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Leth, Julie M; Mertens, Haydyn D T; Leth-Espensen, Katrine Zinck; Jørgensen, Thomas J D; Ploug, Michael

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Did evolution create a flexible ligand-binding cavity in the urokinase receptor through deletion of a plesiotypic disulfide bond?

The urokinase receptor (uPAR) is a founding member of a small protein family with multiple Ly6/uPAR (LU) domains. The motif defining these LU domains contains five plesiotypic disulfide bonds stabilizing its prototypical three-fingered fold having three protruding loops. Notwithstanding the detailed knowledge on structure-function relationships in uPAR, one puzzling enigma remains unexplored. Why does the first LU domain in uPAR (DI) lack one of its consensus disulfide bonds, when the absence of this particular disulfide bond impairs the correct folding of other single LU domain-containing proteins? Here, using a variety of contemporary biophysical methods, we found that reintroducing the two missing half-cystines in uPAR DI caused the spontaneous formation of the corresponding consensus 7–8 LU domain disulfide bond. Importantly, constraints due to this cross-link impaired (i) the binding of uPAR to its primary ligand urokinase and (ii) the flexible interdomain assembly of the three LU domains in uPAR. We conclude that the evolutionary deletion of this particular disulfide bond in uPAR DI may have enabled the assembly of a high-affinity urokinase-binding cavity involving all three LU domains in uPAR. Of note, an analogous neofunctionalization occurred in snake venom α-neurotoxins upon loss of another pair of the plesiotypic LU domain half-cystines. In summary, elimination of the 7–8 consensus disulfide bond in the first LU domain of uPAR did have significant functional and structural consequences.

The urokinase-type plasminogen activator receptor (uPAR) is an extracellular membrane protein composed of three homologous Ly6/uPAR-type (LU) domains and a C-terminal glycosylphosphatidylinositol (GPI) membrane anchor (1). It serves to focalize plasminogen activation on cell surfaces via its high-affinity binding to the urokinase-type plasminogen activator (uPA) (2). In so doing, it facilitates extravascular fibrin surveillance reducing the adverse effects of chronic inflammation caused by unremitting fibrin deposition (3). Besides promoting pericellular proteolysis, the uPA-uPAR interaction also stimulates cell adhesion and migration via direct and indirect interactions with vitronectin and integrins (4–9). Elegant transgenic mouse models show that the interaction between uPA and uPAR promotes hepatic fibrin clearance (3) and improves neuronal recovery after either cerebral ischemia (10, 11) or spinal cord injury (12). Notwithstanding these beneficial effects, the uPA-uPAR interaction may also elicit detrimental pathological effects, particularly in relationship to chronic inflammation. In genetic mouse models, the interplay between uPA and uPAR augments the pathogenesis of collagen-induced arthritis (13, 14). In line with these causal correlations, high plasma levels of shed uPA/uPAR predict poor prognosis in several pathologic conditions with inflammatory lesions, e.g. bacterial infections (15, 16), kidney disease (17, 18), and invasive and metastatic solid cancers (19). The latter association spurred a considerable interest in developing uPAR-specific targeting strategies intended for use in cancer therapy (20–24). These initiatives are now being supplemented by the development of uPAR-targeting probes for noninvasive imaging of uPAR expression using either (i) positron emission tomography to guide patient staging (25–27) or (ii) near-IR fluorescence to guide precision cancer surgery by improving margin resection (28–30).

Crystal structures of uPAR solved in complex with its natural protein ligands (Fig. 1, C and D) (31–34), small molecule antagonists (21, 35, 36), or antibodies (31, 37) reveal that all three LU domains in uPAR combine to form a compact globular structure. This assembly creates (i) a large and hydrophobic uPA-binding cavity comprising elements from all three LU domains and (ii) a smaller peripheral binding site for the somatomedin B (SMB) domain of vitronectin at the interface between the first (DI) and second (DII) LU domain in uPAR. Although these binding sites are nonoverlapping they do, nevertheless, interact cooperatively. Prior uPA-binding thus increases uPAR’s affinity...
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for vitronectin (7) and in so doing it leads to increased cell adhesion and migration (5, 8, 9). We showed previously that the dynamic assembly of the LU domains in uPAR enables this allosteric regulation of ligand binding (5, 38). Combining our biophysical and functional data led us to conclude that uPA occupancy drives uPAR into a more closed and compact conformation and this increases the affinity for SMB (5). Locking uPAR permanently in this compact conformation, by introducing a nonnative disulfide bond between DI and DIII, bypassed the cooperativity of uPA binding and generated a constitutive high-affinity binding site for SMB (6, 39). Our data, furthermore, suggested that uPAR DI plays a dominating role in this flexibility (38). Given that this particular domain differs from the otherwise invariant LU-consensus motif, inasmuch as it lacks one of the 5 plesiotypic disulfide bonds (40), we speculate that the loss of this structural constraint could have been instrumental for evolving high-affinity uPA binding and gaining cooperativity in vitronectin binding. This proposition is not unprecedented as snake venom α-neurotoxins (representing secreted single LU-domain proteins) developed potent neurotoxicity toward synapsid targets and underwent neofunctionalization after deletion of a plesiotypic disulfide bond (41). In this study, we therefore introduced the lacking consensus disulfide bond in human uPAR DI and studied the consequences thereof on the structural flexibility and ligand binding properties of intact uPAR.

Results

Loss of a consensus disulfide bond in uPAR DI

Sequence alignments of the three homologous LU domains in human uPAR clearly show that the 5-disulfide bond signature, considered a plesiotypic trait of ancient three-fingered neurotoxins (41), is maintained in both uPAR DII and uPAR DIII (Fig. 1A). This primordial disulfide pattern was first identified in the basal-type α-neurotoxins (e.g. denmotoxin) that are present in venoms of nonfront-fanged snakes (Colubridae) that feed preferentially on nonmammalian prey (41, 42). In accordance with this feeding behavior, their α-neurotoxins have only weak affinity for mammalian nicotinic acetylcholine receptors (41, 42). In the more advanced elapid snakes, the α-neurotoxins gained high affinity for mammalian acetylcholine receptors via deletion of the 2–3 LU consensus disulfide bond in loop 1 (41, 42), as illustrated in Fig. 1A. Intriguingly, this sequence alignment and our experimental disulfide assignments (31, 32, 35, 40) reveal that uPAR DI unexpectedly lacks another of the invariant disulfide bonds that defines the ancient LU protein domain fold. In this case, the missing disulfide (denoted 7–8 in Fig. 1) is located at the base of loop 3 connecting β-strands E and F (Fig. 1, E and F) and is absent from DIs of all known mammalian orthologues of uPAR (Fig. S1). The deletion of this particular disulfide bond in uPAR during evolution is remarkable given that single LU domain-containing proteins, such as GPIHBP1, CD59, and κ-bungarotoxin require this disulfide bond for their correct protein folding and function (43–45).

Reintroducing the missing disulfide bond in uPAR DI

To assign the most probable site(s) for reintroducing the primordial 7–8 consensus disulfide bond in uPAR DI, we examined primary sequence alignments of individual LU domains. This clearly pinpointed Thr51 and Val70 as the most promising candidate pair (Fig. 1A, Figs. S1 and S8). This notion was further substantiated by the pairwise Cβ–Cβ atom distances in seven different crystal structures available for uPAR in complex with various ligands (Table S5). The Cβ–Cβ distances of these Thr51–Val70 pairs were 6.4 ± 0.5 Å, which is slightly longer than the distances for the bona fide 7–8 consensus disulfide bonds present in DI (4.0 ± 0.2 Å) and DIII (3.9 ± 0.4 Å). Nonetheless, evaluations focused only on minimizing structural perturbations highlighted yet another possible candidate pair, as the Cβ–Cβ atoms for Lys28 and Val70 were only 5.0 ± 0.4 Å apart.

Based on these considerations, we chose to express both uPAR T51C·V70C and uPAR K50C·V70C (residues 1–283) in Droso-
hila S2-cells and purify the secreted proteins. To confirm the oxidation status of the introduced cysteine residues (i.e. validating that they are indeed engaged in disulfide bond formation), we subjected uPAR to limited proteolysis with chymotrypsin under nondenaturing conditions. We optimized the conditions to hydrolize predominately the Tyr87–Ser88 peptide bond in the linker region between DI and DII and to a lesser extent the Tyr57–Arg58 peptide bond located in loop 3 of DI. Mass spectrometry confirmed that all cysteine residues in these protein preparations were engaged in disulfide bonding (Table 1, Fig. S2).

Introducing the 7–8 disulfide bond desensitizes DI to enzymatic deglycosylation and limited proteolysis in intact uPAR

We suspected that the inherent flexibility of DI as well as its assembly with DII and DIII were likely to be perturbed by the constraints introduced by the additional disulfide bond. Several independent lines of experimental evidence support this assumption.

Introducing the 7–8 disulfide bond rendered the glycan attached to DI (Asn52) resistant toward PNGase F-mediated release, when incubating intact uPAR with the deglycosidase under nondenaturing conditions. As shown in Fig. 2A, PNGase F readily removed the glycan attached to DI from full-length uPARwt, but was unable to do so in uPAR T51C·V70C. Likewise, uPAR K50C·V70C was a very poor substrate for PNGase F. Notably, a similar resilience toward PNGase F was acquired by uPARwt when it was driven into its closed conformation either by uPA binding (46) or by insertion of a remote interdomain disulfide bond (uPAR H47C·N259C, Fig. 1G) (6). The accessibility of the glycan on Asn52 to PNGase F was thus markedly compromised by the structural constraints originating from the introduction of the 7–8 disulfide bond between β-strands IE and IF (Fig. 1F) and this recapitulated the properties observed for uPAR in complex with uPA. All glycans attached to DII or DIII proved resistant to PNGase F in all tested uPAR variants.

Probing the various uPAR mutants by limited proteolysis with chymotrypsin revealed that the cleavage efficiency of the exposed linker region between DI and DII (Tyr87–Ser88) was largely unaffected by the constraints from the additional disulfide bonds or by occupancy with GDF (Fig. 2B). In contrast, the sensi-
Activity toward a secondary cleavage site at Tyr57–Arg58 within loop 3 proved markedly different between the various uPAR mutants. Although uPART51C-V70C appeared slightly more prone to this cleavage compared with uPARwt, uPARH47C-N259C was refractory. We suspect that this resistance reflects that DI remains fully integrated with DIIDIII in uPARH47C-N259C despite having a cleaved linker region between DI and DII (6). Maintaining the compact globular three-domain assembly of intact uPAR would thus shield Tyr57–Arg58 from proteolysis. Aligned with that proposition, we observed that GFD occupancy also prevents the cleavage at this position (Fig. 2B). An unexpected fuzziness in the electrophoretic mobility of the 1–57 fragment from uPART50C-V70C complicated the kinetic evaluation of the second chymotrypsin cleavage event in this mutant. The origin of this aberrant mobility remains unclear. Mass spectrometry of uPART50C-V70C digested for the 24 h showed only the expected mass of DI with one internal pep-

**Figure 1. Sequence alignment of LU domains in human uPAR and snake venom α-neurotoxins.** A shows an alignment of primary sequences for the three LU domains in uPAR (Homo sapiens, Q03405) and the single LU domains in the snake venom toxins: demotoxin (Boiga dendrophilia, DQ366293), erabutoxin a (Laticuda semifasciata, P60775), and α-cobratoxin (Naja kauthia, P01391). Linker regions and extensions are omitted from the alignment (their presence is indicated by ). Half-cystines are highlighted in yellow boxes along with their disulfide connectivity. Arrows indicate Thr51 and Val70 in uPAR DI. Residues located in the ligand-binding interface in crystal structures of ATF-uPAR (34) and α-cobratoxinAChBP (71) complexes are highlighted in green, as are residues important for neurotoxicity of erabutoxin a (72). The crystal structure of uPAR is shown in B as a cartoon representation (DI, cyan; DII, wheat; DIII, blue). C shows the ATF-uPAR complex with uPAR in a gray surface representation and ATF (containing GFD and a kringle domain) in cartoon representation. D shows the ATF-uPAR-SMB complex. E shows the LU domain in uPAR DI (residues 1–77) with β-sheets in cyan and disulfide bonds as yellow sticks. F shows the same structure tilted 90° to illustrate their structural constraint on the β-sheets and the proximity of the N-linked glycosylation site (Asn52) and the lacking 7–8 consensus disulfide bond. F shows the positions of the introduced disulfide bonds: Thr51–Val70 (*) and His47–Asn259 (**). Protein structures were created with PyMol (Schrödinger, LLC) using the PDB code 3BTI.
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| Table 1 Verification of disulfide bond status by MS |
|-----------------------------------------------|
| uPAR DI (residues 1–87) was excised from intact uPAR 1–283 by limited chymotrypsin digestion and the molecular masses were determined by LC-ESI-MS and maximum entropy (MaxEnt1) deconvolution of the charge state distributions of the proteins (settings, Gaussian FWHM: 1.0 and resolution: 0.25 Da/channel). The calculated masses are from the primary sequences including a paucimannosidic glycan (Man3GlcNAc2Fuc; 1,038.5 Da) tethered to Asn52 in S2-cells produced uPAR.

| uPAR1–87 | Measured mass | Calculated mass | ΔMass |
|----------|---------------|----------------|-------|
| wt       | 10,792.00     | 10,792.04      | 0.04  |
| K50C–V70C| 10,769.05     | 10,769.01      | 0.04  |
| T51C–V70C| 10,796.15     | 10,796.08      | 0.07  |
| C6S–C12S | 10,752.20     | 10,761.93      | 18.27 |

* This mass difference represents uPAR1–87 with a hydrolyzed peptide bond (Tyr57–Arg58), but the fragments (1–57 and 58–87) remain covalently attached via the introduced disulfide bond between Cys53 and Cys70.

The elution profiles from the size exclusion column revealed: (i) that uPAR DI released by limited proteolysis from uPARwt remains partly associated to DIIDIII (Fig. 3A); (ii) that prior occupancy with GFD greatly enhances this association (Fig. 3C); and (iii) that introduction of the 7–8 consensus disulfide bond completely eliminates the noncovalent association of DI and DIIDIII (Fig. 3, B and D).

Flexible in the globular three-domain assembly of uPAR by small angle X-ray scattering

To gain further insights into possible differences in the interdomain flexibility between the various disulfide-constrained uPAR conformers, we performed small angle X-ray scattering (SAXS) analyses. We analyzed both uPAR and uPAR-ATF complexes to compare the domain flexibility before and after ligand-induced compaction of the receptor. To maximize sample monodispersity, we fractionated uPAR and uPAR-ATF complexes by size exclusion chromatography before collecting batch scattering data by SAXS. From the normalized scattering data, we first derived the intra-particle distance distribution function, \( p(r) \) providing model-independent information on the shape parameters (Fig. 4). Comparison of the different unoccupied uPAR mutants revealed that only uPAR\(^{H47C-N259C}\) exhibits a symmetrical, bell-shaped \( p(r) \) function indicative of a compact and globular structure with a radius of gyration \( R_g \) of 22.4 ± 0.1 Å and a maximal particle dimension \( D_{max} \) of 70 ± 5 Å (Table S2). In contrast, reintroducing the missing 7–8 consensus disulfide in uPAR DI did not lead to large scale changes in the overall shape parameters, the \( p(r) \) functions for uPAR\(^{K50C-V70C}\) and uPAR\(^{T51C-V70C}\) are almost superimposable onto that of uPAR\(^{wt}\) (Fig. 4B). Upon ATF binding, all uPAR disulfide conformers compacted into similar sized particles with a \( D_{max} \) of 90 Å (Fig. 4D, Tables S1–S4). A small shift to higher distances in the \( p(r) \) function of the uPAR\(^{wt}\)-ATF complex is observed relative to ATF complexes with the uPAR disulfide mutants, but whether this reflects a significant structural difference is unclear. Nonetheless, transforming the scattering data into dimensionless Kratky plots provides a clear ranking of the unoccupied uPAR disulfide conformers into three groups according to their degree of flexibility. This transformation of the SAXS data are particularly useful to obtain a semi-quantitative analysis of the propensity of a given protein to adopt a globular fold (represented by a bell-shaped curve.
with peak at 1.104 for $qR_g = \sqrt{3}$ or be intrinsically disordered (represented by a hyperbolic curve with a plateau around $qR_g$ values of 1.5–2.0). From the Kratky plots presented in Fig. 5A, uPAR\textsuperscript{wt} exhibits the greatest flexibility and uPAR\textsuperscript{H47C-N259C} is stable and globular. Both uPAR variants with an intact 7–8 disulfide bond show an intermediate flexibility (Fig. 5A). In the presence of ATF, this difference in flexibility largely disappears, as illustrated by the Kratky plots in Fig. 5B. SANS-driven ensemble modeling of the scattering data by the ensemble optimization method (EOM) yielded a similar conclusion regarding the flexibility of unoccupied uPAR (Fig. 5, C and D). In this analysis, a homogeneous ensemble of relatively compact structures provides an excellent fit to the scattering data for uPAR\textsuperscript{H47C-N259C} with low flexibility metrics ($R_{\text{flex}}$ of 47%, relative to the threshold of randomness: 85%, determined from the random pool) calculated from the probability distributions (47). In contrast, a heterogeneous ensemble comprising both extended and compact conformations with a relatively high $R_{\text{flex}}$ of 81% is required to fit the scattering data for uPAR\textsuperscript{wt}. Intermediate ensemble compositions with $R_{\text{flex}}$ values of 74 and
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73% provide good fits to uPAR$^{T51C-V70C}$ and uPAR$^{K50C-V70C}$, respectively. These properties are also evident from the variations in shape observed in the refined ab initio surface envelopes reconstructed from the experimental SAXS data (Fig. S4). Extended models are generated for the highly flexible variants (uPAR$^{wt}$, uPAR$^{T51C-V70C}$, and uPAR$^{K50C-V70C}$), in contrast to the compact models generated for the less flexible uPAR$^{H47C-N259C}$ and all uPAR-ATF complexes.

Dynamics of uPAR DI

Previously we showed with hydrogen-deuterium exchange MS (HDX-MS) that uPAR DI undergoes a pronounced change in flexibility during the compaction of intact uPAR that occurs on uPA binding (38). Due to the covalent tethering of βE and βF by the 7–8 consensus disulfide bond in LU domains, we suspected that it could stabilize the βE-sheet between strands E and F in uPAR DI and this could in part be reconciled with the reduced flexibility observed in the Kratky plot of the SAXS data (Fig. 5A). To probe this proposition further, we performed a continuous deuterium labeling of the different uPAR disulfide variants in the presence or absence of saturating levels of GFD and determined the deuterium uptake values by MS after pepsin digestion with special emphasis on peptide(57–66). The deuterium uptake plots for this peptide reveals a considerable variability in the flexibility of βE and βF in the different disulfide variants (Fig. 6A). Interestingly, the ranking of the deuterium uptake when uPAR is driven into its compact state by uPA binding (38). Due to the covalent tethering of βE and βF by the 7–8 consensus disulfide bond in LU domains, we suspected that it could stabilize the β-sheet between strands E and F in uPAR DI and this could in part be reconciled with the reduced flexibility observed in the Kratky plot of the SAXS data (Fig. 5A). To probe this proposition further, we performed a continuous deuterium labeling of the different uPAR disulfide variants in the presence or absence of saturating levels of GFD and determined the deuterium uptake values by MS after pepsin digestion with special emphasis on peptide(57–66). The deuterium uptake plots for this peptide reveals a considerable variability in the flexibility of βE and βF in the different disulfide variants (Fig. 6A). Interestingly, the ranking of the deuterium uptake recapitulates to some extent the flexibility assigned

Figure 4. Comparison of molecular shape parameters of the uPAR disulfide variants by SAXS. A and B show the SAXS data and real-space distance-distribution functions of the uPAR disulfide variants. C and D show the SAXS data and real-space distribution functions of the uPAR variants in complex with ATF. Shown are experimental data (circles) along with fits of representative ab initio models, reconstructed using DAMMIF (dotted lines), the corresponding shape models are shown in Fig. S4. The lower panels in A and C show the error-weighted residual differences between the model fits and the experimental data.
by Kratky plots of the scattering data, *i.e.* $u_{\text{PART51C-V70C}}$ $\cong u_{\text{PARwt}} > u_{\text{PARK50C-V70C}} > u_{\text{PARH47C-N259C}}$. Yet one notable difference was apparent. The deuterium uptake for residues 57–66 differed significantly between the two $u_{\text{PAR}}$ variants with introduced 7–8 disulfide bonds, $u_{\text{PART51C-V70C}}$ exhibiting the far greater uptake thus resembling $u_{\text{PARwt}}$ (Fig. 6A). All isotope envelopes are unimodal at the shortest exchange time (10 s) in the presence and absence of GFD (Fig. S6), demonstrating that the amount of misfolded protein is negligible (48). This in turn signifies that the faster exchange kinetics in $u_{\text{PARTS1C-V70C}}$ as compared with $u_{\text{PARK50C-V70C}}$ reflects increased dynamics rather than irreversible protein misfolding (Fig. S6).

These differences in exchange rates of the 57–66 fragment among the tested $u_{\text{PAR}}$ disulfide conformers is almost erased in the corresponding $u_{\text{PAR}}$-GFD complexes (Fig. 6B). The compaction of $u_{\text{PAR}}$ by ligand binding thus reduces the macromolecular interdomain flexibility as well as the intradomain flexibility in DI, as monitored by SAXS and HDX-MS, respectively. Deuterium uptake plots for other regions in $u_{\text{PAR}}$ are shown in Fig. S5, but none of those show as prominent effects as peptide(57–66) (reporting on $\beta$IE and $\beta$IF).

**Binding kinetics of $u_{\text{PA}}$ to the $u_{\text{PAR}}$ disulfide variants**

To determine the kinetic rate constants for the $u_{\text{PA}}$-$u_{\text{PAR}}$ interactions by surface plasmon resonance, we developed a cap-
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![Image of deuterium uptake plots](image)

Figure 6. Domain flexibility of loop 3 in DI of the various uPAR disulfide variants tested by HDX-MS. A shows the deuterium uptake plots for the peptic peptide (57–66) from unoccupied uPAR at 25°C and pH 7.4. Shown are data from uPAR<sup>wt</sup> (blue), uPAR<sup>H47C-N259C</sup> (red), uPAR<sup>K50C-V70C</sup> (green), and uPAR<sup>T51C-V70C</sup> (orange). For comparison the data for uPAR<sup>wt</sup>-GFD complexes are shown (light gray). The peptide display the large differences observed between the different uPAR variants. B shows the corresponding deuterium uptake plots for the GFD-bound uPAR. In this graph, the uptake of unoccupied uPAR<sup>wt</sup> is shown (light gray). The hatched black line represents the peptide derived from the fully deuterated protein. C highlights the position of the peptic peptide (57–66) in intact uPAR (blue) where it represents the β-hairpin formed by β-strands βE and βF (using PDB code 3BT1). Uptake values are shown with standard deviations. Deuterium uptake plots for other peptic peptides are shown in Fig. S5.

**SMB binding to the various disulfide variants of uPAR**

To assess the low-affinity binding between the SMB domain of vitronectin and uPAR by surface plasmon resonance (SPR) in solution, we used a slightly different format. In this setup, 2-fold serial dilutions of SMB reached equilibrium binding with uPAR or uPAR-ATF complexes in solution before they were captured on the sensor surface by the anti-uPAR mAb R24 (Fig. 8A). The virtue of this system is that it uses relatively high concentrations of uPAR and ATF (100 and 150 nM), which makes certain that ATF saturates uPAR despite some of the analyzed disulfide mutants having impaired uPA binding, e.g. uPAR<sup>T51C-V70C</sup>. In accordance with previous studies (5–7), uPAR<sup>wt</sup> bound SMB with a K<sub>D</sub> of 3 μM, whereas no binding of SMB to uPAR<sup>R91D</sup> could be detected (Fig. 7B).

As shown in Table 2 and Fig. 7, the introduction of the 7–8 consensus disulfide bond in uPAR DI led to a minor decline in the affinity of SMB for intact uPAR. With one exception, all uPAR mutants exhibited an increased affinity for SMB upon saturation with ATF. As reported previously (6), the affinity of uPAR<sup>H47C-N259C</sup> for SMB did not change upon ATF binding as it already presents the higher affinity without bound ATF (Fig. 7C). The present SPR platform is thus capable of detecting the allosteric effect of ATF saturation on the SMB binding to uPAR in solution, albeit the magnitude of this effect is slightly less pronounced compared with the previous solid-phase based detection systems (6, 7). Nonetheless, this analysis clearly showed that the allosteric impact of ATF on SMB binding remain intact in both uPAR<sup>T51C-V70C</sup> and uPAR<sup>K50C-V70C</sup> showing that reintroduction of the 7–8 disulfide bond in uPAR DI does not uncouple the beneficial effect of ATF on SMB binding. In fact, binding of ATF managed to compensate for the lower inherent affinity of uPAR<sup>T51C-V70C</sup> and uPAR<sup>K50C-V70C</sup> for SMB and restored an affinity comparable with that of uPAR<sup>wt</sup>-ATF (Fig. 7, B versus D and E).

**Impact of deleting the 2–3 LU disulfide bond in uPAR DI**

Finally, we explored the impact of deleting the 2–3 disulfide bond in uPAR DI, the event that presumably led to neofunctionalization of this LU domain in snake venom α-neurotoxins (41). In all our functional tests, the uPAR<sup>C6S-C12S</sup> mutant behaved essentially as uPAR<sup>wt</sup> despite having only three of the five plesiotypic LU domain disulfides in DI. We found that (i) DI associate with DIIDIII after chymotrypsin cleavage of the linker region in uPAR<sup>C6S-C12S</sup> (Fig. 3F); (ii) the affinity of uPAR<sup>C6S-C12S</sup> for uPA and SMB is comparable with those of uPAR<sup>wt</sup> (Figs. 7F and 8F); and (iii) ATF-binding stimulates SMB binding to the same extent as uPAR<sup>wt</sup> (Fig. 8F). These data indicate that deletion of the 2–3 disulfide bond in...
uPAR DI has no deleterious effect on the known ligand interactions with uPAR.

Discussion

Loss and gain of disulfide-bonded cysteine residues (half-cystines) occurs very infrequently during evolution. One possible reason for this low frequency is that missense mutations of consensus half-cystines rarely survive selection and become permanently integrated in the genomes due to the deleterious effects of the reactive free thiol group in the partner half-cystine (49). Interestingly, one study found that in 99% of the cases, where missense mutations of half-cystines were maintained during evolution, both pairs of half-cystines were in fact replaced in concert (50). Circumstantial evidence from human genetics on LU domains align well with this proposition. One example illustrating this relationship is the pathological outcome of natural missense mutations in GPIHBP1. This protein has a single LU domain and it plays an essential role in intravascular triglyceride hydrolysis by shuttling the lipoprotein lipase to the capillary lumen (51). Individuals with a dysfunctional GPIHBP1 develop severe chylomicronemia. Notably, the majority of the missense mutations in human GPIHBP1 causing disease involves the replacement of single half-cystines (43) or introduction of an unpaired single cysteine (52). A similar pattern emerges for disease-causing mutations in the secreted single LU-domain protein SLURP1, where the dysfunctional protein is associated with a human skin disorder called mal de Meleda (53).

Notwithstanding the need for a concerted replacement of both half-cystines to eliminate a given disulfide bond, such rare events have in fact occurred in the evolution and diver-

Figure 7. Kinetics of uPAR-ATF interactions as assessed by surface plasmon resonance. A shows the principle in our capture protocol for assessing the rate constants of an oriented uPAR-ATF interaction by three rounds of single cycle kinetics. Initially, amine-coupled rabbit anti-mouse IgG (RAM) captures the high-affinity anti-uPAR mAb ATN-615 (31), which subsequently captures 50 nM uPAR yielding a binding stoichiometry of approximately two uPAR molecules bound per ATN-615. Finally, injections of five serial 2-fold dilutions of ATF without intermitting regeneration yields the binding curves. The inset shows a cartoon representation of the experimental setup. The double blank referenced sensorgrams are shown for uPAR野生 (B), uPARH47C-N259C (C), uPARK50C-V70C (D), uPART51C-V70C (E), and uPARC6S-C12S (F). Different colors of the sensorgrams represents different ATF concentrations used for the single cycle setup: 0.03–0.5 nM (blue), 0.06–1.0 nM (red), 0.12–2.0 nM (green), and 0.25–4.0 nM (purple). The thin black line superimposed on each curve represents the experimental fit to a simple bimolecular interaction and the corresponding residuals are in the bottom of the panels.
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Table 2
Kinetics of uPAR-ATF interactions and \( K_D \) of SMB binding

Analyses with single cycle protocols provided association (\( k_{on} \)) and dissociation (\( k_{off} \)) rate constants for the interactions between ATF in solution and different uPAR mutants captured on mAb ATN-615. This setup uses 3–4 rounds of single cycle injections each including five serial 2-fold dilutions of ATF1–135, which in combination covers the concentration range of 0.03–2 nM. Fitting with non-linear regression to a simple bimolecular interaction model yielded the kinetic rate constants and the stoichiometry was calculated as the molar ratio between captured ligand and calculated \( R_{max} \) for the analyte. Fig. 7 shows the corresponding sensorgrams. Equilibrium binding of SMB to uPAR and uPAR-ATF complexes were measured by SPR after capture to the mAb R24 as described in Fig. 8. Standard deviations (shown as ±) refer to parameters derived directly from the fitting procedures.

| uPAR         | \( k_{on} \) \( \times 10^6 \) M\(^{-1}\) s\(^{-1} \) | \( k_{off} \) \( \times 10^{-4} \) s\(^{-1} \) | \( K_D \) \( \mu M \) | Stoichiometry | \( K_D \) \( \mu M \) |
|--------------|-----------------------------------------------|-----------------------------------------------|----------------|---------------|----------------|
| Wt           | 11 ± 0.009                                    | 2.0 ± 0.002                                   | 0.019          | 0.94          | 3.3 ± 0.1     |
| T51C–V70C    | 10 ± 0.021                                    | 8.4 ± 0.002                                   | 0.019          | 0.88          | 4.8 ± 0.2     |
| K50C–V70C    | 15 ± 0.011                                    | 7.2 ± 0.005                                   | 0.050          | 0.67          | 5.9 ± 0.6     |
| H47C–N259C   | 12 ± 0.008                                    | 1.8 ± 0.001                                   | 0.015          | 0.94          | 2.0 ± 0.1     |
| C65–C12S     | 8.4 ± 0.015                                   | 2.5 ± 0.004                                   | 0.030          | 0.75          | 3.6 ± 0.1     |
| H47A         | 7.6 ± 0.053                                   | 2.0 ± 0.001                                   | 0.027          | 0.72          | 3.3 ± 0.1     |
| K50A         | 13 ± 0.011                                    | 4.1 ± 0.003                                   | 0.031          | 0.67          | 3.5 ± 0.1     |
| T51A         | 7.7 ± 0.005                                   | 6.0 ± 0.004                                   | 0.078          | 0.79          | 3.9 ± 0.1     |
| V70A         | 4.3 ± 0.002                                   | 2.2 ± 0.001                                   | 0.050          | 0.56          | 3.9 ± 0.1     |
| R91D         | 18 ± 0.002                                    | 2.2 ± 0.002                                   | 0.013          | 0.88          | 3.9 ± 0.1     |
| D140A        | 11 ± 0.009                                    | 17 ± 0.015                                    | 0.170          | 0.67          | 2.8 ± 0.1     |
| N259A        | 8.1 ± 0.006                                   | 1.3 ± 0.001                                   | 0.016          | 0.79          | 2.9 ± 0.1     |

Figure 8. Equilibrium binding of SMB to uPAR and uPAR-ATF complexes by surface plasmon resonance. A shows the principle in our capture protocol for assessing the equilibrium dissociation constant (\( K_D \)) between uPAR or uPAR-ATF complexes and SMB in solution. In brief, the amine-coupled anti-uPAR mAb R24 captures 100 nM uPAR or 100 nM uPAR-ATF complexes in the presence of 2-fold serial dilutions of SMB (0.3–10 \( \mu M \); replicates at 1.25 and 2.5 \( \mu M \)) in a multicycle setup. The inset shows the section of the sensorgrams where the dose-dependent binding of SMB is recorded. The SMB-binding isotherms for uPAR and uPAR-ATF complexes are shown for uPAR\(^{WT} \) and uPAR\(^{R91D} \) (B), uPAR\(^{H47C–N259C} \) (C), uPAR\(^{K50C–V70C} \) (D), uPAR\(^{T51C–V70C} \) (E), and uPAR\(^{C65–C12S} \) (F). Solid lines represent SMB-binding isotherms for uPAR and dotted lines those for the corresponding uPAR-ATF complexes.
Consensus disulfide bonds in LU domains

The evolution of snake venom α-neurotoxins clearly emphasizes this relationship, where the deletion of the 2–3 disulfide bond allegedly coincided with neofunctionalization and development of high potency toward synapsid neuronal acetylcholine receptors (41). One possible mechanism underlying this association is that the deletion of this particular disulfide bond would relax the LU domain scaffold, which allowed subsequent exploitation of new binding interfaces.

In the present study, we found that elimination of the 2–3 disulfide bond in the first LU domain of uPAR did not have notable functional consequences. Another report showed that elimination of this disulfide bond in the single LU domain protein CD59 also had no impact on its complement regulatory function (44). Based on these studies, it would thus appear that the plesiotypic 2–3 disulfide bond is not essential for the overall structural integrity of LU domain proteins, which would explain why its deletion could cause functional diversification in α-neurotoxins.

Paradoxically, the one plesiotypic LU domain disulfide bond that actually is lost in uPAR is the 7–8 disulfide in DI. In contrast to the 2–3 disulfide, this disulfide bond appears to be indispensable for the correct folding and function of single LU domain proteins in general (45, 51, 53, 54). Furthermore, the deletion of this particular disulfide bond occurs in very few proteins and exclusively in the first LU domain of proteins with 2 or more LU domains e.g. uPAR/PLAUR (31, 35, 40), CA4A/LYPD3 (55, 56), and Haldisin/LYPD5 (57). Bearing in mind that flexibility between the individual LU domains in uPAR plays an important role for ligand binding (5, 6, 38) it becomes highly pertinent to ask the question: did deletion of this plesiotypic 7–8 disulfide in uPAR DI facilitate the evolution of a high-affinity ligand-binding cavity for uPA by enabling a dynamic assembly of its three LU domains?

To address this question, we first needed to identify the putative positions of the two deleted half-cystines. Our comprehensive alignments of more than 50 annotated orthologous uPAR sequences from the class of Mammalia revealed an invariant distance (5 amino acids) between the 6th and 7th cysteines in the second and third LU domains of all orthologues (not shown, but Fig. 1A, and Figs. S1 and S8 provide representative alignments). Moreover, the 5th and 7th positions after the 6th consensus cysteine were invariably a lysine and a glycosylated asparagine in uPAR DI (equivalent to Lys50 and Asn52 in human uPAR DI). In contrast, the 6th position, the assumed position of the deletion, of the 7–8 disulfide bonds were preserved. Importantly, uPAR-like orthologues annotated from the class of Reptilia including lizards, snakes, turtles, and crocodilians (Fig. S8) faithfully replicate these properties. It is therefore beyond any reasonable doubt that the correct position, from an evolutionary point view, for the introduction of the missing 7–8 disulfide bond in human uPAR DI is indeed Thr51 and Val70.

The corresponding uPAR<sub>T51C-V70C</sub> mutant expressed well in S2-cells and all cysteines were engaged in the expected disulfide bonding in the purified protein. Nonetheless, our biochemical and biophysical analysis on the purified uPAR<sub>T51C-V70C</sub> revealed that the disulfide constrained DI behaves very different to that of uPAR<sup>wt</sup>. Among other features, we observed with size exclusion chromatography that the introduction of the 7–8 disulfide abrogated the interdomain interaction between DI and DII in uPAR<sub>T51C-V70C</sub> (Fig. 3). Importantly, we show that the penalty for introducing the missing disulfide in uPAR DI is a 40-fold reduction in its affinity for uPA, which would provide uPAR<sub>T51C-V70C</sub> with a <em>K<sub>D</sub></em> that is at least 40-fold above the plasma concentration of pro-uPA in humans (58). It would therefore appear that the evolutionary deletion of the plesiotypic 7–8 disulfide bond in uPAR DI was essential for creating a high-affinity binding site for uPA via a flexible assembly of all three LU domains in uPAR. This proposition is well aligned with studies on the co-evolution of uPA and uPAR (32, 59). Based on crystal structures of human and murine uPAR-ATF complexes and extensive mutagenesis (3, 31, 32, 60), the functional hotspot residues in the β-hairpin of GFD for uPAR binding is well characterized in these species. The molecular basis for the species selectivity in the uPAR-uPA interaction between primates and nonprimate mammals is represented by the concerted replacement of the Asn22<sup>uPAR</sup> → Tyr23<sup>uPA</sup> and Trp30<sup>uPAR</sup> → Arg31<sup>uPA</sup> dyad in human and mouse uPA (32) (Fig. S9). Moreover, the majority of the important residues in the uPAR-binding motif of GFD from mammals are also conserved within the class of Reptilia (Fig. S9A) and this class is also the earliest class where a bona fide uPAR orthologue with three LU domains can be traced (59). In this phylogenetic class, uPAR DI has already lost its plesiotypic 7–8 disulfide (Fig. S8). In the class of Sarcopterygii, a uPAR-like protein with three LU domains has recently been identified in the African lungfish <em>Protopterus</em> (59). This protein is unique in the sense that it retains the full signature of an ancient LU domain with all five plesiotypic disulfide bonds present in all three domains (Fig. S8). Examination of the β-hairpin of GFD in the corresponding uPA orthologue reveals that the binding motif known from <em>Mammalia</em> and <em>Reptilia</em> has not yet evolved (Fig. S9A). It is therefore possible that this interaction is not operational in species belonging to lobe-finned fishes, but further functional studies on purified proteins is required before a definitive conclusion can be made.

Evolution of the low-affinity interaction between uPAR and the SMB domain of vitronectin follows a slightly different path. Although the hotspot residues in SMB for the vitronectin interaction with human uPAR is highly conserved all the way to <em>Sarcopterygii</em> (Fig. S10A), the hotspot residue in uPAR (i.e. Arg91) in the linker region between DI and DII is only conserved within <em>Mammalia</em> (Fig. S10B). It is therefore likely that the co-evolution, which shaped the uPAR-binding site in SMB, originally was driven by the interaction between vitronectin and the cognate uPA-inhibitor plasminogen activator inhibitor type 1, in which the hotspot residue for SMB-binding (i.e. Arg101) is conserved in lobe-finned fishes, bony fishes, and cartilaginous fishes (61). Later in evolution, the class of <em>Mammalia</em> presumably developed the uPAR-vitronectin interaction through convergent evolution.
Consensus disulfide bonds in LU domains

Experimental procedures

Chemicals and purified proteins

Recombinant human uPAR (residues 1–283) and a panel of mutants thereof were expressed and secreted by Drosophila melanogaster S2 cells and purified from the culture supernatants by affinity chromatography (62). The SMB domain of human vitronectin (residues 1–48) with a C-terminal His6 purification tag was expressed by Pichia pastoris and purified and characterized as described (63). Human pro-uPA1–411 (as the catalytic inactive S356A mutant) and its receptor binding N-terminal fragment ATF1–143 were produced by D. melanogaster S2 cells and purified by affinity chromatography (60). The growth factor-like domain of human uPA, GDF (residues 1–48), was a kind gift from Dr. S. Rosenberg. Mouse monoclonal anti-human uPAR antibodies were produced either locally (R24) (5) or was a kind gift from Dr. A. Mazar (ATN-615) (31). Recombinant peptide-Nε-(acetyl-β-glucosaminyl)-asparagine amidase (PNGase F; E.C. 3.5.1.52) was from Roche Applied Science (25 units/μg) and α-chymotrypsin was from Worthington.

Generation of uPAR1–87 by limited proteolysis with chymotrypsin

The N-terminal LU domain of human uPAR (DI, residues 1–87) was excised by limited proteolysis with chymotrypsin using conditions that preferentially led to hydrolysis of the Tyr87–Ser88 peptide bond in uPAR. In brief, 45 μg of intact uPAR was incubated with 9 ng of chymotrypsin (E:S of 1:5000) in 54 μl of PBS, pH 7.4, for 120 min at 25 °C and the released uPAR DI1–87 was isolated by size exclusion chromatography with a SuperdexTM 75 HR10/300 columnTM (GE Healthcare) operated at 0.5 ml/min in PBS (40). Time course experiments were conducted at higher chymotrypsin ratios (E:S of 1:750) for up to 24 h, which led to an additional cleavage at Tyr57–Arg58 in uPARwt.

Deglycosylation of intact uPAR under native conditions

Purified uPAR variants (10 μg in 10 μl of PBS) were incubated for 1 h at 25 °C with 1 unit of PNGase F. To discriminate between effects on DI and DIIDIII, the PNGase F-treated samples were further incubated for 1 h with 2 ng of chymotrypsin (1:5000 w/w) to cleave the Tyr87–Ser88 peptide bond in the linker region. Heating at 95 °C in SDS-PAGE sample buffer containing 20 mM DTT stopped the reaction. After cooling and addition of 50 mM iodoacetamide, samples were analyzed by SDS-PAGE and visualized by Coomassie G250 staining.

Surface plasmon resonance

We determined the binding kinetics for the uPA-uPAR interactions with SPR measurements on a Biacore T200TM system (GE Healthcare). To accomplish this, we immobilized a polyclonal rabbit anti-mouse immunoglobulin antibody (GE Healthcare) as the first capture layer (30 μg/ml in 10 mM sodium acetate, pH 5.0) on a CM5 sensor chip using N-hydroxysuccinimide and N-ethyl-N-(3-(diethylamino)propyl)-carbodiimide. This yielded a surface density of 1100 resonance units (RU), which corresponds to 7.3 fmol of mAb/mm2 (assuming one RU ~ 1 pg/mm2). Injection of 1 μmol ethanolamine inactivated excess NHS-esters. To prepare the second capture layer, we injected 50 nM of a high-affinity mouse monoclonal anti-uPAR mAb (ATN-615) for 350 s at 20 μl/min in the active flow cell only, which led to ~0.8 fmol of mAb/mm2 (120 RU). The last capture step involved a 300-s injection of 50 nm uPAR at 20 μl/min in both flow cells, which led to a capture level of 1.6–2.0 uPAR molecules bound per ATN-615 in the active flow cell. Kinetic rate constants for the various uPA-uPAR interactions were determined with single cycle protocols by which five serial 2-fold dilutions of the interaction partner (ATF) were injected for 200 s without intervening regeneration and followed by a longer dissociation phase (1,000–3,000 s dependent on the dissociation rate constant koff). These real-time interactions were measured at 50 μl/min in 10 mM HEPES, 150 mM NaCl, 3 mM EDTA, and 0.05% (v/v) surfactant P-20 at pH 7.4 at 20 °C. Three consecutive injections of 10 mM glycine/HCl, pH 1.7, at the end of each single cycle regenerated the chip.

Fitting of the double blank referenced data by nonlinear regression to a bimolecular interaction model, assuming pseudo-first order reaction conditions, yielded the association (koni) and dissociation (koff) rate constants, the Kd (koff/koni), as well as the binding capacity (Rmax). We used the evaluation software supplied with the instrument for global fitting (BiacoreT200 EvaluationTM 3.0).

Our experimental protocol for measuring equilibrium binding between SMB and uPAR or uPAR-ATF complexes relies entirely on steady state conditions in solution between 100 nm uPAR ± 150 nm ATF and serial 2-fold dilutions of SMB (0.3–10 μM). Binding of SMB was monitored by the mass increase of uPAR or uPAR-ATF complexes during capture by the anti-uPAR mAb R24 as a function of the added SMB concentration (37). We used amine chemistry to immobilize R24 directly on a CM5 chip at a density of 651 RU (4.3 fmol of mAb/mm2). We designed this atypical binding protocol to minimize the effects of the vastly different inherent stabilities of the bimolecular uPAR-ATF and uPAR51C-V70C-ATF complexes (see Table 2). Using concentrations of uPAR and ATF that were at least 100-fold above the highest measured Kd, we made sure that uPAR was saturated with ATF during the steady state binding to SMB in solution.

Hydrogen-deuterium exchange

Purified uPAR was diluted in PBS to a final concentration of 30 μM in the absence or presence of 2-fold molar excess of GFD1–48, and subsequent incubation for 15 min at 25 °C secured complete complex formation. To initiate isotopic exchange, samples were diluted 10-fold in PBS exchange D2O buffer (10 mM Na2HPO4, 150 mM NaCl in D2O, pHread 7.2), which resulted in 90% D2O (v/v) in the final labeling solutions. Aliquots of 60 pmol of uPAR were withdrawn after 10, 100, and 1000 s of incubation at 25 °C. We used acidification and lowered the temperature to efficiently prevent further exchange by adding 1 volume of ice-cold quench buffer (0.1 M Na2HPO4, 0.8 M tris-(2-carboxyethyl)phosphine, 2 M urea in H2O, pH 2.5) to the withdrawn sample and kept it on ice for 3 min to allow reduction of disulfide bonds. Subsequently, the quenched samples were snap frozen in liquid nitrogen and stored at ~80 °C until analysis. All samples were labeled in triplicates except
uPAR^{T51C-V70C}, which was measured in duplicates. To prepare an undeuterated control, the samples were prepared as described above except we replaced the PBS exchange D_{2}O buffer with the corresponding protiated solvent. We prepared a full-deuterium control by incubating the sample in the exchange buffer for 72 h at 37 °C before quenching.

**Determining deuterium uptake by UPLC-ESI-MS**

Quenched and reduced uPAR samples (60 pmol) were thawed and immediately injected into a cooled (0 °C) nanoACQIUTY UPLC reversed-phase chromatographic system equipped with HDX technology (Waters, Milford, MA), using an ice-cold syringe to minimize back-exchange. The sample entered a 100-µl injection loop prior to the on-line digestion of the protein, which was carried out by a 2.0 × 20-mm column (IDEX Upchurch Analytical Guard Column, Oak Harbor, WA) packed with agarose-immobilized pepsin (Thermo Scientific Pierce, Rockford, IL), located in a compartment with a temperature of 20 °C. The on-line digestion occurred at a flow rate of 300 µl/min in 0.23% (v/v) formic acid in H_{2}O and the generated peptic peptides were trapped on a short guard column and desalted for 3 min (ACQUITY UPLC BEH C18 VanGuard Pre-Column, 1.7 µm, 2.1 × 5 mm, Waters). Subsequently, the peptides were separated at a flow rate of 40 µl/min on a 1.0 × 50-mm analytical column (ACQUITY UPLC BEH C18, 1.7 µm, 1.0 × 50 mm, Waters) with a 12-min linear gradient from 5–50% (v/v) acetonitrile containing 0.23% (v/v) formic acid. In some experiments, we used a longer analytical column (100 mm).

Eluted peptides were introduced into a quadrupole TOF mass spectrometer (Synapt G2, Waters Corp.) by electrospray ionization. Acquired MS spectra was lock-mass corrected against leucine enkephalin and calibrated against sodium iodide.

Extensive cleaning reduced carryover between runs to less than 5%. In brief, we washed the injection loop with 200 µl of 50% (v/v) MeOH in H_{2}O containing 0.23% (v/v) formic acid, followed by 200 µl of 0.23% (v/v) formic acid followed by a blank gradient.

The peptic peptides from uPAR were identified by collision-induced dissociation with data independent (MS^{3}) acquisition mode. Protein Lynx Global Server 3.0 (PLGS) software (Waters) searched and identified the peptic peptides from uPAR. Data processing with DynamX 3.0 (Waters) determined the deuterium content in each peptide.

**Small-angle X-ray scattering**

Synchrotron radiation X-ray scattering data were collected on the EMBL P12 beamline of the storage ring PETRA III (DESY, Hamburg) and EMBL X33 beamline of the storage ring DORIS (DESY, Hamburg) (Tables S1–S4), using PILATUS 2M and 1M pixel detectors (DECTRIS, Switzerland), respectively. Batch experiments of uPAR and uPAR-ATF complexes (purified by size exclusion chromatography) were measured in 20 mM PBS, 5% glycerol, pH 7.4, ± 50 mM NaSO_{4}, while flowing through a temperature controlled capillary (P12: 1.2-mm inner diameter, X33: 1.7-mm inner diameter) at 10 °C. Twenty frames of 0.05-s exposure time (P12) or four frames of 30-s exposure time (X33) were collected. The sample-to-detector distance was 2.7 m (P12) and 3.1 m (X33), covering a range of momentum transfer 0.002 Å^{-1} ≤ s ≤ 0.5 Å^{-1} and 0.008 Å^{-1} ≤ s ≤ 0.6 Å^{-1} (s = 4πsinθ/λ, where 2θ is the scattering angle, and λ = 1.24 Å is the X-ray wavelength), respectively. Based on comparison of successive frames, no detectable radiation damage was observed. Data from the detector were normalized to the transmitted beam intensity, averaged, placed on absolute scale relative to water, and the scattering of buffer solutions subtracted. All data manipulations were performed using PRIMUSqt and the ATSAS software package (64). The forward scattering I(0) and radius of gyration, R_{g} were determined from Guinier analysis (65) assuming that at very small angles (s ≤ 1.3/R_{g}) the intensity is represented as I(s) = I(0)exp(−(sR_{g})^{2}/3). These parameters were also estimated from the full scattering curves using the indirect Fourier transform method implemented in the program GNOM (66), along with the distance distribution function p(r) and the maximum particle dimensions D_{max}.

Molecular masses of solutes were estimated from I(0) by computation of partial specific volume and the contrast between the glycosylated protein sequence and the chemical components of the solution using the MULCH server (http://smb-research.smb.usyd.edu.au/NCVWeb/)^{3} (67). Computation of theoretical scattering intensities was performed using the program CRYSOL (68).

**Ab initio shape determination**

Low resolution shapes were reconstructed from SAXS data using the programs DAMMIF (69), which represents the macromolecule as a densely packed interconnected configuration of beads or chain-like ensemble of dummy residues, respectively, that best fits the experimental data I_{exp}(s) by minimizing the discrepancy,

$$\chi^{2} = \frac{1}{N-1} \sum_{s} \frac{[I_{exp}(s) - c \cdot I_{calc}(s)]^2}{\sigma(s)}$$

(Eq. 1)

where N is the number of experimental points, c is a scaling factor, and I_{calc}(s) and \sigma(s) are the calculated intensity and the experimental error at the momentum transfer s, respectively. Multiple modeling runs were conducted to verify the stability of the solution, and to establish the most typical 3D reconstructions according to a spatial discrepancy measure using DAMAVER (70).

**Ensemble modeling**

Crystal structures of uPAR and uPAR-ATF complex (PDB IDs 2FD6, 3U74, and 3U73) were used as template for flexible modeling of the uPAR domain structures in solution. Glycosylation was introduced into the models using the GLYCOSYLATION routine of ATSAS (64). The program EOM (47) was used to generate a pool of 10,000 uPAR conformations with flexible linkers defined between the three LU domains. A generic algorithm was then employed to select subsets of conformers from the pool that best fit the experimental scattering data for each

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Consensus disulfide bonds in LU domains

construct. The flexibility metric, $R_{flex}$, was determined for each ensemble as described previously (47), where $R_{flex} = 100\%$ indi-
cates maximum flexibility/uncertainty. SANS data has been deposited at the SASBDB (www.sasdbd.org)\textsuperscript{3} with accession codes: SASDAT4, SASDAX4, SASDAU4, SASDF82, SASDF92, SAS DFA2, SASDFB2, and SASDFC2.

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Did evolution create a flexible ligand-binding cavity in the urokinase receptor through deletion of a plesiotypic disulfide bond?

Julie M. Leth, Haydyn D. T. Mertens, Katrine Zinck Leth-Espensen, Thomas J. D. Jørgensen and Michael Ploug

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