient is induced by an infrared laser. MST was measured using a Monolith NT.115 instrument (NanoTemper Technologies) at 24 °C. Instrument parameters were adjusted to 20 % LED power and medium MST power. Data of independently pipetted measurements were analyzed (MO.Affinity Analysis software version 2.3, NanoTemper Technologies) using the signal from an MST-on time of 1 s (hot region from 4 s to 5 s). Relative fluorescence is used to quantify binding via MST. To obtain ΔF , the baseline F value (F is

the fluorescence) is subtracted from all data points of the same curve. The baseline F value is equivalent to the mean F value of the unbound target, usually in capillaries 14-16, and is given by the MO.Affinity Analysis software as 'unbound' value when a fit is performed. Thus, by definition, ΔF is 0 in the unbound state.

The fraction bound corresponds to the value of the baseline corrected normalized fluorescence (Δ F) divided by the curve amplitude of each data point. The fraction bound (FB) is described by the equation:

 $F_{norm} = (1-FB)F_{norm unbound} + (FB)F_{norm bound}$

where Fnorm unbound is the normalized fluorescence of the unbound state and Fnorm bound is the normalized fluorescence of the bound state. From FB, the value $Kd = [A]^*[B]/[AB]$ of a binding reaction between a partner A and a partner B to form the complex AB can be derived as described in (36). In **Fig. S4**, where no binding is present, Fnorm was used to compare the curves and Fnorm is defined as the F1/F0 ratio where F0 corresponds to the normalized fluorescence prior to MST activation (cold region) and F1 is the normalized fluorescence after MST activation (hot region from 4 s to 5 s).

Plasma clotting assay

The effect of VC1 on blood coagulation was examined by plasma clotting assay. Varying amounts of VC1 in 10 µl of 20 mM HEPES, 100 mM NaCl pH 7.1 were added to 100 µl of citrated human plasma (Vericon 1 Normal Control Plasma, HART Biologicals) and incubated for 30 min at room temperature. After addition of 10 µl of aPTT reagent (Ultrasense EA, HART) and incubation for 2 min at 37 °C, the mixture was centrifuged at 6000 rpm at room temperature for 1 min. 110 µl of supernatant were carefully transferred to a microtiter plate. The clotting reaction was startedby the addition of 20 µl of 25 mM CaCl₂. The absorbance at 405 nm was monitored for 10 min every 5 s using a microplate reader (EnSight Multimode Plate Reader, PerkinElmer).

Data availability: MS proteomics data have been deposited to the ProteomeXchange Consortium (http://proteomecentral.proteomexchange.org) via the PRIDE partner repository (55) with the dataset identifier PXD020355. All remaining data are contained within the article.

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String accession	Accession	Description	Gla domain	MW [kDa]	positive identification
F2	P00734	Prothrombin	+	70	8
C4BPA	P04003	C4b-binding protein alpha chain		67	8
PROS1	P07225	Vitamin K-dependent protein S	+	75.1	8
C4A	P0C0L4	Complement C4-A		192.7	8
F9	P00740	Coagulation factor IX	+	51.7	6
A2M	P01023	alpha-2-macroglobulin		163.2	6
VTN	P04004	Vitronectin		54.3	5
THBS1	P07996	Thrombospondin-1		129.3	5
ITIH2	P19823	Inter-alpha-trypsin inhibitor heavy chain H2		106.4	5
COMP	P49747	Cartilage oligomeric matrix protein		82.8	5
IGHG2	P01859	Immunoglobulin heavy constant gamma 2		35.9	4
IGHM	P01871	Immunoglobulin heavy constant mu		49.4	4
PROZ	P22891-2	Isoform 2 of Vitamin K-dependent protein Z	+	47	4
C3	P01024	Complement C3		187	3
IGHA1	P01876	Immunoglobulin heavy constant alpha 1		37.6	3
FGA	P02671-1	Fibrinogen alpha chain		94.9	3
F5	P12259	Coagulation factor V		251.5	3
FRK	P42685	Tyrosine-protein kinase FRK		58.2	3
ITIH3	Q06033-1	Inter-alpha-trypsin inhibitor heavy chain H3		99.8	3
TTN	Q8WZ42-12	Isoform 12 of Titin		3992.1	3

Table 1. Human	plasma	proteins	interacting	with V	C1 in p	pull-down-assays

The bound material of VC1 pull-down assays from plasma samples was analyzed by LC/MS. Out of 295 proteins identified and listed in the **Dataset S1**, 20 were selected using a cut-off of \geq 3 positive identifications over 8 pull-down assays.

Peptide Sequence ^a	PSMs ^b	XCorr ^c	Charge ^d	MH+ [Da] ^e	ΔM [ppm] ^f
¹⁰¹ AAcLEGNcAEGLGTNY ¹¹⁶	2	3.64	2	1699.70293	-1.29
¹⁰⁵ EGNcAEGLGTNYR ¹¹⁷	2	3.35	2	1440.61601	-0.88
¹⁷⁸ RQEcSIPVcGQDQVTVAmTPR ¹⁹⁸	2	6.49	3	2448.14023	-1.11
¹⁹⁹ SEGSSVNLSPPLEQcVPDR ²¹⁷	2	3.94	2	2070.97295	-1.55
²⁰⁷ SPPLEQcVPDR ²¹⁷	2	3.00	2	1297.61797	-2.01
²⁰⁷ SPPLEQcVPDRGQQY ²²¹	2	5.31	2	1773.82146	-0.60
²⁴⁸ HQDFNSAVQLVENF ²⁶¹	2	3.75	2	1647.77519	-0.62
²⁵² NSAVQLVENF ²⁶¹	2	3.61	2	1120.56145	-1.71
²⁷⁶ VAGKPGDFGY ²⁸⁵	2	3.10	2	1010.49156	-2.63
²⁸⁹ NYcEEAVEEETGDGLDEDSDR ³⁰⁹	2	4.87	2	2431.93169	0.07
²⁹¹ cEEAVEEETGDGLDEDSDR ³⁰⁹	2	5.11	2	2154.82280	-1.15
³³⁰ GSGEADcGLRPL ³⁴¹	2	3.73	2	1231.57231	-1.07
³⁶⁴ IVEGSDAEIGMSPW ³⁷⁷	2	3.37	2	1490.67912	-2.77
³⁹¹ cGASLISDR ³⁹⁹	2	2.95	2	978.46599	-1.40
⁴¹⁸ TENDLLVR ⁴²⁵	2	3.17	2	959.51494	-0.77
⁴³⁵ ERNIEKISML ⁴⁴⁴	1	3.21	2	1232.66509	-1.39
⁴⁷⁵ SDYIHPVcLPDR ⁴⁸⁶	4	3.30	3	1471.69828	-1.09
⁴⁷⁸ IHPVcLPDRETAASL ⁴⁹²	2	3.25	2	1678.85649	-1.01
⁵²⁴ QVVNLPIVER ⁵³³	2	2.91	2	1166.68779	-1.24
⁵²⁴ QVVNLPIVERPVcK ⁵³⁷	4	3.87	2	1650.93450	-0.94
⁵⁵⁴ KPDEGKR ⁵⁶⁰	2	2.90	2	829.45293	0.29
⁵⁶⁰ RGDAcEGDSGGPF ⁵⁷²	2	4.19	2	1324.52056	-1.34
⁵⁶¹ GDAcEGDSGGPF ⁵⁷²	2	3.24	2	1168.41997	-1.07
⁶¹³ IQKVIDQFGE ⁶²²	2	3.69	2	1176.62395	-1.72

Table 2: Identification of the 75-kDa polypeptide as prothrombin

MS fingerprinting of the ~75 kDa band excised from the gel of the VC1-bound material fraction separated by SDS-PAGE as shown in **Fig. 1A**.

^aThe amino acid residues refer to the precursor sequence of prothrombin from *Homo sapiens* (UniProt accession number P00734, amino acids 1-622). Sequence coverage was 35.37%. Lowercase c indicates cysteine carbamidomethylation and lowercase m indicates methionine oxidation.

^bPSMs: peptide spectrum matches (total number of identified peptide sequences including those redundantly identified).

^cXCorr score values from Sequest algorithm.

^dCharge state of the precursor ion.

^eMH+ [Da] represents the singly charged mass of the precursor ions in Daltons.

^fDifference between the theoretical mass of the peptide and the experimental mass of the precursor ion.

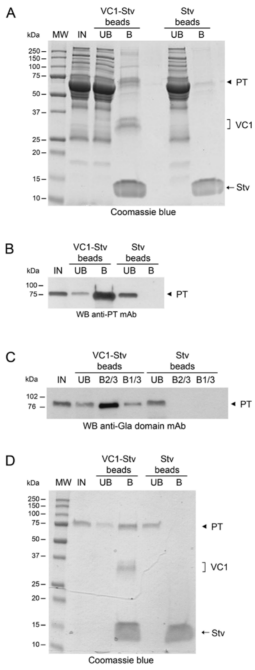


Figure 1. Prothrombin (PT) is the most abundant RAGE-interactor detected in human plasma by VC1 pull-down assay. (A) SDS-PAGE of the fractions from pull-down assays carried out in 20 mM HEPES, 100 mM NaCl, 0.5 % Triton X-100 pH 7.1. IN: input; UB: unbound fraction; B: bound material obtained after boiling the beads for 5 min in SDS sample buffer. The arrowhead indicates the 75 kDa-polypeptide identified as PT and present in the VC1-STV beads and absent in the control (STV-beads). VC1 and STV are also indicated. Proteins were stained by Coomassie blue. (B) Western blot analysis using Anti-PT mAb of fractions from a VC1 pull-down assay. (C) Western blot analysis of a VC1 pull-down assay using Anti-Gla domain mAb. B2/3 and B1/3 indicate that 2/3 and 1/3 of the total bound material were loaded. (D) VC1 Pull-down assay using purified PT instead of plasma. The indicated samples were analyzed by SDS-PAGE and proteins were stained by Coomassie blue.

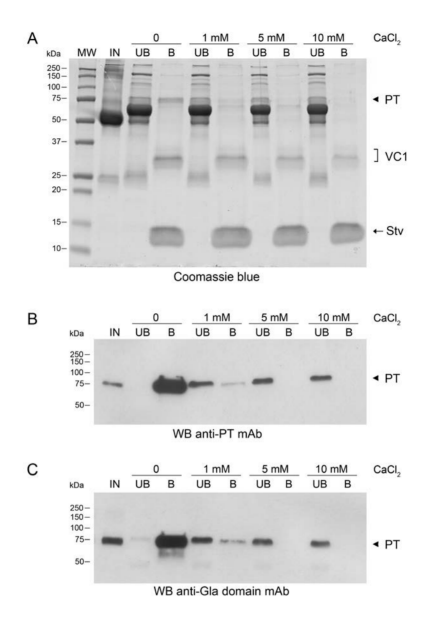


Figure 2. Competition in PT binding to VC1 by CaCl₂. The pull-down assays were carried out in 20 mM HEPES, 100 mM NaCl, 0.5 % Triton X-100 pH 7.1, before and after addition to plasma of increasing concentrations of CaCl₂. IN: input; UB: unbound fraction; B: bound material obtained after boiling the beads for 5 min in SDS sample buffer. (A) Proteins were analyzed by SDS-PAGE and stained by Coomassie blue. (B) Western analysis of the fractions of pull-down assays using anti-PT mAb. (C) Western analysis of the fractions from pull-down assays using anti-Gla mAb. Arrowhead: prothrombin (PT). VC1: VC1 domain; Stv: streptavidin.

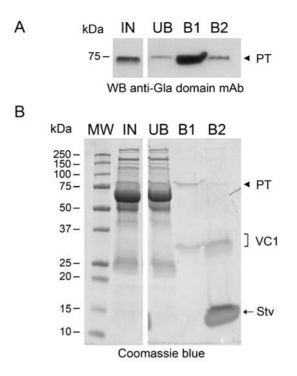


Figure 3. Calcium at high concentration displaces PT from the complex with VC1. (A) Western analysis with anti-Gla domain mAb of fractions from VC1 pull-down assays carried out in 20 mM HEPES, 100 mM NaCl, 0.5 % Triton X-100 pH 7.1. IN: input; UB: unbound fraction; B1: bound material obtained after incubating the beads with 50 mM CaCl₂; B2: residual bound material obtained by boiling beads from B1 for 5 min in SDS sample buffer. (B) Coomassie stained gel of the pull-down assays obtained in the condition described in (A). The arrowhead indicates prothrombin.

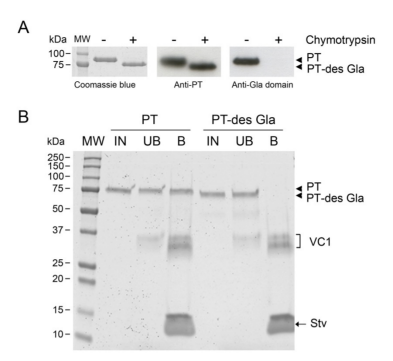


Figure 4. The Gla domain mediates the interaction of PT with VC1 in pull-down assay. (A) Proteolytic treatment of purified PT removed the Gla domain and produced PT des-Gla. (B) Coomassie-stained gel of VC1 pull-down assays performed in 20 mM HEPES, 100 mM NaCl, 0.5 % Triton X-100 pH 7.1 (no CaCl₂), in the presence of PT or PT des-Gla. Fractions were labeled as IN: input; UB: unbound fraction; B: bound material obtained after boiling the beads for 5 min in SDS sample buffer.

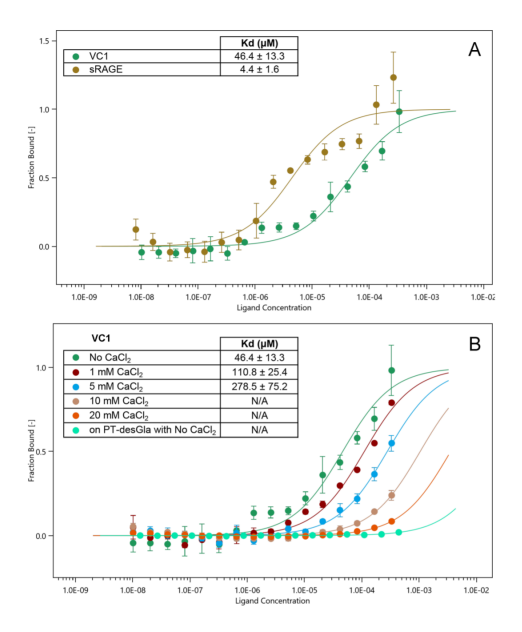


Figure 5. Molecular interaction of PT with VC1 and with sRAGE. (A) MST measurements were carried out in 20 mM HEPES pH 7.1, 100 mM NaCl, 0.05% Tween 20 and standard capillaries with a constant concentration of fluorescently labeled prothrombin (100 nM) mixed with varying amounts of unlabeled VC1-His-Strep (final concentrations from 10.2 nM to 333 μ M) or sRAGE (from 80.6 nM to 264 μ M). After 30 min-incubation at 24 °C, the samples were loaded into Monolith NT.115 capillaries (NanoTemper Technologies) and measurements were performed using the Monolith NT.115 (NanoTemper Technologies) at 20 % LED power and medium MST power. The values of the dissociation constant (K_d) derived from the curves for the PT-VC1 interaction (n = 5 independent measurements) and for the PT-sRAGE interaction (n = 4 independent measurements) are indicated in the inset. (B) MST measurements with VC1 were performed as in (A) but in the presence of increasing concentrations of CaCl₂ or in the presence of fluorescently labeled PT des-Gla instead of PT (n = 3 independent measurements). N/A: not applicable.

In A and B, the fraction bound is plotted against the log_{10} of VC1 or sRAGE molar concentrations. The error bars on individual data points represent the standard deviations among the repeats.

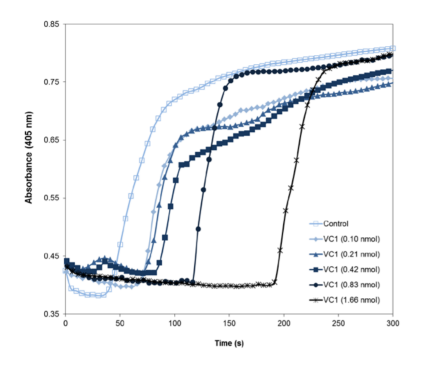


Figure 6. Effects of VC1 on plasma clotting induced by Ca^{2+} . Plasma-citrate was pre-incubated with various amounts of VC1 (0.01-1.66 nmol in 100 µl of plasma) and the intrinsic pathway was activated by the addition of aPTT reagent (10 µl). After addition of 20 µl of 25 mM CaCl₂, plasma clotting was monitored by measuring the absorbance at 405 nm using a microtitre plate reader. Time course curves are representative of three independent experiments.