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Ana Cristina Sotomayor-Pérez, Daniel Ladant, Alexandre Chenal. Calcium-induced folding of intrinsically disordered repeat-in-toxin (RTX) motifs via changes of protein charges and oligomerization states.. *Journal of Biological Chemistry*, 2011, 286 (19), pp.16997-17004. 10.1074/jbc.M110.210393 . pasteur-01508724

HAL Id: pasteur-01508724

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Submitted on 14 Apr 2017

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Calcium-induced Folding of Intrinsically Disordered Repeat-in-Toxin (RTX) Motifs via Changes of Protein Charges and Oligomerization States^{*[5]}

Received for publication, December 8, 2010, and in revised form, February 26, 2011. Published, JBC Papers in Press, March 15, 2011, DOI 10.1074/jbc.M110.210393

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Ligand-induced disorder-to-order transition plays a key role in the biological functions of many proteins that contain intrinsically disordered regions. This trait is exhibited by so-called RTX (repeat-in-toxin) motifs found in many virulence factors secreted by numerous Gram-negative pathogenic bacteria: RTX proteins are natively disordered in the absence of calcium but fold upon calcium binding. The adenylate cyclase toxin (CyaA) produced by *Bordetella pertussis*, the causative agent of whooping cough, contains ~40 RTX motifs organized in five successive blocks separated by non-RTX flanking regions. This RTX domain mediates toxin binding to its eukaryotic cell receptor. We previously showed that the last block of the RTX domain, block V, which is critical for CyaA toxicity, exhibits the hallmarks of intrinsically disordered proteins in the absence of calcium. Moreover, the C-terminal flanking region of CyaA block V is required for its calcium-induced folding. Here, we describe a comprehensive analysis of the hydrodynamic and electrophoretic properties of several block V RTX polypeptides that differ in the presence and/or length of the flanking regions. Our results indicate that the length of the C-terminal flanking region not only controls the calcium-induced folding but also the calcium-induced multimerization of the RTX polypeptides. Moreover, we showed that calcium binding is accompanied by a strong reduction of the net charge of the RTX polypeptides. These data indicate that the disorder-to-order transition in RTX proteins is controlled by a calcium-induced change of the polypeptide charges and stabilized by multimerization.

The past decade has seen a fundamental reappraisal of the protein structure-to-function paradigm because it became evident that a significant fraction of polypeptide sequences code

for proteins, commonly designated as natively or intrinsically disordered proteins, that are lacking ordered structures under physiological conditions (1–8). Numerous intrinsically disordered proteins are nevertheless able to acquire ordered conformations upon binding to specific ligands, and in most cases, these disorder-to-order transitions are linked to key functional properties of these proteins. Elucidation of the biochemical factors and physicochemical parameters that control the transition between these different states of the polypeptides is therefore essential to understand the biological functions of intrinsically disordered proteins.

We have recently begun to explore the biophysical mechanisms at work in this process in a newly described class of intrinsically disordered protein that contains so-called RTX repeated motifs and in which the disorder-to-order transition is triggered by the binding of calcium ions (9–11). These motifs are found in many virulence factors secreted by numerous Gram-negative pathogenic bacteria (12, 13). The original name (repeat in toxin) was first coined to group four virulence factors secreted by *Escherichia coli*, *Bordetella pertussis*, *Pasteurella hemolytica*, and *Actinobacillus* sp., that contain a series of tandemly repeated RTX sequences at their C-terminal end. Since then, many other RTX proteins have been identified in more than 250 bacterial strains (14), and they cover a vast array of biological activities, many of them involved in host-pathogen interactions (13, 14). In most cases, the RTX sequences are located at the C-terminal end of the polypeptides, appended to functional domains endowed with specific biochemical activities such as pore-forming cytolytic activity, proteases, lipases, etc. (13, 14). Despite this functional diversity, most known RTX proteins are thought to be secreted by a dedicated type I secretion system and to require calcium to exert their biological activities.

The prototype sequence of the RTX motif is GGXG(N/D)DX(U)X, where X represents any amino acid, and (U) represents any large hydrophobic residue such as Ile, Leu, Val, Phe, and Tyr (13). The number of RTX motifs in RTX proteins may vary from 5 to more than 50. Structural data available on some RTX-containing proteins showed that these sequences fold in the presence of calcium into an idiosyncratic parallel β -helix structure, where the first six residues, GGXGXD, of each RTX motif form a calcium-binding turn and the last three residues XUX make a short β -strand (15, 16). The assembly of consecutive turns and β -strands builds up a right-handed parallel β -helix that is stabilized by calcium binding. In contrast, in the

^{*} This work was supported by CNRS Institut Pasteur Projet Transversal de Recherche Grant PTR#374 and the Agence Nationale de la Recherche Programme Jeunes Chercheurs Grant ANR-09-JJC-0012.

^[5] The on-line version of this article (available at <http://www.jbc.org>) contains supplemental text, Table S1, and Fig. S1.

¹ Supported by the Allocation de Thèse de la Région Ile-de-France Ph.D. program.

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absence of calcium, these RTX sequences appeared to be largely disordered as revealed in particular by our biophysical studies of the adenylate cyclase (CyaA) toxin from *B. pertussis*, the causative agent of the whooping cough (17, 18). CyaA contains ~40 RTX motifs grouped in five consecutive blocks (named blocks I–V) separated by non-RTX-flanking regions. This RTX-containing domain (RD)⁴ is located at the C-terminal moiety of CyaA (residues 1006–1706) and is implicated in the calcium-dependent binding of the toxin to a specific receptor, the CD11b/CD18 integrin, at the surface of the host target cells. We indeed showed that RD is intrinsically disordered in the absence of calcium but acquires a stable β -rich fold upon calcium binding (9).

It has been proposed that the calcium-induced disorder-to-order transition in RTX motifs may be essential for the secretion of the corresponding polypeptides by their dedicated type I secretion system. The disordered conformations of RTX motifs in the low calcium environment of the bacterial cytosol may facilitate the protein secretion through the type I secretion system machinery, whereas in the external medium, calcium binding could trigger the folding of the proteins into their active conformations for host-pathogen interactions (9, 19–21).

To gain further insights into the mechanisms of calcium-induced folding of the CyaA RTX domain, we further characterized a region of RD that encompasses the last block (block V) of RTX motifs and that was found to be essential for the cytotoxic activity of CyaA (22–24). We demonstrated that the RTX motifs *per se* were not able to fold in the presence of calcium and that the polypeptide segments at the C termini of the tandemly repeated RTX motifs were essential for the calcium-induced folding (10), yet the molecular basis allowing this C-terminal flanking region to induce the folding of the RTX motifs remained elusive.

Here, we have investigated the electrokinetic and hydrodynamic properties of several RTX polypeptides derived from block V of RD by a combination of complementary biophysical approaches. Our results indicate that the C terminus-flanking region is not only involved in the calcium-induced folding but can also modulate the multimerization of the RTX polypeptides.

Moreover, we showed that calcium binding induced a strong reduction of the mean net charges of the RTX polypeptides. This indicated that the disorder-to-order transition of RTX proteins could be controlled by calcium-induced changes in protein valence.

EXPERIMENTAL PROCEDURES

Reagents—The experiments were done at 25 °C in 20 mM Hepes, 20 mM NaCl, pH 7.4 (Buffer A). All reagents were of the highest purity grade.

Polypeptides Production and Purification—RTX polypeptide production and purification have been described elsewhere (10). Protein preparation is further described in the [supplemental materials](#).

⁴The abbreviations used are: RD, RTX-containing domain; QELS, quasi-elastic light scattering; SEC, size exclusion chromatography; TDA, Tetra detector array; AUC, analytical ultracentrifugation.

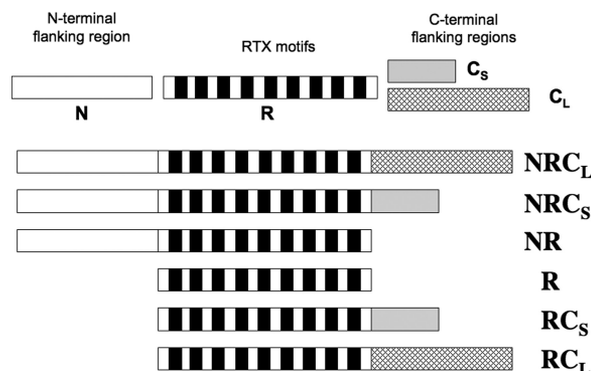


FIGURE 1. Scheme of the six block V polypeptides used in this study. Several regions can be distinguished along the primary sequence of CyaA block V (residues 1487–1680). From left to right on the scheme are: the N-terminal flanking region (N) for residues 1487–1529 (white); the RTX motif region (R; each of the nine RTX motifs is represented by a black box) for residues 1530–1630; the proximal C-terminal flanking region for residues 1631–1652 (the short C region, C_S, gray); and the long C-terminal flanking region (C_L, hatched) for residues 1631–1680.

Spectroscopic Methods—Quasi-elastic light scattering (QELS) experiments (11) and electrophoretic mobility measurements were done using a NanoZS instrument (Malvern). Fluorescence measurements were performed with a FP-6200 spectrofluorimeter (Jasco), and CD spectra were recorded on an Aviv model 215 circular dichroism spectropolarimeter, as described previously (10). The estimation of protein shape and hydration using size exclusion chromatography (SEC) connected on-line to a Tetra detector array (TDA) (Viscotek Ltd., a Malvern Company) in conjunction with analytical ultracentrifugation (Beckman XL-I; Beckman Coulter) is described elsewhere (9, 10, 25, 26). A further description of CD, anisotropy, QELS, and electrophoretic mobility experiments is provided in the [supplemental materials](#).

RESULTS

The polypeptides studied in this work have been described previously (10) and are schematized in Fig. 1. They all encompass the core of the last block (block V) of RTX motifs (residues 1530–1630, noted R) from CyaA but differ in the presence of either the N-terminal (residues 1487–1529, noted N) or the C-terminal (C) flanking region, the latter being further subdivided into short C_S (residues 1631–1652) and long C_L (residues 1631–1680) regions. These polypeptides were produced and purified as described previously (see “Experimental Procedures” and Ref. 10 for details).

Calcium-induced Changes of RTX Protein Hydrodynamic Radii—We first measured the hydrodynamic radii (R_H) of the six RTX polypeptides by QELS. In the apo-state (*i.e.* in the absence of calcium), all RTX polypeptides showed hydrodynamic radii higher than expected for folded proteins of similar molecular masses (compare anhydrous to hydrodynamic radii in Table 1), in agreement with our previous results, which revealed that all RTX polypeptides are intrinsically disordered in the apo-state (10). We previously showed, and confirmed here, that the polypeptides harboring the C-terminal flanking region C_L or C_S, *i.e.* NRC_L, RC_L, NRC_S, and RC_S, underwent a calcium-induced folding, whereas the NR and R polypeptides, lacking a C-terminal extension, did not (Ref. 10

TABLE 1

Hydrodynamic radii of RTX polypeptides measured by quasi-elastic light scattering

The experimental conditions were as follows: temperature, 25 °C; apo-states, buffer A; in the presence of calcium, buffer A, 2 mM CaCl₂ for holo-NRC_L, holo-RC_L, and holo-RD; buffer A, 5 mM CaCl₂ for NRC_S, RC_S, NR, and R. NA, not applicable (because of the heterogeneity of the samples). The S.D. of the frictional ratio is within 10% of the value.

	NRC _L	RC _L	NRC _S	RC _S	NR	R	RD
R_H (nm) apo in the absence of calcium	3.4 ± 0.1	3.2 ± 0.1	3.4 ± 0.1	2.9 ± 0.1	3.0	2.7 ± 0.1	7.2 ± 0.3
R_H (nm) holo in the presence of calcium	2.5 ± 0.1	2.2 ± 0.2	NA	3.1 ± 0.1	3.0	2.7 ± 0.1	3.1 ± 0.1
R_0 anhydrous (nm) Apo	1.81	1.68	1.72	1.58	1.64	1.47	2.83
R_0 anhydrous (nm) Holo	1.81	1.68	1.72	1.58	1.64	1.47	2.75
f/f_0 Apo	1.88	1.93	1.97	1.84	1.82	1.84	2.54
f/f_0 Holo	1.38	1.32	NA	1.97	1.82	1.84	1.13

and supplemental Fig. S1). Accordingly, the addition of calcium did not change the hydrodynamic radii of NR and R (Table 1), whereas it triggered a large decrease of the R_H of NRC_L and RC_L, indicative of a strong compaction of their hydrodynamic volumes upon acquisition of secondary and tertiary structures.

Strikingly, the R_H of holo-RC_S was not significantly different from that of the apo-form, although the CD spectra clearly indicated that RC_S acquired secondary and tertiary structures in the presence of calcium (Table 1 and supplemental Fig. S1). Unfortunately, we could not measure the R_H of holo-NRC_S as a significant fraction of the protein aggregated in these conditions, precluding an accurate analysis of the QELS data. In contrast, the QELS data of apo- and holo-RC_S were robust and good enough to accurately determine their translational diffusion coefficients that were also found (see below) to be in agreement with AUC data. It was therefore intriguing that the intrinsically disordered apo-RC_S state could fold in the presence of calcium (supplemental Fig. S1) without a concomitant decrease of its hydrodynamic radius (Table 1). To better describe the calcium-induced folding reaction of RC_S, we further investigated its hydrodynamic properties by SEC-TDA, by analytical ultracentrifugation (AUC) in velocity mode, by fluorescence anisotropy, and by electrophoretic mobility.

The SEC-TDA experiments were performed on a Superdex 200 column at 25 °C. The chromatographs of RC_S in the apo-state and in the holo-state (in the presence of 5 mM calcium) are shown in Fig. 2A. The retention volume of RC_S was similar in the absence and in the presence of calcium (Fig. 2A and supplemental Table S1), suggesting that both states exhibited similar hydrodynamic volumes, in agreement with the QELS data. The right angle light scattering signals (RALS, green profiles), combined with the protein concentration determined by the deflection refractometer (black profiles), provided on-line measurements of the molecular mass of each eluting species. As shown in Fig. 2B, the molecular mass determined for the apo-state of RC_S was 15.1 kDa, as expected for a monomeric species (Fig. 2B, green dashed trace), whereas the molecular mass of the holo-state (Fig. 2B, green solid trace) was 42 kDa. Hence, these data indicated that RC_S formed a trimer upon calcium binding (supplemental Table S1). To further estimate the equilibrium between the monomer and the trimer, holo-RC_S was loaded on the SEC-TDA at three different concentrations (70, 35, and 7 μM). Irrespective of the protein concentration, holo-RC_S eluted as a single peak exhibiting a molecular mass of 42 ± 2 kDa (data not shown). No other oligomeric states (*i.e.* monomer, dimer, tetramer, etc.) could be detected, suggesting that the trimeric state was the most stable form of holo-RC_S, if not the only

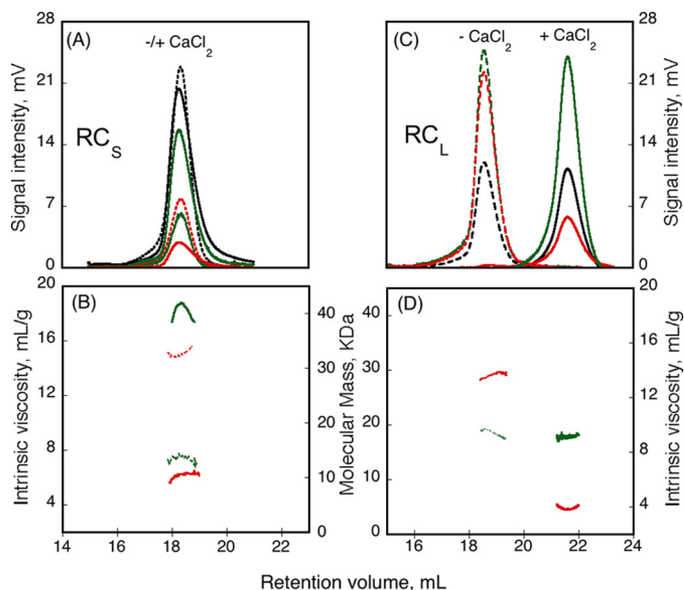


FIGURE 2. Hydrodynamic properties of RC_S and RC_L polypeptides analyzed by SEC-TDA. RC_S (A) and RC_L (C) polypeptides, in the absence (dashed line) or in the presence (solid line) of 5 mM CaCl₂, were analyzed by size exclusion chromatography on a Superdex 200 connected to a Tetra detector array as described under "Experimental Procedures." The traces correspond to the right angle light scattering (green lines), differential refractometer (black lines), and differential viscometer (red lines). The experimental conditions were: buffer A at 25 °C. The molecular mass (green lines) and intrinsic viscosity (red lines) are given for RC_S (B) and RC_L (D) in the absence (dashed lines) or in the presence of calcium (solid lines).

possible one. To rule out any possible artifact, we analyzed in parallel the RC_L protein by SEC-TDA, and as shown in Fig. 2 (C and D), we found that, in agreement with our previous results (10), RC_L is monomeric in both states, and its R_H decreased dramatically upon calcium binding (Table 1). Finally, the SEC-TDA experiments also provided the intrinsic viscosity (measured by on-line differential pressure transducers) of each eluting species. As shown in Fig. 2B, the intrinsic viscosity of RC_S strongly decreased from 14.7 to 6.4 ml·g⁻¹ upon calcium binding (Fig. 2B, red traces, and supplemental Table S1). This change of intrinsic viscosity indicates that RC_S underwent a change of hydration and/or shape upon calcium binding (see below).

Hydrodynamic Parameters of RTX Proteins—We then measured the sedimentation coefficients of the apo- and holo-states of RC_S by AUC (i) to independently confirm the results obtained by QELS and SEC-TDA and (ii) to further determine the respective contribution of the viscosity increment and hydration parameters in the intrinsic viscosity values of the different species. Velocity experiments provided sedimentation

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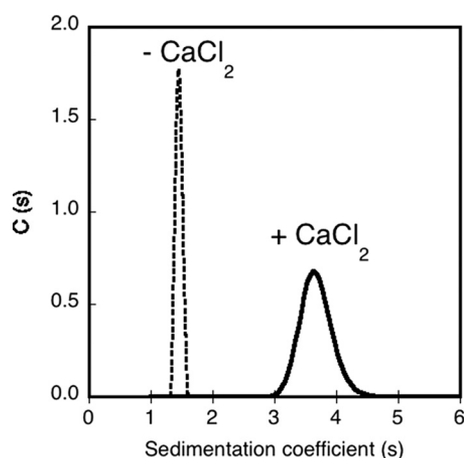


FIGURE 3. **Analytical ultracentrifugation analysis of RC_S.** Sedimentation coefficient distribution, deduced from the fitted curves, of RC_S in the absence (dashed line) or in the presence of 5 mM CaCl₂ (solid line). The experimental conditions were: 20 μM RC_S and buffer A at 25 °C.

coefficients of 1.46 and 3.65 S for apo-RC_S and holo-RC_S, respectively (Fig. 3 and supplemental Table S1). These data, together with those acquired by QELS and SEC-TDA, provided independent measurements of S , R_H , and molecular mass (supplemental Table S1) that are related to each other according to the Svedberg equation $S = M(1 - \rho\bar{v})/(6\pi\eta N_A R_H)$. All of the three experimental parameters, S , R_H , and M , were found to be remarkably consistent for both the apo- and the holo-states of RC_S. Indeed, when two of these experimentally determined parameters, e.g. R_H and M , were used to calculate (using the Svedberg equation) the third one (S), the theoretical value obtained was very close to the one experimentally measured. This further confirmed the robustness and accuracy of the measured hydrodynamic parameters. Altogether, the hydrodynamic analysis indicated that apo-RC_S was monomeric with a R_H of 2.9 nm, whereas holo-RC_S folded into a stable and compact trimeric conformation exhibiting a R_H of 3.1 nm.

From the intrinsic viscosity of both apo- and holo-RC_S, we then determined the respective contribution of the shape factor and the time-averaged apparent hydration. The method to estimate these parameters has been described elsewhere (9, 10, 25, 26). Briefly, the molecular mass, the intrinsic viscosity, and the R_H (or alternatively the sedimentation coefficient; see “Experimental Procedures”) are required to calculate the shape factor and the protein hydration using the Einstein viscosity relation. The viscosity increment, ν , is a hydrodynamic function (i.e. a shape factor) from which the values of the semi-axes a/b of an ellipsoid can be inferred (27). The time-averaged apparent hydration, δ , defined as the mass of solvent bound to and dragged by the protein along its Brownian motion, is related to the exclusion volume of the protein, i.e. the volume defined by the shearing plane between the protein and the bulk solvent.

As shown in supplemental Table S1, the viscosity increment values, and hence the shape of apo-RC_S and holo-RC_S, were similar. Therefore, the difference between the intrinsic viscosity of apo-RC_S (14.7 ml/g) and holo-RC_S (6.4 ml/g) was mainly due to a difference of hydration (see “Experimental Procedures”). Indeed, apo-RC_S was characterized by a high time-averaged apparent hydration, close to 3 g/g (supplemental Table S1), a value

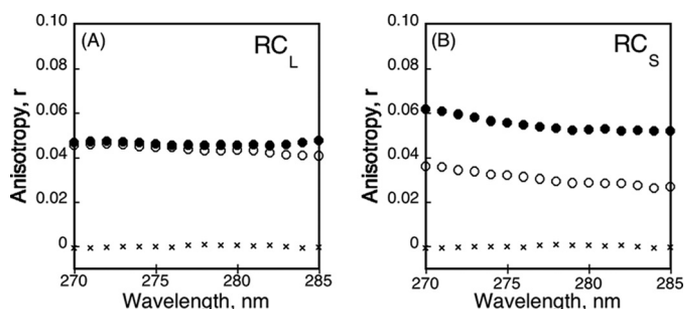


FIGURE 4. **Tryptophan fluorescence anisotropy of RC_L and RC_S.** The fluorescence anisotropy was recorded at an emission wavelength of 350 nm for RC_L (A) and RC_S (B) polypeptides in the absence (○) or in the presence (●) of calcium (2 mM for RC_L and 5 mM for RC_S). In both graphs, L-Trp is represented (×). The experimental conditions were: 2 μM buffer A at 25 °C.

much higher than the theoretical one (0.43 g/g) computed from the protein sequence using the Sednterp software. RC_S underwent a strong dehydration upon calcium binding, although its hydration value (1 g/g) still remained higher than the computed one (supplemental Table S1). The calcium-induced dehydration of RC_S was also evidenced by the changes of the frictional ratio (supplemental Table S1). The frictional ratio ff/f_0 (defined in Table 1 as the ratio of radii, $ff/f_0 = R_H/R_0$) is related, through the Perrin equation (see “Experimental Procedures”), to the hydrodynamic shape function (the Perrin factor) and the time-averaged apparent hydration, δ . Interestingly, the decrease of the frictional ratio from 1.8 for apo-RC_S to 1.35 for holo-RC_S (supplemental Table S1) was essentially assigned to a change of hydration, whereas the shape function was not affected. Similar observations have been reported for the other RTX proteins (Table 1 and Ref. 10), indicating that the strong calcium-dependent dehydration may be a general feature of these polypeptides.

RTX Protein Tryptophan Fluorescence Anisotropy Studies—We further analyzed the properties of RC_S and RC_L by steady-state tryptophan fluorescence anisotropy, a sensitive tool to explore the flexibility of a fluorescent probe. As shown in Fig. 4A, the anisotropy values of apo-RC_L and holo-RC_L were similar and quite low yet higher than that of L-tryptophan used as standard. The low anisotropy value of apo-RC_L was not unexpected because the polypeptide adopted intrinsically disordered conformations in which the tryptophans may be freely moving. However, the fact that the anisotropy of RC_L did not increase upon calcium binding was more surprising, because the tryptophan probes experienced steric constraints from the surrounding amino-acids in the folded holo-RC_L state. We hypothesized that the effects of local constraints on the tryptophan anisotropy may be compensated by the lower hydrodynamic radius, allowing a faster motion (related to the rotational diffusion coefficient of the protein) of holo-RC_L as compared with apo-RC_L. In other words, the effects of the local constraints on RC_L anisotropy (expected increase) may be counterbalanced by the faster rotation of the protein (decreasing anisotropy), thus explaining the absence of a global anisotropy change between the apo- and holo-states. Alternatively, this effect may be due to a nonuniform folding of holo-RC_L: a tryptophan-containing region may remain locally partially unfolded even in the presence of calcium. This scenario is possible

because one of the RC_L tryptophans is located in the C-terminal region.

In contrast, the anisotropy of RC_S was modulated by calcium binding as shown in Fig. 4B. In the apo-state, the anisotropy of RC_S was low likely because of the absence of constraints on the tryptophan residues in this unfolded state, as noted above for apo-RC_L. At variance with RC_L, calcium binding triggered a significant increase of the anisotropy of RC_S. This increase of anisotropy of holo-RC_S may reflect the restriction of the tryptophan side chain flexibility as a result of structural constraints within the folded holo-RC_S. However, because the hydrodynamic volumes of the trimeric holo-RC_S (R_H , 3.1 nm) and of the monomeric apo-RC_S (R_H , 2.9 nm) were similar, the macromolecular dimension of RC_S did not change and therefore should not contribute to any variation of anisotropy. Hence, only the tryptophan side chain flexibility (related to the local tertiary contacts) ultimately contributed to the anisotropy changes of RC_S. Altogether, these data suggest that the anisotropy of RC_L was not modified upon calcium binding due to the compensating effects of local and global changes, whereas that of RC_S increased upon calcium binding due to a unique effect of the folding-induced constraints on the tryptophans.

Calcium-induced Changes of the Electrophoretic Mobilities of RTX Proteins—We recently proposed that electrostatic interactions could play a critical role in the folding and stability of RTX proteins (11). To provide a direct experimental evidence of a charge effect on the unfolded-to-folded transition of the RTX proteins, we characterized the electrophoretic mobility (μ_e), *i.e.* the velocities in a given electric field, of RC_L and RC_S in both the apo- and holo-states. Combined with the above-determined hydrodynamic properties, these data allowed us to estimate the mean net charge (*i.e.* the protein valence) of the polypeptides in the apo- and holo-states. The number of calcium ions bound per protein was then inferred, assuming that the difference in charges between the apo- and the holo-state should correspond to the number of charges brought by the calcium ions bound to the protein in the holo-state.

The approach used to convert the electrophoretic mobility into a number of charges is based on the study of Basak and Ladisch (28) (see “Experimental Procedures”). These authors empirically determined, for a large set of proteins, a relationship between the electrophoretic mobility of a protein (μ_e), the number of charges (N_c), its molecular mass (M), and its frictional ratio (f/f_0) (see the equation under “supplemental materials”). Briefly, this relation indicates that the electrophoretic mobility is proportional to the number of charges and inversely correlated to both the frictional ratio and the molecular mass of the protein. Because we already determined the molecular masses and the frictional ratios of RC_L and RC_S in both of their apo- and holo-states, the measurement of the electrophoretic mobility values should allow us to estimate the number of charges of each species.

The electrophoretic mobility values measured in fast field reversal mode for RC_L and RC_S are shown in Fig. 5. These electrophoretic mobility values, together with the molecular masses and frictional ratios determined previously, are also reported in Table 2. As expected for proteins binding several calcium ions (two charges per cation), large variations of the

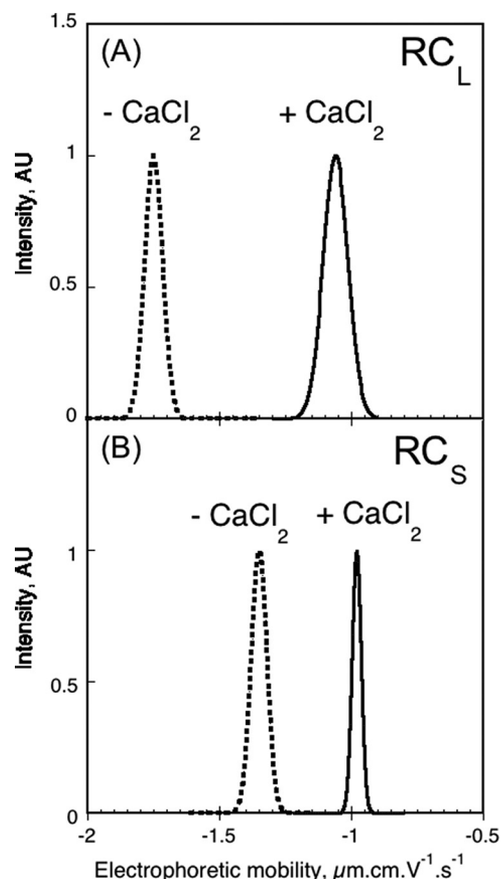


FIGURE 5. **Electrophoretic mobility distribution of RC_L and RC_S.** Electrophoretic mobility of RC_L (A) and RC_S (B) polypeptides, in the absence (*dashed line*) or in the presence (*solid line*) of 2 mM (A), 5 mM (B) CaCl₂. The experimental conditions were: 70 μM RC_S, 130 μM RC_L, and buffer A at 25 °C.

electrophoretic mobility were evidenced between the apo- and the holo-state of each protein. Upon calcium binding, the electrophoretic mobility of RC_L changed from -1.75 to -1.06 $\mu\text{m}\cdot\text{cm}\cdot\text{V}^{-1}\cdot\text{s}^{-1}$ (Fig. 5A). This corresponded to a number of charges of -23.6 for apo-RC_L and -9.1 for holo-RC_L (Table 2). The charge difference between the holo- and apo-forms ($\Delta N_c = -9.1 - (-23.6) = 14.5$) provided an estimation of the number of charges neutralized by the binding of the divalent cation, implicating that approximately seven calcium ions were bound to holo-RC_L ($N_{\text{Ca}^{2+}} = \frac{1}{2}\Delta N_c = 7.25$; Table 2). The electrophoretic mobility of RC_S decreased from -1.35 to -0.98 $\mu\text{m}\cdot\text{cm}\cdot\text{V}^{-1}\cdot\text{s}^{-1}$ upon calcium binding (Fig. 5B). From these values, we estimated a number of charges of -14.9 for the monomeric apo-state and of -15.5 for the trimeric holo-state (Table 2), this latter value corresponding to an averaged charge of -5.2 (*i.e.* $-15.5/3$) per monomer of holo-RC_S. Therefore the charge difference between holo-RC_S and apo-RC_S, ΔN_c , was 9.7 ($\Delta N_c = -5.2 - (-15.4)$) suggesting that approximately five calcium ions were bound per polypeptide in the trimeric holo-RC_S ($N_{\text{Ca}^{2+}} = \frac{1}{2}\Delta N_c = 4.86$ in Table 2).

DISCUSSION

Previous studies have shown that RTX-containing polypeptides are remarkable models to explore the biophysical principles underlying the ligand-induced disorder-to-order transition in proteins (9–11, 24, 29–31). We indeed showed that a

TABLE 2

Electrophoretic mobility of RC_L and RC_S polypeptides

The experimental conditions were as follows: temperature, 25 °C; apo-states, buffer A; in the presence of calcium, buffer A, 2 mM CaCl₂ for holo-RC_L; buffer A, 5 mM CaCl₂ for RC_S.

Parameters	RC _L	RC _L + Calcium	RC _S	RC _S + Calcium
Molecular mass (kDa)	17.1	17.1	14.1	42.3
Frictional ratio (f/f_0) ^a	1.93	1.32	1.83	1.35
μ_e ($\mu\text{m}\cdot\text{cm}\cdot\text{V}^{-1}\cdot\text{s}^{-1}$) ^b	-1.75 ± 0.08	-1.06 ± 0.11	-1.35 ± 0.07	-0.98 ± 0.04
N_c	-23.6 ± 1.1	-9.1 ± 0.4	-14.9 ± 0.7	-15.5 ± 1.1
$N_c/\text{monomer}$	-23.6 ± 1.1	-9.1 ± 0.4	-14.9 ± 0.7	-5.2 ± 0.3
$\Delta N_c = N_{c\text{Holo}} - N_{c\text{Apo}}$ ^c	14.5 ± 1.2		9.7 ± 0.7	
Number of calcium ($N_{\text{Ca}^{2+}}$) ^{c,d}	7.25 ± 0.6		4.86 ± 0.4	
N_c from the amino acid sequence ^e	-23		-19	

^a Experimental data from analytical ultracentrifugation.

^b Experimental data from electrophoretic mobility.

^c Parameters calculated as described under "Experimental Procedures."

^d Number of calcium, $N_{\text{Ca}^{2+}} = (\Delta N_c)/2$.

^e The number of charges, N_c , is calculated as follows: $N_c = |nK + nR| - |nD + nE|$.

700-residue-long polypeptide, corresponding to the RD of the CyaA toxin, is natively disordered (yet fully soluble) in the absence of calcium and folds upon calcium binding into a stable, β -rich structure in a highly cooperative manner (9, 11). Further biophysical studies of a smaller region from the CyaA RTX domain, encompassing a single block of RTX repeats (block V), have shown that the folding of RTX motifs in the presence of calcium is critically dependent upon the adjacent polypeptide segment immediately downstream to the RTX sequences (10, 31). How this C-terminal flanking region and calcium binding could contribute to the folding of the RTX motifs remains to be elucidated.

To address these questions, we have performed here a comprehensive analysis of the hydrodynamic properties of several RTX polypeptides, all containing the RTX core (R) of block V of repeats but differing in their N- or C-terminal flanking regions. We examined more thoroughly the two polypeptides, RC_S and RC_L, made of the RTX core (R) followed by either a short (C_S) or a long (C_L) C-terminal flanking region. Our prior studies have shown that both polypeptides fold into a β -rich structure in the presence of calcium but with a significant difference in affinity for this cation (10).

In the absence of calcium, all of the RTX polypeptides were found to be monomeric and highly hydrated, and they exhibited high hydrodynamic radii, as expected for intrinsically disordered proteins. Upon calcium-induced folding, RC_L experienced a strong compaction as shown by a large decrease of its hydrodynamic radius (Table 1), whereas it remained monomeric (Fig. 2, C and D, and Ref. 10). In contrast, the RC_S polypeptide folded in the presence of calcium into a trimeric state that exhibited roughly the same hydrodynamic radius as the unfolded monomeric apo-form. Hence, these results indicated that the C-terminal flanking region is not only implicated in the calcium-induced folding of the RTX β -helix, but that it also affects the quaternary structure of the polypeptide. Importantly, we could not detect any monomeric or dimeric forms of RC_S, in the presence of calcium, suggesting that holo-RC_S is only stable as a trimer.

The striking difference in behavior of these two polypeptides has important implications for the folding reaction of intrinsically disordered RTX motifs into calcium-loaded β -structures. It has been previously reported that β -helices have the propensity to multimerize (29, 32–36). It has been suggested that intra-

molecular interactions within the β -helices are too weak to stabilize the structure, whereas additional intermolecular interactions may provide further energy to stabilize the assembly of β -helix proteins (16, 32). Moreover, Lilie *et al.* (29) showed that a synthetic polypeptide made of several consensus RTX motifs was able to form a β -helix in the presence of calcium and polyethylene glycol but only upon polymerization, suggesting that intermolecular interactions were indeed essential to stabilize the putative β -roll structure of these synthetic RTX peptides (29).

Our present results strongly support the idea that a β -helix structure made of RTX motifs is thermodynamically weakly stable *per se*. In the RC_S polypeptide, the short C-terminal flanking region may not be sufficient to stabilize the β -helix fold within a monomeric species, whereas the thermodynamic contribution of intermolecular interactions within a trimer of RC_S may provide additional energy to maintain a β -helix conformation. In other words, by restricting the degrees of freedom of the polypeptides, the trimeric state may provide a significant entropic contribution to stabilize the β -helix structure. This may explain why we could not detect any monomeric or dimeric holo-RC_S by SEC-TDA even at the lowest protein concentration tested. The holo-state may only exist as a trimer, which provides the favorable environment to stabilize the calcium-loaded structure. We propose that monomeric RC_S mainly populates the intrinsically disordered state. In the presence of calcium, monomers may experience transient exchanges between calcium-loaded partially folded and unfolded states. Upon diffusion/collision, intermolecular interactions between partially folded RC_S could take place and lead to the formation of a stable trimeric complex. At variance, in RC_L the long C terminus-flanking region may provide enough intramolecular contacts to stabilize the holo-protein in a monomeric form.

The propensity of the RTX β -helices to be stabilized via intermolecular interactions may be pertinent as well for the folding of the whole RD domain that contains five blocks of RTX motifs. Indeed, Iwaki *et al.* (23) initially reported that the C-terminal region of RD encompassing the RTX block V could functionally complement *in vitro* a truncated CyaA (lacking the last 75 residues) to restore its cytotoxic activities. Later, Bejerano *et al.* (22) showed that this C-terminal region of RD could associate in a calcium-dependent manner with the trun-

cated CyaA. Our present results suggest that block V, which can fold as an autonomous domain in the presence of calcium, might provide stabilizing contacts to the upstream blocks (I–IV) of RTX motifs and trigger the calcium-dependent folding of the whole RD domain. Interestingly, we recently found that calcium-induced folding of RD is a highly cooperative process (11). This indicates that, *in vitro*, the loading of the different calcium-binding sites (many of them deviating from the canonical RTX sequences) occurs in a tightly concerted manner. Hence, block V may serve as a nucleus to trigger the folding of the whole RD domain, assuming that, when secreted by the type I secretion machinery, the C-terminal extremity of CyaA is reaching first the calcium-rich extracellular medium.

Our present study also highlighted some important biophysical properties of the RTX polypeptides regarding, in particular, their calcium-induced folding process, which is pertinent for the disorder-order equilibrium of proteins in general. We first compared and described quantitatively the hydrodynamic properties of the apo- and holo-states of both RC_L and RC_S by using the expression that relates the hydrodynamic volume to the molecular mass, the partial specific volume, the time-averaged hydration, and the buffer density.

$$V_H N_A = M(\bar{v} + (\delta/\rho)) \quad (\text{Eq. 1})$$

As previously reported (10), apo- and holo-RC_L exhibited a large difference in their retention volumes on SEC. This variation resulted from a decrease of both the hydrodynamic volume (V_H in the left part of the equation) and the hydration (δ in the right part of the equation) of the protein upon calcium binding. RC_L yet remained a monomer in both the apo- and the holo-state. The response of RC_S to calcium binding was found here to be drastically different. RCs also experienced a large reduction of its hydration upon calcium binding, but at variance with RC_L, it was essentially compensated by an increase in molecular mass, from monomer to trimer, whereas the overall hydrodynamic volume remained unchanged. A plot of the hydrodynamic radii of the RTX polypeptides as a function of their molecular masses, according to Uversky (37), revealed that all of the apo-forms lie between premolten globule conformations and natively unfolded coils, whereas the holo-forms adopt compact conformations (Fig. 6). Also, whereas NRC_L, RC_L, and RD proteins experienced a calcium-induced decrease of their hydrodynamic radius (*i.e.* change along the *y* axis to reach the region of compact proteins), RC_S responded to calcium by a change of its apparent molecular mass (horizontal shift on the *x* axis).

Finally, we attempted to further define the biophysical principles that might control the calcium-induced folding of RTX proteins. We suggested recently that electrostatic interactions might play an important role in the disorder-to-order transition of RD (11). We proposed that, in the apo-form, electrostatic repulsions between the charged Asp residues of the RTX motifs might force the polypeptide chain to adopt expanded and disordered conformations. Calcium binding to the RTX motifs could screen the Asp negative charges, thus allowing the polypeptide chain to collapse and fold into a compact structure.

To directly test this model, we experimentally measured here the changes of the mean net charge induced by the binding of

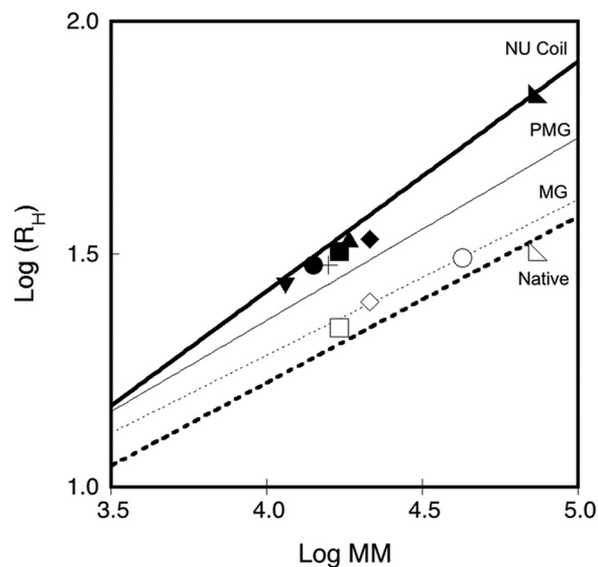


FIGURE 6. Dependences of the hydrodynamic radius on protein molecular mass. The plot shows the hydrodynamic radii (R_H) of the apo-state (closed symbols) and the holo-state (open symbols) of RD (▲), NRC_L (◆), RCL (■), NRC_S (▲), RC_S (●), NR (+), and R (▼) polypeptides as a function of their respective molecular masses. The logarithmic dependences of the R_H on protein molecular mass for native (bold dashed line), molten globule (thin solid line), premolten globule (thin solid line), and natively unfolded proteins (bold solid line) conformations were computed according to the equations described by Uversky (Ref. 37 and the references therein).

calcium to the RTX proteins. Electrophoretic mobility measurements, in conjunction with AUC, QELS, and SEC-TDA (see “Experimental Procedures” for details), allowed us to estimate the mean net charge of both the apo- and the holo-state of RC_S and RC_L. From the mean net charge values, we calculated that the RC_L polypeptide binds ~7 calcium ions, whereas the trimeric holo-RC_S binds ~15 calcium ions, corresponding to roughly 5 calcium ions/monomer on average. These results are in good agreement with our prior calcium binding measurements (10, 24). Interestingly, the somewhat lower number of calcium-binding sites in RC_S may indicate that in the trimeric form of holo-RC_S, the packing constraints may distort or prevent the folding of certain RTX motifs (as revealed by the lowest $n-\pi$ band intensity of RC_S as compared with RC_L in [supplemental Fig. S1](#)). The large changes in the mean net charge of the RTX polypeptides upon calcium binding are interesting to compare in terms of a charge-hydrophobicity phase plot (38). As shown in Fig. 7, the apo-proteins range in the intrinsically disordered region of the charge-hydrophobicity phase space, whereas the strong decrease of the mean net charge upon calcium binding shifts the holo-proteins well below the boundary between unfolded and folded proteins (Fig. 7). To our knowledge, these are the first experimental measurements of ligand-induced changes of polypeptide charges associated with a disorder-to-order transition.

Concluding Remarks—We previously proposed that the intrinsically disordered state of the CyaA RTX motifs in the bacterial cytoplasm may favor the secretion of the toxin by the type I secretion system (9, 10), whereas the calcium-induced folding of the RD domain in the extracellular milieu is critical for the CyaA cytotoxic activity. We showed here that RTX blocks could fold and stabilize upon multimerization, a

Ca-induced Charge Changes and Oligomerization of RTX Motifs

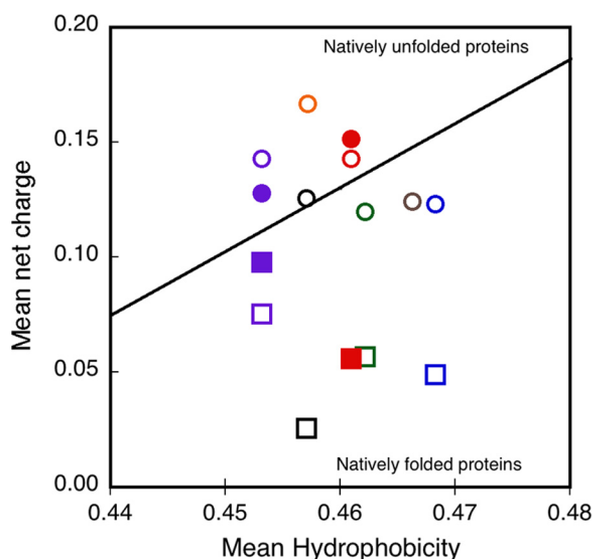


FIGURE 7. Dependence of the mean net charge and the mean hydrophobicity of the block V polypeptides. This plot presents the charge hydrophobicity phase space for both the apo-state (circles) and the holo-state (squares) of RD (black), NRC_L (blue), RC_L (red), NRC_S (green), RC_S (purple), NR (gray), and R (orange) polypeptides. The filled symbols correspond to the mean net charge values of RC_S and RC_L that were experimentally determined herein. The open symbols correspond to the mean net charge values of the RTX polypeptides that were theoretically computed from the amino acid sequence and considering that block V polypeptides and the full RD protein should bind 7 and 35 calcium ions, respectively (24). The theoretically computed charge value of holo- RC_L (open red square) is hidden by the experimental measurement of the protein charge (closed red square). The oblique line represents the boundary between intrinsically disordered and folded proteins according to Uversky *et al.* (38).

cooperative process that might contribute to improve the secretion efficiency. We further showed that the apo-forms of RTX block V polypeptides adopt a premolten globule state that switches to a native state in the presence of calcium. This transition appeared finely tuned from a thermodynamic point of view to respond to calcium binding in physiological conditions. Moreover, we showed that the high number of calcium bound per protein allows switching from an intrinsically disordered, highly charged apo-form to a folded holo-state with a reduced mean net charge. Our data suggest that the folding of the RTX proteins is controlled by a charge change effect caused by the variations of physiological concentrations of calcium. The simplicity of this calcium-induced, charge-driven disorder-to-order transition may favor a robust and chaperone-independent protein folding process adapted to the biogenesis of the RTX proteins.

Acknowledgments—We thank Bertrand Raynal for expertise and help on AUC and for the critical reading of the manuscript. We also thank J. D'Alayer for the N-terminal sequencing of the proteins and Johanna C. Karst for fruitful comments on this manuscript.

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