Interdomain Flexibility in Full-length Matrix Metalloproteinase-1 (MMP-1)*

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The presence of extensive reciprocal conformational freedom between the catalytic and the hemopexin-like domains of full-length matrix metalloproteinase-1 (MMP-1) is demonstrated by NMR and small angle x-ray scattering experiments. This finding is discussed in relation to the essentiality of the hemopexin-like domain for the collagenolytic activity of MMP-1. The conformational freedom experienced by the present system, having the shortest linker between the two domains, when compared with similar findings on MMP-12 and MMP-9 having longer and the longest linker within the family, respectively, suggests this type of conformational freedom to be a general property of all MMPs.

Matrix metalloproteinases (MMP) are extracellular hydrolytic enzymes involved in a variety of processes including connective tissue cleavage and remodeling (1–3). All 23 members of the family are able to cleave simple peptides derived from connective tissue components such as collagen, gelatin, elastin, etc. A subset of MMPs is able to hydrolyze more resistant polypeptide substrates, such as cross-linked elastin, and partially degraded collagen forms, such as gelatin and type IV collagens (4). Intact triple helical type I–III collagens is only attacked by collagenases MMP-1, MMP-8, and MMP-13 and by MMP-2 and MMP-14 (5–12). Although the detailed mechanism of cleavage of single chain peptides by MMP has been largely elucidated (13–19), little is known about the process of hydrolysis of triple helical collagen. In fact, triple helical collagen cannot be accommodated in the substrate-binding groove of the catalytic site of MMPs (9).

All MMPs (but MMP-7) in their active form are constituted by a catalytic domain (CAT) and a hemopexin-like domain (HPX) (20–22). The CAT domain contains two zinc ions and one to three calcium ions. One zinc ion is at the catalytic site and is responsible for the activity, whereas the other metal ions have structural roles. The isolated CAT domains retain full catalytic activity toward simple peptides and single chain polypeptide substrates such as elastin, whereas hydrolysis of triple helical collagen also requires the presence of the HPX domain (9, 23–25). It has been shown that the isolated CAT domain regains a small fraction of the activity of the full-length (FL) protein when high amounts of either inactivated full-length proteins or isolated HPX domains are added to the assay solution (9). Finally, it has been shown that the presence of the HPX domain alone alters the CD spectrum of triple helical collagen in a way that suggests its partial unwinding (26, 27). It is tempting to speculate that full-length collagenases attack collagen by first locally unwinding the triple helical structure with the help of the HPX domain and then cleaving the resulting, exposed, single filaments (9, 28).

Until 2007, three-dimensional structures of full-length MMPs had been reported only for collagenase MMP-1 (29–31) and gelatinase MMP-2 (32). The structures of the two proteins are very similar and show a compact arrangement of the two domains, which are connected by a short linker (14 and 20 amino acids, respectively). It is difficult to envisage that rigid and compact molecules of this type can interact with triple helical collagen in a way that can lead to first unwinding and then cleavage of individual filaments. It has been recently suggested that such concerted action could occur much more easily if the two domains could enjoy at least a partial conformational independence (9). Slight differences in the reciprocal orientation of the CAT and HPX domains of MMP-1 in the presence (29) and absence (30, 31) of the prodomain were indeed taken as a hint that the two domains could experience relative mobility (29).

Two recent solution studies have shown that conformational independence is indeed occurring in gelatinase MMP-9 (33) and elastase MMP-12 (34), whereas the x-ray structure of the latter (34) is only slightly less compact than those of MMP-1 (29–31) and MMP-2 (32). Among MMPs, MMP-9 features an
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exceptionally long linker (68 amino acid) (33, 35), which in fact constitutes a small domain by itself (the O-glycosylated domain) (33), and therefore, this inspiring observation can hardly be taken as evidence that conformational freedom is a general characteristic of the two-domain MMPs. MMP-12 features a much more normal 16-amino acid linker, thereby making more probable a general functional role for this conformational freedom (34). However, both MMP-9 and MMP-12 retain their full catalytic activity against their substrates even when deprived of the HPX domain (9). Therefore, the question remains of whether conformational freedom is also a required characteristic for those MMPs that are only active as full-length proteins, i.e., collagenases. Interestingly, the three collagenases (MMP-1, MMP-8, and MMP-13) have the shortest linker (14 amino acids) among all MMPs. Demonstrating or negating the presence of conformational freedom in one of these collagenases would therefore constitute a significant step forward to formulate mechanistic hypotheses on their collagenolytic activity.

Our recent studies on MMP-12 in solution (34) have shown that a combination of NMR relaxation studies and small angle x-ray scattering (SAXS) is enough to show the presence and the extent of the relative conformational freedom of the two domains of MMPs. Here we apply the same strategy to full-length MMP-1 and show that sizable conformational freedom is indeed experienced even by this prototypical collagenase, although somewhat less pronounced than that observed for MMP-12.

EXPERIMENTAL PROCEDURES

Preparation of Protein Samples—The cDNA encoding the sequence (Asn-106–Asn-469) of the MMP-1 full-length protein was amplified from TrueClone cDNA (OriGene) by PCR and cloned into the pET21 (Novagen) expression vector using NdeI and XhoI (New England Biolabs) as restriction enzymes. One additional methionine at position 105 was present in the final expression product. The recombinant vector was transformed into Escherichia coli strain BL21(DE3) CodonPlus RIPL (Stratagene), and colonies were selected for ampicillin and chloramphenicol resistance. The bacteria were grown in 2 × YT medium. When a cell density corresponding to 0.6 A was reached, the expression of the protein was induced by adding 0.5 mM isopropyl-β-D-thiogalactoside, and the incubation at 37 °C was continued for another 5 h. The full-length MMP-1 precipitated in the inclusion bodies, and these were solubilized, after lysis of the cells, in a solution of 8 M urea, 20 mM dithiothreitol, and 20 mM Tris-HCl, (pH 8). The protein was diluted with a buffer containing 6 M urea, 10 mM CaCl₂, 0.1 mM ZnCl₂, 20 mM cystamine, 20 mM Tris-HCl (pH 8) and refolded by using a multistep dialysis against solutions containing 50 mM Tris-HCl (pH 8), 4 M urea, 10 mM CaCl₂, 0.1 mM ZnCl₂, 5 mM B-mercaptoethanol, 1 mM 2-hydroxyethyl disulfide, then against a solution containing 50 mM Tris-HCl (pH 7.2), 2 M urea, 10 mM CaCl₂, 0.1 mM ZnCl₂, 0.3 M NaCl, and then against the same solution without urea. The protein was purified by size-exclusion chromatography on the HiLoad 26/60 Superdex 75 pg (Amersham Biosciences). For expression of 15N- and 13C-enriched protein samples at concentrations ranging between 0.3 and 0.7 mM (pH 7.2). For FL-MMP-1, all NMR experiments were performed on samples at a concentration of 0.3 mM (pH 7.2). NMR experiments were performed at 298 and 306 K and acquired on Bruker AVANCE 900, AVANCE 800, AVANCE 700, AVANCE 600, and DRX 500 spectrometers equipped with triple resonance cryo-probes. All spectra were processed with the Bruker TOPSPIN software packages and analyzed by the program CARA (Computer Aided Resonance Assignment, ETH Zürich) (38). The backbone resonance assignment of HPX domain was obtained by the analysis of HNCA, HNCACB, and CBCA-(CO)NH spectra performed at 500 and 900 MHz, whereas the assignment of the 2H-, 15N-, and 13C-enriched FL-MMP-1 was obtained by the analysis TROSY-HNCA, TROSY-HNCA/B, TROSY-HNCO/CAB performed on an 800-MHz spectrometer. The obtained assignments are reported in supplemental Table S1 for the full-length protein, in supplemental Table S2 for the catalytic domain, and in supplemental Table S3 for the hemopexin-like domain.

The protein assignment and the mobility measurements on FL-MMP-1 were performed on the NNGH-inhibited, E219A mutant, due to its high stability to the self-hydrolysis. Mobility
measurements on the catalytic domain were performed on the NNGH-inhibited form of the protein.

*Rg*, *Rg* and NOE Measurements—The experiments for the determination of 15N longitudinal and transverse relaxation rates and 1H-15N NOE were recorded at 306 K and 700 MHz on 15N-enriched samples. The 15N longitudinal relaxation rates (*R1*) were measured using a sequence modified to remove cross correlation effects during the relaxation delay (39). Inversion recovery times ranging between 2.5 and 3000 ms, with a recycle delay of 3.5 s, were used for the experiments. The 15N transverse relaxation rates (*R2*) were measured using a Carr-Purcell-Meiboom-Gill (CPMG) sequence (39, 40) with delays ranging between 8.5 and 237.4 ms for the CAT domain, between 8.5 and 203.5 ms for the HPX domain, and finally between 8.5 and 135.7 ms for the FL-MMP-1 protein with a refocusing delay of 450 μs. The relaxation data are reported in supplemental Table S4.

The relaxation data were processed using standard procedures and extrapolated to infinite dilution using the program PRIMUS (42). The data were processed using the Guinier approximation (43), assuming that the maximum dimension, *Dmax*, and the interatomic distance distribution functions, (p(r)), were also computed using the program GNOM (44). The scattering from the high resolution models was computed using the program CRYOSOL (45). Given the atomic coordinates, the program predicts the theoretical scattering pattern and fits the experimental intensity by adjusting the excluded volume of the particle and the contrast of the hydration layer to minimize the discrepancy

\[ \chi^2 = \frac{1}{N-1} \sum \left( \frac{I_{\text{exp}}(s_j) - I_{\text{calc}}(s_j)}{\sigma(s_j)} \right)^2 \]

where *N* is the number of experimental points, *c* is a scaling factor, *I_{\text{exp}}(s_j)* and *I_{\text{calc}}(s_j)* are the experimental and calculated intensities, respectively, and \(\sigma(s_j)\) is the experimental error at the momentum transfer *s*. To assess the conformational variability of MMP-1, an ensemble optimization method (EOM) was used (46), allowing for the coexistence of multiple conformations in solution. About 10,000 randomized models of FL-MMP-1 differing by the conformation of the interdomain linker were generated, and their scattering patterns were computed using the program RanCh of the EOM package. These models formed a pool of possible structures for which the scattering patterns were computed by CRYOSOL. The EOM program employs a genetic algorithm to select from the pool a small number (usually about 20) of representative structures such that the average scattering from the selected ensemble fits the experimental data. Multiple runs of EOM were performed, and the results were averaged to provide quantitative information about the flexibility of the protein in solution (in particular, about the *Rg* distribution in the selected ensembles).

**RESULTS AND DISCUSSION**

Full-length human MMP-1 and its CAT-(106–261) and HPX-(274–469) domains were expressed as described under “Experimental Procedures.” In both FL-MMP-1 and its CAT domain, an E219A mutation was introduced. Such mutation ensures a dramatic lowering of the catalytic activity (13, 19), which is needed in the present investigation to prevent self-hydrolysis. In addition, the strong active site-directed hydroxamic inhibitor NNGH (15) was always added. Finally, when needed, cadmium(II) was substituted for zinc(II) in the catalytic site to further reduce the residual activity as it was already shown for FL-MMP-12 and its CAT domain (34). 15N-, 13C-, 15N-, and 2H-13C-15N-enriched samples were used for NMR investigations. The samples were stable under the NMR experimental conditions for several months. For the present mobility studies, the assignment of the 1H-15N HSQC spectra of the three constructs was needed. The two-dimensional and three-dimensional experiments that were used for this purpose are described under “Experimental Procedures.” The 1H-15N-TROSY-HSQC spectra of the three constructs, acquired at 306 K, are superimposed in Fig. 1. The spectral quality, especially for the FL and HPX domains, is better at 306 K than at 298 K. This prompted us to perform all subsequent work at 306 K. Moreover, the latter temperature is closer to the physiological value of 310 K, so the resulting mobility data will be more meaningful.
Analysis of the Chemical Shift Differences between FL-MMP-1 and Its Isolated Domains—For clarity, the spectrum of the full-length protein in Fig. 1 is from a triply labeled $^{2}$H, $^{13}$C, $^{15}$N sample, which displays much sharper lines. From visual inspection, it already appears that the FL spectrum is largely the superposition of the CAT and HPX spectra as the majority of the signals in the FL spectrum overlap with either a CAT or an HPX signal. There are a number of additional signals in the FL spectrum that are attributed to the portion of the interdomain linker (262–275), which is missing in both the CAT and the HPX isolated domains. Furthermore, a few more signals undergo a modest shift on passing from the isolated domains to the FL protein. These signals could arise from contacts between each domain and the linker, contacts between the two domains, or both. In the latter two cases, evidence of interdomain contacts would imply that either the FL length structure is rigid or, that even in the presence of conformational freedom, “closed” compact structures contribute to the overall description of the molecule.

The assignments for the three constructs are reported in supplemental Tables S1–S3. From a comparison of the $^{1}$H and $^{15}$N chemical shifts in the $^{1}$H-$^{15}$N-TROSY-HSQC spectra for the FL and the two isolated domains, a mean shift difference plot (47) was generated (Fig. 2). It appears that the number of peaks experiencing chemical shift differences is rather limited and clustered in the 112–115, 139–149, 244–261, and 277–316 regions. Fig. 3A shows the experimental x-ray three-dimensional structure of the FL protein (30) as well as of its isolated domains and linker, exploded to show the regions of contact between CAT and HPX domains which is missing in both the CAT and the HPX isolated domains. Furthermore, a few more signals undergo a modest shift on passing from the isolated domains to the FL protein. These signals could arise from contacts between each domain and the linker, contacts between the two domains, or both. In the latter two cases, evidence of interdomain contacts would imply that either the FL length structure is rigid or, that even in the presence of conformational freedom, “closed” compact structures contribute to the overall description of the molecule.

Relaxation Measurements—Measurements of longitudinal ($R_1$) and transverse ($R_2$) relaxation rates of backbone amide nitrogens at 700-MHz $^{1}$H Larmor frequency and 306 K have been performed on $^{15}$N-enriched samples of both the isolated CAT domains and the isolated HPX domains as well as of FL-MMP-1. Estimates of $R_1$ and $R_2$ values for these three constructs under the chosen experimental conditions of magnetic
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field and temperature were obtained using the program HydroNMR (48), and the three-dimensional structures of the constructs were taken from the x-ray structure of the FL protein (30). The experimental and theoretical $R_1$ and $R_2$ values are reported in Fig. 4, A–E, together with the corresponding data for MMP-12 (Fig. 4, F–L) (34) at the same field and 298 K, for comparison purposes. If the experimental data do not agree with the calculated data, outside the experimental error, then they are inconsistent with the structural model. As far as the catalytic domain is concerned (left side), it is clear that the $R_1$ and $R_2$ data for the isolated domain are in excellent agreement with the theoretical expectations. On the contrary, the experimental $R_1$ data for the CAT domain, when it is part of the FL protein, are sizably higher, and the $R_2$ data are sizably lower, with respect to the theoretical values for a rigid FL protein. Higher $R_1$ and lower $R_2$ values, taken together, indicate that the two domains behave as if they belonged to a lower molecular weight protein; that is, the CAT domain possesses some degree of motion, which is independent of the motion of the FL protein as a whole.

In principle, the same should hold for the HPX domain. Indeed, the $R_1$ and $R_2$ data for the HPX domain in the full-length protein are also higher and lower, respectively, than the theoretical estimate, but here the difference is less marked than for the CAT domain. A plausible explanation lies in the fact that the $R_1$ and $R_2$ data for the isolated HPX domain show the opposite behavior, i.e. the $R_1$ values are lower and the $R_2$ values are higher than expected. This is typically encountered in the presence of self-aggregation. The HPX domain does possess a rather hydrophobic surface, which could allow for the existence of a fraction of dimeric or higher order aggregation species in solution. Partial aggregation brought about by the HPX domain in the FL protein would of course artificially attenuate the effect of interdomain mobility on $R_1$ and $R_2$. The hypothesis of HPX aggregation is easy to check by changing the protein concentration. Indeed, for isolated HPX solutions of higher concentration, the discrepancy with the calculated data sizably increases (data not shown). In conclusion, the HPX data (i) demonstrate the presence of aggregation but also (ii) confirm interdomain mobility in the full-length protein, which is still apparent from $R_1$ and $R_2$ data despite the attenuation of the effect due to aggregation.

A further proof of the interdomain mobility is provided by the very small NOE values of four adjacent residues in the linker region (267, 268, 270, and 271) (Fig. 4E). Comparison with the MMP-12 data (Fig. 4, F–L), collected at lower temperature, shows that interdomain mobility in MMP-1, although certainly present, is less extensive than in MMP-12. This is possibly related (i) to the shorter linker (14 versus 16 residues) and (ii) to the more extensive interdomain contact present in the x-ray structure of active FL-MMP-1 with respect to that observed in the x-ray structure of MMP-12.

A Paramagnetic Probe Monitors Interdomain Surface Exposure—Paramagnetic probes dissolved in protein solutions have been previously used (i) to identify surface residues to help structure determination (49–56), (ii) to determine the orientation of membrane-bound peptides on the membrane surface (57–59), and (iii) to monitor the formation of protein-protein
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complexes by detecting residues that become shielded from the probe upon complex formation (4, 60, 61). In analogy with the latter strategy, we have used the gadolinium complex Gd(DTPA-BMA) (57) to probe the CAT-HPX interface in the FL protein. An inversion recovery HSQC sequence was used to measure the $R_1$ values of the backbone NH protons of FL-MMP-1 in the absence ($R_{1\text{dia}}$) and in the presence ($R_{1\text{para}}$) of 1.4 mM Gd(DTPA-BMA), as detailed under “Experimental Procedures.” The residues for which the relaxation rate difference $R_{1\text{para}} - R_{1\text{dia}}$ is higher than a threshold value of 0.7 s$^{-1}$ are reported in supplemental Table S5. Among them, on the HPX side, residue Leu-314 should be shielded by the CAT domain; on the CAT side, residue Asp-231 should be shielded by the linker, and residues Asp-245, Val-246, and Gln-247 should be shielded by the HPX domain. On the contrary, all of them experience paramagnetic effects particularly strong for Asp-245 (2.7 s$^{-1}$) and Val-246 (2.4 s$^{-1}$). These effects are inconsistent with the compact x-ray structure of FL-MMP-1 and confirm that the protein must spend a non-negligible fraction of time in an extended conformation so that the CAT-HPX interface becomes accessible to the probe.

SAXS Measurements—The processed x-ray scattering pattern from FL-MMP-1 is shown in Fig. 5. The experimental radius of gyration $R_g$ is $29 \pm 1$ Å. This value significantly exceeds that calculated ($R_g = 25$ Å) from the x-ray structure of active FL-MMP-1 (30). Moreover, the scattering pattern calculated from the active FL-MMP-1 model using CRYSOL (45) fails to fit the experimental data (discrepancy $\chi = 4.22$, Fig. 5, curve 2). Interestingly, the fit to the FL-MMP-12 model is better (discrepancy $\chi = 1.8$, Fig. 5, curve 3). If a distribution of FL-MMP-1 conformations, including compact and extended ones, is taken into consideration, neither individual models nor averaging over the random pool allowed one to fit the SAXS data satisfactorily ($\chi > 1.9$ in both cases). The representative ensembles selected to fit the data give information about the preferable conformations of the protein. To assess the preferable conformations in solution, the EOM method (46) was used. Given a representative pool of (random) structures, the method employs a genetic algorithm to select the ensembles from this pool that best fit the experimental data, as explained under “Experimental Procedures.” Several EOM runs yielded reproducible ensembles neatly fitting the experimental data with discrepancy $\chi$ less than 1.2, and a typical fit provided by the ensemble selected by EOM is given in Fig. 5, curve 4. All the fits from different EOM runs are graphically indistinguishable from Fig. 5, curve 4. The $R_g$ distributions of the particles in the initial pool and in the selected ensembles are compared in Fig. 5 (inset). The former distribution is rather broad and covers the $R_g$ range from about 20 to 45 Å, corresponding to extremely compact and completely extended domain configurations, respectively. In contrast, the $R_g$ distribution of the selected ensembles displays a relatively sharp peak around $R_g = 25–26$ Å, including about 70% of the particles in the selected ensembles. Visual inspection of the models having $R_g$ values in the peak range indicates, not unexpectedly, that their shape is similar to that of the FL-MMP-1 structure in the crystal (30) (although with varying interdomain orientations). However, an extended tail at higher $R_g$ values up to around 40 Å, accounting for about 30% of the particles, is apparent from the multiple EOM reconstructions. These results indicate that the crystallographic conformation of FL-MMP-1 (30) is largely present also in solution, but in addition, the protein experiences noticeable flexibility with a significant amount of extended conformations in equilibrium with the closed one(s). We also tried to explore the possibility of a two-state exchange situation allowing for only two conformations in the mixture. In such a case, two structures with an $R_g$ of around 25 and 32 Å are selected. It is interesting that FL-MMP-1 (30) appears much more compact than FL-MMP-12 (34), as shown by the comparison in Fig. 5, inset, indicating that the two extra residues in the linker region of FL-MMP-12 increase its flexibility.

Concluding Remarks and Biological Implications—The present data demonstrate that FL-MMP-1 shows relative mobility of its catalytic and hemopexin-like domains, as recently observed for FL-MMP-12 (34). As in the latter case, the reorientation of the backbone NH vectors with respect to the magnetic field occurs on a time scale that is faster than the rotational time of the whole molecule (62). The amplitude of the motion is probably smaller for FL-MMP-1 than it is for FL-MMP-12, as judged from the SAXS data that suggest that the molecules spend two-thirds of the time in a conformation that is more or less as compact as the solid state structure but is ample enough that the residues at the interface between...
the two domains are significantly exposed to a paramagnetic probe in solution. Besides MMP-12, relative mobility of the CAT and HPX domains has also been recently suggested for MMP-9, where the linker between the two domains is much longer and constitutes a domain by itself (O-glycosylated domain) (63). In the case of MMP-9, it has been argued (33) that the long O-glycosylated domain may be flexible and may mediate protein-substrate interactions (64, 65). However, the present finding of flexibility in FL-MMP-1 has further biological implications because MMP-1 is a collagenase that catalysis has often been invoked as a means to unwind the domain is crucial for the function. Notably, the possibility of the presence of the HPX domain is crucial for the fact. Notably, the possibility of reorienting the HPX with respect to the CAT domain during catalysis has often been invoked as a means to unwind the collagen triple helix in preparation for the catalytic cleavage (66).

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