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Julien Lefèvre, Philippe Savarin, Pierre Gans, Loïc Hamon, Marie-Jeanne Clément, et al.. Structural Basis for the Association of MAP6 Protein with Microtubules and Its Regulation by Calmodulin.: Microtubule and calmodulin binding on Mn modules of MAP6. Journal of Biological Chemistry, American Society for Biochemistry and Molecular Biology, 2013, 288 (34), pp.24910-22. <10.1074/jbc.M113.457267>. <inserm-00858655>

> HAL Id: inserm-00858655 http://www.hal.inserm.fr/inserm-00858655

> > Submitted on 7 Jul2014

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Microtubule and calmodulin binding on Mn modules of MAP6

Structural Basis for the Association of MAP6 with Microtubules and its regulation by Calmodulin

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*Running title: Microtubule and calmodulin binding on Mn modules of MAP6

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Keywords: Calcium; Calmodulin; Cold; MAP6; Microtubules; NMR; STOP; Tubulin

Background: Microtubules are cold-sensitive but some cold-stable microtubules are observed in specific cells due to the presence of MAP6.

Results: Structural data detail how a MAP6 fragment stabilizes microtubules and how calmodulin regulates its activity.

Conclusion: MAP6 may stabilize microtubules by bridging adjacent tubulin heterodimers, an activity sterically hindered by calmodulin.

Significance: This work provides a better understanding of cellular microtubule stabilization and its regulation by calmodulin.

SUMMARY

Microtubules are highly dynamic $\alpha\beta$ -tubulin polymers. In vitro and in living cells, microtubules are most often cold and nocodazole sensitive. When present, the MAP6/STOP family of proteins protects microtubules from cold and nocodazole induced depolymerization but the molecular and structure determinants by which these proteins stabilize microtubules remain under debate. We show here that a short protein fragment from MAP6-N which encompasses its Mn1 and Mn2 modules (MAP690,177) recapitulates the function of the full-length MAP6-N protein toward microtubules, i.e. its ability to stabilize microtubules in vitro and in cultured cells in ice-cold condition or in the presence of nocodazole. We further show for the first time, using biochemical assays and NMR spectroscopy, that these effects result from the binding of MAP6₉₀₋₁₇₇ to microtubules with a 1:1 MAP6₉₀₋₁₇₇:tubulin heterodimer stoichiometry. NMR data demonstrate that the binding of MAP6₉₀₋₁₇₇ to microtubules involve its two Mn modules but that a single one is also able to interact with microtubules in a closely similar manner. This suggests that the Mn modules represent each a full microtubule binding domain and that MAP6 proteins may stabilize microtubules bv bridging tubulin heterodimers from adjacent protofilaments or within a protofilament. Finally. we demonstrate that Ca²⁺-calmodulin competes with microtubules for MAP6₉₀₋₁₇₇ binding and that the binding mode of MAP690-177 to

microtubules and Ca²⁺-calmodulin involves a common stretch of amino acid residues on the MAP6₉₀₋₁₇₇ side. This result accounts for the

In certain cell types such as neurons, fibroblasts or glial cells, a subpopulation of microtubules (MTs) remains stable in depolymerizing conditions such as exposure to the cold or depolymerizing drugs (1-5). Search for cellular factors responsible for this stabilization led to the discovery of a class of microtubule-associated proteins named MAP6 (also called STOP) (6-8). In mammals, MAP6 isoforms are encoded by a single gene (map6) and result from mRNA splicing and alternative promoter use (9-11). The physiological role of MAP6 proteins is not yet fully understood but phenotypic and cellular analyses of MAP6 null mice indicated that MAP6 proteins are involved in a number of neuronal functions. MAP6 null mice present synaptic plasticity defects in and neurotransmission associated with severe behavioral disorders (12). Interestingly, these disorders can be alleviated by drugs like neuroleptics suggesting that these mice may represent a valuable animal model for the study of schizophrenia (13). In line with these findings, genetic studies have pointed out a possible link between MAP6 and schizophrenia (14) suggesting that *map6* could be considered as a candidate gene that predispose to schizophrenia and thus be used as a potential biomarker for its early detection (15,16). Besides, MAP6 proteins were also found to be implicated in adult olfactory neurogenesis (17) and MAP6-F (the fibroblastic MAP6 isoform) was proposed to be a temperature sensor that protects MTs from temperature variations in animals during episodes of torpor or hibernation (18).

MAP6-N (the neuronal MAP6 isoform) is the largest MAP6 isoform and is mainly expressed in mature neurons. This 952 residues protein contains several repeated motifs called Mn and Mc modules which are linked to its ability to stabilize MTs (Fig. 1A) (19). The Mn modules allow MAP6 to protect MTs against cold- and nocodazole-induced depolymerization while the Mc modules permit MAP6 to stabilize MTs in condition. MAP6-N contains cold two consecutive Mn modules in its N-terminal part (Mn1 and Mn2) and an additional one isolated in the middle of its sequence (Mn3). The Mc

regulation of microtubule stability in cold condition by Ca^{2+} -calmodulin.

modules are clustered in the central domain as five highly conserved repetitions organized in tandem. Whereas Mn modules are well conserved across species (Fig. 1B), the homology between Mn1, Mn2 and Mn3 is low and they share only a few residues (Fig. 1C). It is worth noting that the number and the organization of Mn and Mc modules can be different among the MAP6 isoforms and this may lead to functional variations toward the propensity of MAP6 proteins to stabilize MTs (8,18,19). In addition, it was shown that the MT stabilization activity of MAP6 proteins is inhibited by calmodulin (CaM) (20,21).Accordingly, peptide arrays revealed that MAP6-N possesses several CaM binding sites, some of which being found in the Mn and Mc modules suggesting that these modules have a dual propensity to interact both with MTs and CaM (19). Very few studies addressed the structural basis of the interaction of MAP6 with MTs and of its interplay with CaM. An NMR study of the interaction between CaM and a peptide model corresponding to the consensus Mc motif showed an unusual interaction mode with a very limited structural impact on the CaM structure (22).

In the present work, we undertook biochemical and structural analyses of the interaction of MAP6-N with MTs and CaM to enlighten the mechanism by which it stabilizes MTs and the mechanisms by which such stabilization is regulated by CaM. As MAP6-N has a high molecular weight, we focused this work on a minimal MAP6-N fragment (residues 90-177, named herein MAP6₉₀₋₁₇₇) which recapitulates the MT stabilization and CaM binding properties of MAP6 proteins. We show here that the MAP6₉₀₋₁₇₇ fragment, which comprises the Mn1 and Mn2 modules, interacts tightly with MTs and has a powerful MT stabilization activity in vitro and in cultured cells. It also interacts with CaM in a Ca²⁺-dependent manner. Using a combination of biochemical and NMR experiments, we determined at the residue scale the location of its MT and CaM binding sites and we clearly demonstrated that Ca²⁺-CaM binding impairs MAP690-177 association with MTs

probably because the Ca²⁺-CaM binding sites substantially overlap the MT binding sites. Furthermore, we examined by NMR the structural impact of MAP6₉₀₋₁₇₇ binding on the Ca²⁺-CaM structure. The results showed dramatic conformational and dynamical changes of Ca²⁺-CaM affecting its overall structure which led us to propose a binding mode for Ca²⁺-CaM on the Mn modules of MAP6 proteins.

EXPERIMENTAL PROCEDURES

Transient transfection and immunofluorescence microscopy

The pCMV-FLAG-MAP690-177 plasmid encodes a fusion protein comprising residues 90 to 177 of Rattus norvegicus MAP6-N and was constructed by PCR using a derivative of the pSG5-STOP plasmid as template (8). HeLa cells were grown at 37 °C with 5% CO₂ in Dulbecco's modified Eagle's medium (DMEM) containing 5% fetal bovine serum and 1% penicillin-streptomycin. 5 µg of pCMV-FLAG-MAP690-177 plasmid was mixed with 5 µL of lipofectamine 2000 (Invitrogen, Carlsbad, USA) and then diluted in 200 µL of opti-MEM I. This mix was left for 20 min at 20°C and then added to the HeLa cells culture medium and gently mixed. Cells were returned to the incubator for 4 h at 37°C and the medium was replaced with fresh full medium. 48 h post-transfection, cells were either exposed to cold (on ice for 30 min) or to nocodazole (20 µM for 30 min at 37°C). Cells were then washed twice with PBS and fixed/permeabilized with paraformaldehyde (30 min at 4°C) and cooled methanol (5 min at -20°C). After washing, cells were incubated 1 h at 37 °C with the blocking solution (20 mM Tris-HCl, 150 mM NaCl, 0.1% Triton, 2% BSA, 0.1% NaN₃, pH 7.4). The MT network and MAP690-177 were revealed by immunofluorescence with E7 anti- α -tubulin mouse antibody (1:3000) and anti-FLAG rabbit antibody (1:160) (Sigma-Aldrich, Saint-Louis, USA). Cells were washed twice with PBS and incubated with goat anti-mouse (Alexafluor 488) and anti-rabbit (Alexafluor 594) antibodies. The cells were finally washed with PBS and examined for fluorescence with a Zeiss microscope using a 63x/1.4 numerical aperture objective.

Protein expression and purification

The MAP6₉₀₋₁₇₇ fragment was overexpressed in a recombinant form in fusion with an N-terminal polyhistidine tag using the pET-46 Ek/LIC plasmid (Novagen-Merck, Darmstadt, Germany). After transformation with the pET-46 Ek/LIC plasmid, E. coli BL21 DE3* cells (Invitrogen, Carlsbad, USA) were grown in LB medium at 37°C with 0.1 mg/mL ampicillin in a 1L flask. $^{15}N-^{13}C$ For uniform isotopic labeling, transformed bacteria were grown in M9 minimal medium containing 0.6 g/L 95% ¹⁵NH₄Cl and 2.2 g/L 95% ¹³C-glucose (Cortecnet, Paris, France) as the sole nitrogen and carbon sources respectively. In all cases, overexpression was induced at $OD_{600 \text{ nm}} = 0.7$ with 1 mM IPTG for 2 h. Bacteria were pelleted by low speed centrifugation (3,200 x g, 10 min) and then resuspended in 10 mL of buffer A (20 mM Tris-HCl, 100 mM NaCl, 1 mM DTT, pH 7.6). After lysis by sonication, the lysate was heated for 25 min at 95°C and then ultracentrifuged at 4°C (100,000 x g, 1 h). The clarified cell lysate was then loaded on a Ni-NTA column (Qiagen, Hilden, Germany) equilibrated with buffer A supplemented with 10 mM imidazole. The proteins were eluted with buffer B (20 mM Tris-HCl, 100 mM NaCl, 1 mM DTT, 300 mM imidazole, pH 7.6). Fractions containing the MAP6₉₀₋₁₇₇ were combined and concentrated by ultrafiltration (Amicon 3 kDa cutoff, Millipore) at 4°C to 0.5 mL and diluted into 4.5 mL buffer C (50 mM MES-KOH, 150 mM NaCl, 1 mM DTT. pН 6.9). The procedure of concentration/dilution was repeated three times with buffer C to eliminate imidazole. Final concentration of MAP6₉₀₋₁₇₇ was determined by amino acid analysis. The result was used to determine the extinction coefficient of MAP690- $_{177}$ ($\epsilon_{280nm} = 2.3 \times 10^4 \text{ M}^{-1} \cdot \text{cm}^{-1}$). The final yield was about 1 mg/L of pure protein. The MAP690-142 fragment (residues 90-142 of Rattus norvegicus MAP6-N) was obtained using the same protocol ($\varepsilon_{280nm} = 9.2 \text{ x } 10^3 \text{ M}^{-1}.\text{cm}^{-1}$) with a final yield of 3 mg/L of pure peptide. All the purification steps were performed in the presence of protease inhibitors cocktails (Roche, Basel, Switzerland and Halt, Thermo Scientific, Brebières, France) complemented with 0.1 mg/mL TPCK and TLCK.

CaM from bovine brain was purchased as a lyophilized powder (Sigma-Aldrich). ¹⁵N-¹³C-labeled CaM was expressed and purified according to Bouvier et al. (22).

Tubulin was purified from sheep brain using the method of Castoldi and Popov (23) and stored in liquid nitrogen. Before use, an additional cycle of polymerization/depolymerization was performed and tubulin was resuspended in 50 mM MES-KOH, 4.1 M glycerol, 1 mM MgCl₂, 0.5 mM EGTA, pH 6.9. Tubulin concentration was determined by spectrophotometry using an extinction coefficient $\varepsilon_{278nm} = 1.2 \times 10^5 \text{ M}^{-1} \text{ cm}^{-1}$.

Binding of MAP6-N fragments to MTs and MT stabilization Assays

For MT co-sedimentation assays, tubulin was polymerized for 30 min at 37°C in 40 mM MES-KOH, 4 mM MgCl₂, 1 mM DTT, 1 mM GTP, 2.7 M glycerol, pH 6.9 (buffer P) with stoichiometric amounts of taxol. 0, 5, 15 or 25 µM of taxol-stabilized MTs were then incubated with 20 µM MAP6₉₀₋₁₇₇ for 15 min at 20°C. MT pellets were collected by centrifugation (25,000 x g, 40 min at 20°C). To obtain MT binding saturation data, co-sedimentation experiments were performed using a range of concentrations of MAP690-177 (0-50 µM) and a fixed amount of taxol-stabilized MTs (10 µM). After 10 min incubation at 20°C, MT-bound MAP690-177 was obtained by centrifugation (25,000 x g, 40 min at 20°C) and estimated by SDS-PAGE using an Odyssey Imaging System (LI-COR Biosciences, Lincoln, NE, USA). The molar binding stoichiometry and dissociation constant (K_d) were determined graphically. The same procedure was used for co-sedimentation assays in the presence of Ca^{2+} -CaM (35 μ M taxolstabilized MTs, 10 µM MAP690-177, 0.2 mM CaCl₂, 0, 40 and 80 µM CaM, incubation for 5 min at 20°C). For MT cold stability assays, 25 µM tubulin was polymerized in buffer P for 30 min at 37°C without (control) or with 20 µM MAP6₉₀₋₁₇₇. After incubation on ice for 10 min, MT pellets and supernatants were separated by centrifugation (25,000 x g, 40 min, 4°C). To evaluate the influence of Ca^{2+} -CaM on the ability of MAP6₉₀₋₁₇₇ to stabilize MTs in cold condition, we used the same procedure in the presence of CaM (35 µM MT, 0.5 mM spermine and 50 µM CaCl₂ were first incubated for 10 min at 37°C with 25 µM MAP690-177 alone or in the presence of 100 uM CaM, followed by an incubation for 10 min on ice). The kinetics of MT polymerization were monitored turbidimetrically at 370 nm (1 cm path) in an Ultrospec 3000 spectrophotometer (GE Healthcare, Fairfield,

USA) equipped with a temperature controller. 35 µM tubulin was polymerized with 0, 0.35, 3.5 and 35 µM MAP6₉₀₋₁₇₇ in buffer P at 37°C. At the steady state, MT destabilization was induced by a cold shock $(4^{\circ}C)$ for 2 min. To verify the integrity of the tubulin after the cold treatment, the temperature was then set again at 37°C. The presence of MTs after the cold shock was examined by atomic force microscopy (AFM) on 10 µL aliquots containing 35 µM MAP6₉₀₋₁₇₇ and sampled at t = 30 min. The samples were deposited on freshly cleaved mica and dried for AFM imaging as described previously (24). All experiments were performed AFM in intermittent mode with a multimode AFM instrument (Digital Instruments, Veeco, Santa Barbara, USA) operating with a Nanoscope IIIa controller (Digital Instruments). Images were collected at a scan frequency of 1.5 Hz and a resolution of 512 x 512 pixels.

Immobilized CaM-binding assay

CaM-agarose beads (Sigma-Aldrich) were equilibrated in 50 mM MES-KOH, 1 mM CaCl₂, pH 6.9 and incubated with pure MAP6₉₀₋₁₇₇ for 30 min at 20°C. Beads were then washed with the same buffer. Flow-through, washes and CaM-beads fractions were analyzed on SDS-PAGE and the proteins revealed by Coomassie blue staining. The Ca²⁺-dependence of this interaction was tested with calcium-free buffer supplemented with 5 mM EGTA.

Isothermal titration calorimetry (ITC)

ITC measurements were performed at 25°C using an ITC₂₀₀ calorimeter (GE Healthcare, Fairfield, USA). Samples were thoroughly degassed before each titration. Titration and sample solutions were prepared in 20 mM MES-KOH, 50 mM NaCl, 1 mM DTT either with 1 mM CaCl₂ or 5 mM EGTA, pH 6.9. Titrations were carried out by injecting 25 consecutive aliquots (1.5 µL) of 200 µM MAP690-177 into the ITC cell (0.2 mL) containing 25 µM CaM. A background titration consisting of the identical titration solution but containing only the buffer in the cell was subtracted from each titration to quantify heat produced by sample dilution. The value obtained was subtracted from the heat of reaction to give the effective heat of binding. The resulting titration data were analyzed using the ORIGIN software (MicroCal Inc., Northampton, USA). The molar binding stoichiometry, binding

constant ($K_a = 1/K_d$) and binding enthalpy (ΔH) were determined by fitting the binding isotherm to a model with one set of sites. For the fit, no constraint in stoichiometry, K_a and ΔH was introduced. Changes in free energy (ΔG) and in entropy (T ΔS) were calculated from $\Delta G = -RT$ ln $K_a = \Delta H - T\Delta S$, where R is the gas constant and T the temperature in Kelvin.

NMR experiments

MAP6₉₀₋₁₇₇ and MAP6₉₀₋₁₄₂ NMR spectra were acquired at 20°C on a Bruker Avance 600 NMR spectrometer using a cryoprobe. All experiments were performed in 50 mM MES-KOH, 150 mM NaCl, 1mM DTT, pH 6.9 on 60 μ L samples using the MATCHTM system (Cortecnet, Paris, France). Three-dimensional NOESY- and TOCSY-¹H-¹⁵N HSQC and HNCO, HNCA, HN(CA)CO, CBCA(CO)NH experiments were performed for backbone assignments on a 500 μ M ¹⁵N- or ¹⁵N-¹³C-labeled MAP6₉₀₋₁₇₇ sample (25). All spectra were processed with Topspin 2.0 and NMRView (26).

 1 H- 15 N HSQC experiments were used to characterize the interaction of MAP6₉₀₋₁₇₇ and MAP6₉₀₋₁₄₂ with tubulin, taxol-stabilized MTs, apoCaM, Ca²⁺-CaM and BSA (negative control). Titration series were performed using 15 N-labeled MAP6₉₀₋₁₇₇ or MAP6₉₀₋₁₄₂ samples dissolved in 50 mM MES-KOH, 150 mM NaCl, 1 mM DTT, pH 6.9 at 6°C for tubulin and taxol-stabilized MTs and 20°C for CaM. Spectra were recorded using 128 x 2048 complex data points in F1 and F2 dimensions with 16 scans per increment. The spectral widths were 1202 and 1500 Hz in the 15 N and 1 H dimensions respectively.

A Ca²⁺-loaded uniformly ¹⁵N-¹³C-labeled CaM sample was prepared at 60 µM in 18 mM Tris-HCl, pH 7.8 and 10% D₂O and studied in Shigemi tube (Shigemi Inc., Allison Park, PA). The formation of the MAP6₉₀₋₁₇₇:Ca²⁺-CaM complex was examined on samples by direct addition of different aliquots of a concentrated solution of unlabeled MAP690-177 (from a 0.6 mM stock solution) to the ¹⁵N-¹³C-labeled Ca²⁺-CaM NMR sample to reach 1.5:1 MAP6₉₀₋₁₇₇:Ca²⁺-CaM stoichiometry. NMR spectra were acquired at 27 °C on an Agilent VNMRS 800 MHz spectrometer using a cryoprobe. BEST-TROSY experiment was used as it provides resolution without losing sensitivity compared to ¹H-¹⁵N HSQC (27). 128 x 930 complex points in the F1

and F2 dimensions were acquired for an experimental time of about 3 h. All data were processed using the NMRPipe software (28). Figures are represented with the VMD software (29).

RESULTS

MAP6₉₀₋₁₇₇ induces MT resistance to cold and nocodazole in HeLa cultured cells. As a prerequisite to biochemical and structural investigation, we verified the functionality of the MAP690-177 fragment toward MT stabilization in cultured HeLa cells. HeLa cells are well suited for this kind of experiments because they do not endogenously express MAP6 proteins and are devoid of cold-stable and drug-resistant MTs (30). HeLa cells were transfected with the pCMV-FLAG-MAP690-177 vector and examined for MT resistance to the cold and nocodazole. Results show that all the transfected cells exhibit a clearly visible MT network whereas nontransfected cells show a diffuse labeling. Additionally, analysis of the distribution of MAP6₉₀₋₁₇₇ in transfected cells shows that it colocalizes with the MT network suggesting that MT stabilization by MAP6₉₀₋₁₇₇ is direct (Fig. 2). As MAP6₉₀₋₁₇₇ thus binds to and stabilizes MTs in a cellular context, the investigation of its biochemical and structural properties appears of interest for the characterization of the full length MAP6 proteins.

MAP6₉₀₋₁₇₇ binds to MTs and induces MT resistance to cold in vitro. As a second step, to distinguish between a direct and indirect binding through putative protein partners, we analyzed the association of MAP690-177 with MTs in vitro using co-sedimentation experiments with taxolstabilized MTs. In the absence of MT, MAP6₉₀-177 was not detectable in the pellet demonstrating that MAP6₉₀₋₁₇₇ is soluble in these conditions. When MAP6₉₀₋₁₇₇ was incubated with increasing amounts of MTs, a concentration-dependent MAP6₉₀₋₁₇₇ binding to MTs was observed (Fig. 3A). This result shows that MAP6₉₀₋₁₇₇ binds directly to MTs. In addition, we found that shearing MTs, which increases the number of MT ends, did not significantly affect MAP6₉₀₋₁₇₇ binding to MTs (data not shown). To further characterize the MAP690-177:MT interaction, MAP6₉₀₋₁₇₇ at different concentrations was then

incubated with a fixed amount of taxol-stabilized MTs (Fig. 3B). MTs were then pelleted and the MAP6₉₀₋₁₇₇ content was analyzed by SDS-PAGE and quantified. The results show that the binding of MAP6₉₀₋₁₇₇ to MTs is saturable with an estimated K_d of 6 μ M and a 1:1 MAP6₉₀₋₁₇₇:tubulin heterodimer stoichiometry.

We then assessed the ability of the $MAP6_{90-177}$ fragment to stabilize MTs against cold. After polymerization of tubulin into MTs at 37°C, MTs were exposed to cold (ice, 10 min) in the absence or presence of MAP690-177. SDS-PAGE analysis shows that the amount of MTs in the pellet increases dramatically in the presence of MAP6₉₀₋₁₇₇ after cold treatment (Fig. 3C). Similar results were observed whether the MAP6₉₀₋₁₇₇ fragment was mixed to tubulin before polymerization or added to assembled MTs. To confirm that the MAP6₉₀₋₁₇₇ fragment truly protected MTs from cold depolymerization, we examined the dynamics of MT assembly by turbidimetry at 370 nm (Fig. 3D). Results show that the kinetics of tubulin polymerization are very similar in the absence or presence of different concentrations MAP6₉₀₋₁₇₇ of demonstrating that MAP690-177 has no impact on the nucleation and elongation steps. In addition, the steady state turbidity value was also similar in both cases suggesting that no particular supraassembly of MTs such as bundles were formed in the presence of MAP6₉₀₋₁₇₇. The cold shock was performed at steady state. In the control induced condition, it а clear MT depolymerization. The effect of this cold shock diminished dramatically in the presence of MAP6₉₀₋₁₇₇, and a total protection was observed when the MAP6₉₀₋₁₇₇:tubulin heterodimer ratio reached a value of about 1:1. To ascertain that the macromolecular objects diffusing light during the cold shock in the presence of MAP6₉₀. ₁₇₇ are really MTs, we performed AFM imaging which shows filamentous structures with length and width in good agreement with MT dimensions (31) (Fig. 3E).

Quantification of the MAP6₉₀₋₁₇₇: CaM interaction. It was previously demonstrated that MAP6-N binds to CaM (8) and that Mn modules are in part implicated in this interaction (19). To further investigate such interaction, we examined the binding of MAP6₉₀₋₁₇₇ to CaM immobilized on beads (Fig. 4A). The results show that MAP6₉₀₋₁₇₇ is able to bind to CaM but only in the

presence of $CaCl_2$. Indeed, when the binding experiment was realized with 5 mM EGTA instead of $CaCl_2$, no binding was observed demonstrating that calcium is necessary for this interaction.

We next carried out ITC experiments to quantify the MAP6₉₀₋₁₇₇:CaM interaction. Figure 4B displays the binding isotherm of MAP6₉₀₋₁₇₇ titration into the Ca²⁺-CaM solution. The best fit of the integrated isotherm indicated that one MAP6₉₀₋₁₇₇ binds to two Ca²⁺-CaM molecules with a K_d value of 0.4 μ M (Fig. 4C). The same ITC experiment carried out with EGTA in place of CaCl₂ showed no detectable interaction.

Identification of MAP6₉₀₋₁₇₇ residues implicated in the interaction with MTs and Ca^{2+} -CaM. Large amounts of pure and soluble MAP6₉₀₋₁₇₇ could be obtained from recombinant expression in E. coli which allowed NMR study of MAP690-177 interactions with its partners at the residue scale. ¹H-¹⁵N HSQC spectrum of MAP6₉₀₋₁₇₇ in aqueous buffer shows well resolved resonances with a weak dispersion of correlation peaks (less than 1 ppm and about 20 ppm on the ¹H and ¹⁵N dimensions respectively) suggesting that under these conditions the MAP690-177 does not fold into any defined tertiary structure (Fig. 5A). NMR assignment of the backbone resonances of MAP6₉₀₋₁₇₇ was achieved (Supp. files, Table 1). Analysis of the chemical shifts deviations from random coil values of ¹³CO and ¹³C α (32) also indicated that MAP690-177 does not contain significant secondary structure (Fig. 5B). In agreement with this, we observed that during the course of overexpression and purification, the MAP6₉₀₋₁₇₇ fragment exhibited a high sensitivity to protease degradation together with a high thermostability, properties that are frequently observed for intrinsically unstructured proteins.

In order to identify MAP6₉₀₋₁₇₇ residues that mediate MT and Ca²⁺-CaM interaction, ¹H-¹⁵N HSQC spectra of free ¹⁵N- MAP6₉₀₋₁₇₇ and bound to unlabeled tubulin, MTs and Ca²⁺-CaM were recorded (Fig. 6A). Because of the large size of these complexes, many ¹H-¹⁵N cross-peaks of residues implicated in the interaction underwent line broadening and disappeared which helped us to point them out. Experiments carried out either with tubulin or MTs gave identical results whereas control experiments performed with BSA showed no modification. A peak by peak analysis using assignment of MAP6₉₀₋₁₇₇ allowed us to define two MT binding sites delimited by

residues 121-138 and 150-175 (Fig. 6A). As the C-terminal tail of tubulin is a major binding site for many MT partners, we tested whether this part of tubulin is implicated in the MAP6₉₀₋₁₇₇:MT interaction. Co-sedimentation assays were carried out in the presence of an excess of a recombinant tubulin fragment corresponding to the C-terminal tail of α -tubulin (α Tub410C) (33) (Supp. files, Fig. S1). Results show that this α Tub410C fragment did not compete with MTs for MAP6₉₀₋₁₇₇ binding. This result was confirmed by NMR which showed that the ¹H-¹⁵N HSQC spectrum of ¹⁵N-MAP6₉₀₋₁₇₇ remained identical in the absence or in the presence of an excess of an excess of unlabeled α Tub410C (data not shown).

We then attempted to locate the Ca^{2+} -CaM binding site on MAP6₉₀₋₁₇₇ using similar ¹H-¹⁵N HSQC mapping experiments (Fig. 6A). As seen for MTs, Ca^{2+} -CaM binding to ¹⁵N-MAP6₉₀₋₁₇₇ induced line broadening due to the large size of the complex. The mapping of the Ca²⁺-CaM binding site points out two regions delimited by residues 121-137 and 151-176 which are very similar to those that participate to the binding of MTs. Surprisingly, this suggests a common binding mode for MTs and Ca²⁺-CaM. Finally, the binding of MAP6₉₀₋₁₇₇ to Ca²⁺-CaM could be reversed with EGTA confirming again the Ca²⁺-dependence of the MAP6₉₀₋₁₇₇:CaM interaction (data not shown).

We then examined by competition experiments whether Ca²⁺-CaM could impair MAP6₉₀₋₁₇₇ binding to MTs (Fig. 6B). Taxol-stabilized MTs were incubated in the presence of a fixed amount of MAP690-177 and increasing concentrations of Ca²⁺-CaM. Free and MT-bound MAP6₉₀₋₁₇₇ were then separated by centrifugation and analyzed by SDS-PAGE and Coomassie blue staining. Results indicate a progressive displacement of MAP690-177 from pellet to supernatant as the concentration of Ca^{2+} -CaM increases. Hence, it appears that MTs and Ca^{2+} -CaM bind to MAP6₉₀₋₁₇₇ in a mutually exclusive manner. This competition was confirmed in functional assays in which the protective effect of MAP690-177 for MT cold depolymerization was tested in the presence of Ca²⁺-CaM (Fig. 6C). In this experiment, Ca²⁺ was required to obtain the holoform of CaM. However, due to the destabilizing effect of Ca²⁺ on MTs, experiments were performed in the presence of excess of spermine which acts as a Ca^{2+} competitive inhibitor for

MT binding (33). Polymerized MTs were incubated on ice alone (control, the two left lanes) or with MAP6₉₀₋₁₇₇ (the two middle lanes) or with MAP6₉₀₋₁₇₇ and Ca²⁺-CaM (the two right lanes). Results show clearly that MAP6₉₀₋₁₇₇ no longer protects MTs in the presence of Ca²⁺-CaM.

Binding of $MAP6_{90-177}$ to Ca^{2+} -CaM induces large conformational changes of the Ca^{2+} -CaM structure. In order to study the structural impact of MAP6₉₀₋₁₇₇ binding on the Ca²⁺-CaM structure, we carried out NMR experiments with purified and ¹⁵N-¹³C-labeled Ca²⁺-CaM. BEST-TROSY spectra of ¹⁵N-¹³C-Ca²⁺-CaM in the absence and presence of unlabeled MAP690-177 were recorded (Fig. 7A). The addition of MAP6₉₀₋₁₇₇ to Ca²⁺-CaM resulted in very large changes of the spectrum profile of Ca²⁺-CaM with modifications of the peaks number and of the chemical shifts distribution. Residues affected by this interaction are reported on the CaM sequence and on a crystallographic structure of Ca^{2+} -CaM (34) (Fig. 7B and 7C). Results show that the perturbations are found mostly in the N-terminal domain and in the central α helix of Ca²⁺-CaM. Indeed, NMR mapping points out a first stretches in the Nterminal domain (residues 13-39) and a second long stretches comprising the end of the Nterminal domain and more than half the central α helix (residues 49-80). The C-terminal domain is affected but in a lesser extent. Together, these observations indicate that the binding of MAP6₉₀₋₁₇₇ induces wide-range conformational and dynamical changes in the Ca2+-CaM structure implicating mostly its N-terminal domain and the central α -helix.

MTs and Ca^{2+} -CaM compete for the same binding site on the Mn1 module. We finally examined the possibility that a single Mn module could interact with MTs and Ca^{2+} -CaM. To that end, we overexpressed and purified a 53 amino acid residues fragment encompassing the Nterminal part and the Mn1 module of MAP6₉₀₋₁₇₇ (MAP6-N residues 90-142, named herein MAP6₉₀₋₁₄₂). We observed that the ¹H-¹⁵N HSQC spectrum of MAP6₉₀₋₁₄₂ superimposes very well with that of the MAP6₉₀₋₁₇₇ fragment which strongly facilitated its assignment. When the MAP6₉₀₋₁₄₂ fragment was studied in the presence of tubulin or MTs, we observed on its ¹H-¹⁵N HSQC spectrum (Fig. 8A) line broadening responsible for the disappearance of some peaks corresponding to the stretch of residues 121-138 with the exception of C135 (Fig. 8B). Interestingly, the ${}^{1}\text{H}{-}^{15}\text{N}$ HSQC spectrum of MAP6₉₀₋₁₄₂ acquired in the presence of Ca²⁺-CaM (Fig. 8A) gave very close results with line broadening and disappearance of peaks corresponding to residue V118 and of the stretch 121-138 except Q129 and S134 (Fig. 8B). Hence, these results show that the Mn1 module, when considered isolated from the other Mn modules, is able to bind to both MTs and Ca²⁺-CaM in a similar manner.

DISCUSSION

MAP690-177 binds and stabilizes MTs - The aim of this work was to better understand the molecular mechanisms underlying the stabilization of MTs by MAP6 and its regulation by CaM using biochemical and structural investigations. To that end, we took advantage of a MAP6-N fragment encompassing its Mn1 and Mn2 modules which recapitulates the functional properties of the full length protein. As a first step, we studied the interaction between MAP6₉₀₋₁₇₇ and MTs. Using co-sedimentation assays, we found that MAP690-177 interacts with MTs with an apparent K_d value of about 6 µM and a 1:1 MAP690-177:tubulin heterodimer stoichiometry (Fig. 3B). To the best of our knowledge, this is the first report on a K_d value regarding the interaction of Mn modules with MTs. Delphin et al. reported a K_d value of about 70 nM for the interaction between the Mc domain of MAP6-F and MTs (18). The lower apparent affinity observed here for MAP690-177 may be related to the fact that this fragment contains only two Mn modules whereas the Mc domain contains five Mc modules. This suggests that the binding of MAP6 to MTs may be cooperative. Co-sedimentation assays (Supp. files, Fig. S1) and NMR experiments (data not shown) using a α-tubulin C-terminal fragment (α Tub410C, residues 410–451 from α 1a-tubulin (33)) indicated that this part of tubulin is not sufficient for the interaction with MAP6₉₀₋₁₇₇. This agrees with previous report showing that, unlike other MAPs, MAP6 can bind significantly to any tubulin isoform whatever its degree of Cterminal modification by polyglutamylation (35).

We observed that the stabilization of MTs against cold and nocodazole by MAP6₉₀₋₁₇₇ does

not induce MT aggregates nor bundles in contrast to what was observed with full length MAP6-N by immunofluorescence microscopy (7). In this later case, the formation of MT bundles may be due either to the presence of multiple Mn and Mc repeats or to a possible multimerization of the protein via stretches of amino acid residues that are located outside the MAP6₉₀₋₁₇₇ region. We next mapped the MAP6₉₀₋ 177 interaction with MTs to two MAP690-177 binding sites defined by residues 121-138 and 150-175 which correlate strongly with Mn1 and Mn2 modules respectively (19). Interestingly, NMR data show that a single Mn module (MAP6₉₀₋₁₄₂) can still interact with MTs suggesting that each Mn module in MAP6 may serve as anchor point for MT binding. Accordingly, each Mn repeat may recognize a conserved binding motif located on α - and β tubulin. From a structural point of view, MT stabilization by MAP6₉₀₋₁₇₇ is thus likely to occur through the reinforcement of the longitudinal contacts between tubulin heterodimers within a protofilament as seen for Tau (36) or lateral contacts between adjacent tubulin heterodimers preventing protofilaments separation like the stabilization mode of doublecortin (37). In addition, we cannot exclude that MAP6₉₀₋₁₇₇ may also stabilize MTs by preventing cold- or nocodazole-induced conformational changes in tubulin by a more complex mechanism.

Association of MAP690-177 with MTs is regulated by $Ca^{2+}-CaM$ - Biochemical and ITC experiments showed that MAP690-177 binds to CaM in a Ca²⁺-dependent manner with a K_d value of 0.4 µM and a 1:2 MAP6₉₀₋₁₇₇:Ca²⁺-CaM stoichiometry (Fig. 4). This K_d value lies in the range of that previously reported for MAP6- $F{:}Ca^{2+}{-}CaM$ (K_d ~ 1.7 and 8.1 $\mu M)$ (19) and Mc:Ca²⁺-CaM (K_d ~ 1.5 and 3.9 μ M) interactions (22,38). Such an affinity is compatible with a physiological interaction of Ca²⁺-CaM with MAP6 for instance after a transient rise in cytoplasmic $[Ca^{2+}]$. Investigation of MAP6₉₀₋ 177:Ca²⁺-CaM interaction by NMR pointed out two MAP6₉₀₋₁₇₇ regions (121-137 and 151-176) which are nearly identical to the MAP690-177:MTs binding sites. This suggests that Ca²⁺-CaM may compete with MTs for MAP6₉₀₋₁₇₇ binding. In agreement with this hypothesis, we demonstrated for the first time using co-sedimentation assays that Ca²⁺-CaM is responsible for a stoichiometric inhibition of MAP690-177 binding to MTs

resulting in a loss of MT stabilization. Altogether, these observations suggest that the regulation of MAP6 binding to MTs by Ca²⁺-CaM is a dynamic process rapidly modulated by the Ca²⁺-charge state of CaM. A similar mechanism was proposed for Tau in the flip-flop model (39). Finally, this part of MAP6-N also comprises S139 which can be phosphorylated by the Ca²⁺-Calmodulin-dependent protein kinase II (CaMKII) which triggers the translocation of MAP6 from MTs to actin (40). Interestingly, this kind of regulation has already been observed for the microtubule-associated protein p35 which is also regulated by site-specific phosphorylations and Ca²⁺-CaM with MT overlapping binding sites (41). Similar mechanisms of CaM regulation were again suggested for Tau (42-44).

Impact of MAP6₉₀₋₁₇₇ on Ca^{2+} -CaM structure, implication for the binding mode - Previous NMR, infrared spectroscopy and differential scanning calorimetry investigations on the interaction of Ca²⁺-CaM with a model peptide corresponding to a single Mc motif revealed an unusual binding mode. Indeed, these studies showed that the Mc peptide interacts with the Cterminal domain of CaM leaving it in an extended conformation with no significant structural change (22)(38). The present data provide strong evidences that the binding mode of MAP690-177 to Ca²⁺-CaM is distinct from that previously observed with Mc modules. However, the present NMR data show that the binding of MAP 6_{90-177} to Ca²⁺-CaM produces large structural and dynamical changes in Ca²⁺-CaM. Interestingly, sequence analysis of the two sites of MAP6₉₀₋₁₇₇ that interact with Ca²⁺-CaM, suggests that they belong to the 1-5-10 class of canonical CaM-binding motifs found in several CaM-regulated proteins such as CaMKII and Synapsin I (45,46) (Fig. 9). This class refers to a group of Ca²⁺-CaM partners whose key hydrophobic bulky residues are spaced by eight

residues with an additional anchoring residue in the middle. Upon binding, both Ca²⁺-CaM and target peptide motif undergo large the conformational rearrangements. The canonical CaM-binding motif adopts а α -helical conformation and the two domains of CaM wraps around it, enclosing it in a hydrophobic channel within the globular core. The data MAP6₉₀₋₁₇₇:Ca²⁺-CaM obtained here on interaction argue for a similar binding mode: i) Ca²⁺ is required for MAP6₉₀₋₁₇₇ binding, a fact that is systematically observed with this type of binding mode; ii) large structural modifications of the Ca²⁺-CaM structure are observed affecting its two domains and its central flexible helix and iii) secondary structure prediction of MAP6₉₀₋₁₇₇ show that residues 117-125 and 148-158 may fold into α -helices. These stretches overlap the 1-5-10 motifs of MAP690-177 and it is possible that they locally fold when in complex with Ca^{2+} -CaM. It is worthy to note that neither the Mc modules nor the Mn3 module of MAP6 (which bind to $Ca^{2+}-CaM$) contain such 1-5-10 motif. This may explain previous reports showing an unusual binding mode for these CaM-binding sites (22) (Pierre Gans, personal communication). Interestingly, some 1-5-10 motifs were shown to be implicated both in Ca^{2+} -CaM and actin binding (47-49) and MAP6-N was shown to interact with actin (40). It would be of interest to probe whether this region is implicated in MAP6-N association with actin and whether Ca²⁺-CaM binding to this region may regulate such process.

In conclusion we report here for the first time structure and functional information on a short region from MAP6 that recapitulates its ability to bind and stabilize MTs upon exposure to cold or nocodazole. The binding of this region to MTs appears modulated by a competitive interaction with Ca²⁺-CaM which further documents the complex regulation of MT assembly by MAP6.

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Acknowledgments- We thank Rabia Letaief for her help for NMR resonances assignment and Imane Hamdi for cell culture experiments. We thank Sylviane Hoos and Patrick England from la Plate-forme de biophysique des macromolécules et de leurs interactions (Institut Pasteur, Paris, France) for help, technical support and fruitful discussions.

FOOTNOTES

*Recipient of a postdoctoral fellowship from the PRES UniversSud

The abbreviations used are: AFM, atomic force microscopy; BSA, bovine serum albumin; CaM, calmodulin; CaMKII, Ca²⁺-Calmodulin-dependent protein kinase II; DMEM, Dulbecco's modified Eagle medium; DTT, dithiothreitol; EGTA, ethylene glycol tetraacetic acid; GTP, guanosine 5'-triphosphate; MAP, microtubules-associated protein; ITC, isothermal titration calorimetry; NMR, nuclear magnetic resonance; MT(s), microtubule(s); SDS-PAGE, sodium dodecyl sulfate polyacrylamide gel electrophoresis; STOP, stable tubule only polypeptide

FIGURE LEGENDS

FIGURE 1. Mn modules from MAP6-N are conserved across species. A. Schematic representation of the MAP6-N protein (Rattus norvegicus). B. Sequence alignment of MAP6-N Mn1 and Mn2 modules from different species obtained with the ClustalW2 software (50). Conservation is based on the AMAS method of multiple sequence alignment analysis and reflects the conservation of physico-chemical properties of amino acids (51). Accession numbers are: Rn *NP_058900* (NCBI), Mum *NP_001041632* (NCBI), Hs *AAI50255* (GenBank), Mam *AFJ70456* (GenBank), Dc *XP_534007* (NCBI), Ss *NP_001167398* (NCBI), Tn *CAF90592* (GenBank), Gg *NP_990250* (NCBI), Xt *NP_001120222* (NCBI). C. Sequence alignment of the three Mn modules of MAP6-N (Rattus norvegicus).

FIGURE 2. MAP6₉₀₋₁₇₇ induces MT resistance to cold and nocodazole in HeLa cells. HeLa cells were transfected with the pCMV-FLAG- MAP6₉₀₋₁₇₇ plasmid. 48 h after transfection, cells were either exposed to ice or to 20 μ M nocadazole for 30 min, fixed/permeabilized and then stained with anti-tubulin and anti-FLAG antibodies (bar = 5 μ m).

FIGURE 3. MAP6₉₀₋₁₇₇ binds in vitro to MTs and induces MT resistance to the cold. A. SDS-PAGE analysis of MT co-sedimentation assay. Increasing concentrations of taxol-stabilized MTs were incubated with a fixed amount of MAP6₉₀₋₁₇₇ (20 μ M) and centrifuged. The pellet fractions containing MTs and co-sedimented MAP6₉₀₋₁₇₇ were run on SDS-PAGE and revealed by Coomassie blue staining. B. Saturation curve of MAP6₉₀₋₁₇₇ fixation to MTs. MAP6₉₀₋₁₇₇ at different concentration was incubated with a fixed amount of taxol-stabilized MTs (10 μ M). MT-bound MAP6₉₀₋₁₇₇ was quantified by SDS-PAGE using an Odyssey Imaging System. Error bars are from three independent experiments. C. SDS-PAGE analysis of MT stabilization assay. 25 μ M tubulin was polymerized at 37°C for 30 min without or with 20 μ M MAP6₉₀₋₁₇₇. After incubation on ice (or 37°C for the control) for 10 min, MT pellets were collected by centrifugation at 4°C (or 37°C for the control) and analyzed by SDS-PAGE and Coomassie blue staining. D. MT formation was assessed by turbidimetry. 35 μ M tubulin was polymerized with varying concentrations of MAP6₉₀₋₁₇₇ at 37°C: 0 (full line), 0.35 (thick dashed line), 3.5 (thin dashed line) and 35 μ M (dot line). MT destabilization was induced by a cold shock at t = 23 min (2 min at 4°C) and temperature was then set again at 37°C. E. AFM imaging. Image was recorded at t = 30 min in the presence of 35 μ M MAP6₉₀₋₁₇₇ (bar = 1 μ m).

FIGURE 4. MAP6₉₀₋₁₇₇ interacts with Ca²⁺-CaM (K_d = 0.4 μ M). A. MAP6₉₀₋₁₇₇ binds only to the holoform of CaM. MAP6₉₀₋₁₇₇ was incubated with CaM-agarose beads in the presence or absence of Ca²⁺. Beads were then washed and flow-through, washes and CaM-beads fractions were analyzed on SDS-PAGE and revealed by Coomassie blue staining. B. ITC analysis of the interaction of MAP6₉₀₋₁₇₇ with Ca²⁺-CaM. Heat signal is shown (top) with the binding isotherm derived from this signal (bottom). C. Parameters obtained from the best fit.

FIGURE 5. Structural analysis of MAP6₉₀₋₁₇₇. A. Assigned MAP6₉₀₋₁₇₇ ¹H-¹⁵N HSQC spectrum in 50 mM MES-KOH, 150 mM NaCl, 1mM DTT, pH 6.9 at 20°C. Some indeterminations remained for

residues marked with triangles (E151 or R166), stars (K156) and rectangles (Q142, E149 or Q155). B. Corresponding secondary chemical shifts for ¹³CO and ¹³C α illustrating the absence of secondary structure.

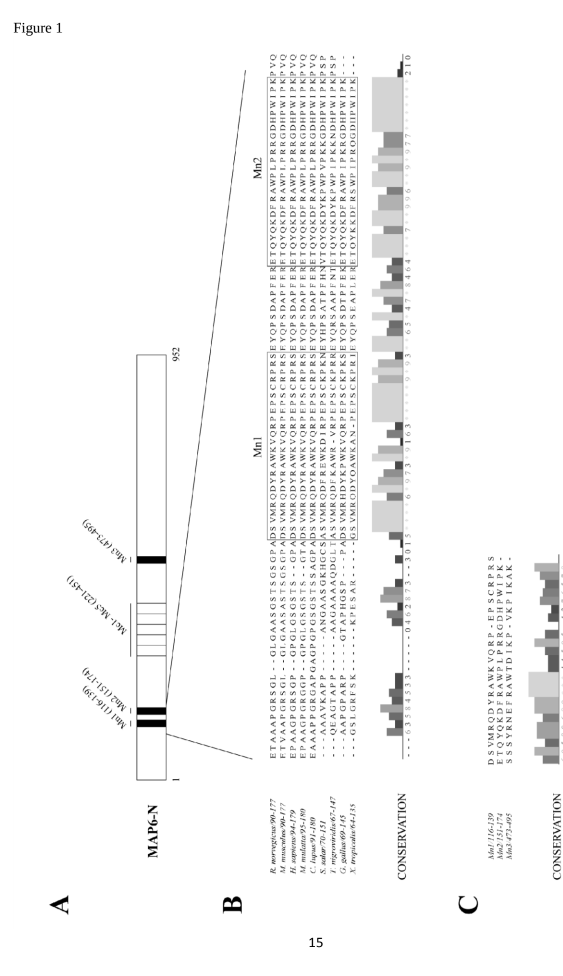
FIGURE 6. MTs and Ca²⁺-CaM compete for the same binding site on MAP6₉₀₋₁₇₇. A. NMR analysis of MAP6₉₀₋₁₇₇:MT and MAP6₉₀₋₁₇₇:Ca²⁺-CaM interactions. ¹H-¹⁵N HSQC spectra of ¹⁵N-labeled MAP6₉₀₋₁₇₇:MT (molar ratio 1:2, 6°C) and ¹⁵N-labeled MAP6₉₀₋₁₇₇:Ca²⁺-CaM (molar ratio 1:3, 20°C) complexes. The MAP6₉₀₋₁₇₇ residues implicated in the binding of MTs and Ca²⁺-CaM are highlighted in bold (proline residues lying in or around the sequences were arbitrarily included). B. Ca²⁺-CaM competes with MTs for MAP6₉₀₋₁₇₇ binding. 35 μ M taxol-stabilized MTs were incubated with 10 μ M MAP6₉₀₋₁₇₇, 0.2 mM CaCl₂ and varying concentrations of CaM for 5 min at 20°C. After centrifugation, pellet and supernatant contents were analyzed by SDS-PAGE and Coomassie blue staining. Stars correspond to experiments where 0.2 mM CaCl₂ was replaced by 0.5 mM EGTA. C. Ca²⁺-CaM inhibits the protective effect of MAP6₉₀₋₁₇₇ on MTs. 35 μ M tubulin was polymerized in the presence of 0.5 mM spermine to counteract the negative effects of Ca²⁺ on MTs. After addition of 50 μ M CaCl₂, polymerized MTs were incubated on ice for 10 min alone (control, the two left lanes) or with 25 μ M MAP6₉₀₋₁₇₇ (the two middle lanes) and 100 μ M CaM (the two right lanes). Pellets and supernatants contents were analyzed by SDS-PAGE and coomassie blue staining.

FIGURE 7. Ca²⁺-CaM undergoes large conformational changes when in interaction with MAP6₉₀₋₁₇₇. A. BEST-TROSY spectra superposition of ¹⁵N-¹³C-labeled Ca²⁺-CaM in the absence (red) and presence of MAP6₉₀₋₁₇₇ (0.5:1 MAP6₉₀₋₁₇₇:Ca²⁺-CaM molar ratio) (gray). Residues that are not significantly affected by the interaction (no line broadening nor significant chemical shift perturbation $(((\Delta\delta^1H)^2 + (\Delta\delta^{15}N/6.5)^2)^{1/2} < 0.09 \text{ ppm}, (52))$ are labeled. B. Sequence of CaM showing the residues affected by the interaction (bold). The proline P67 residue was arbitrarily included. The central α helix separating the N- and C-terminal domains of CaM is underlined. C. Spatial representations of the Ca²⁺-CaM structure (PDB code: 1OSA). Figure on the right was obtained after a 180° rotation around the x axis and a 90° rotation around the z axis. Residues which are not significantly affected by the interaction with z axis. Yellow balls: Ca²⁺ ions.

FIGURE 8. The Mn1 module of MAP6-N (residues 90-142, MAP6₉₀₋₁₄₂) interacts with MTs and Ca²⁺-CaM. A. NMR analysis of MAP6₉₀₋₁₄₂:MT and MAP6₉₀₋₁₄₂:Ca²⁺-CaM interactions. ¹H-¹⁵N HSQC spectra of ¹⁵N-labeled MAP6₉₀₋₁₄₂:MT and ¹⁵N-labeled MAP6₉₀₋₁₄₂:Ca²⁺-CaM complexes (1:1 molar ratio at 6°C and 1:3 molar ratio at 20°C respectively). B. Residues implicated in the binding of MTs and Ca²⁺-CaM are highlighted in bold in the MAP6₉₀₋₁₄₂ sequence (proline residues lying in the sequences were arbitrarily included).

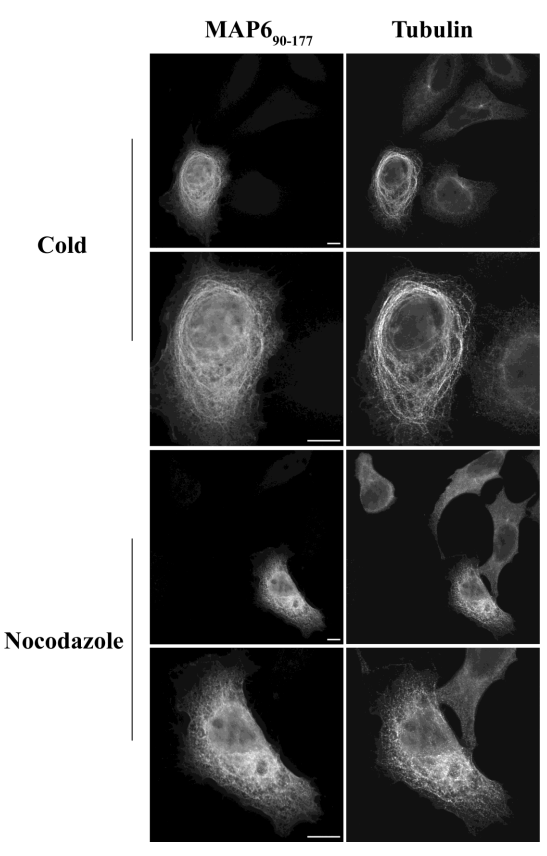
FIGURE 9. CaM 1-5-10 binding motif alignment. Sequence alignment of 1-5-10 class motifs from selected CaM-binding partners. The Ca²⁺-CaM interacting region delimited by NMR MAP6₉₀₋₁₇₇ comprises two canonical CaM binding motifs belonging to the 1-5-10 class.

FIGURES

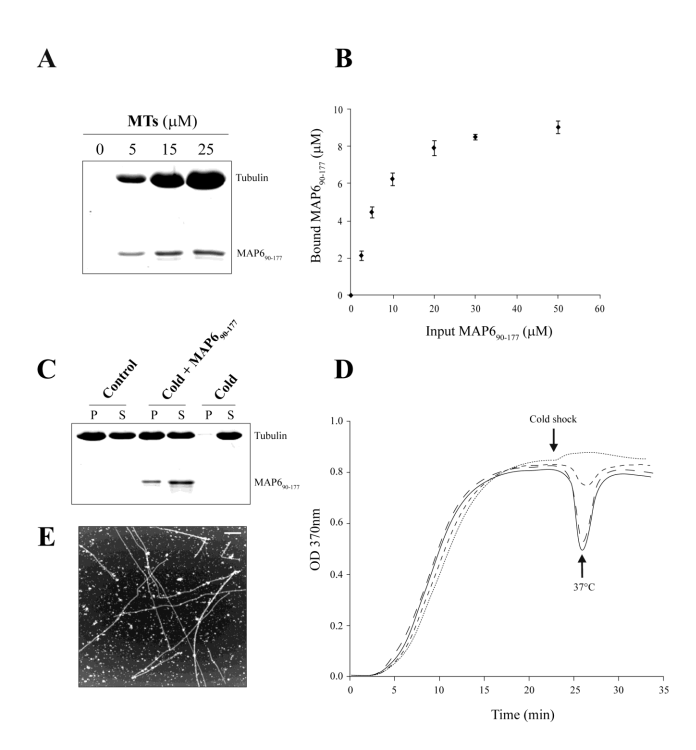


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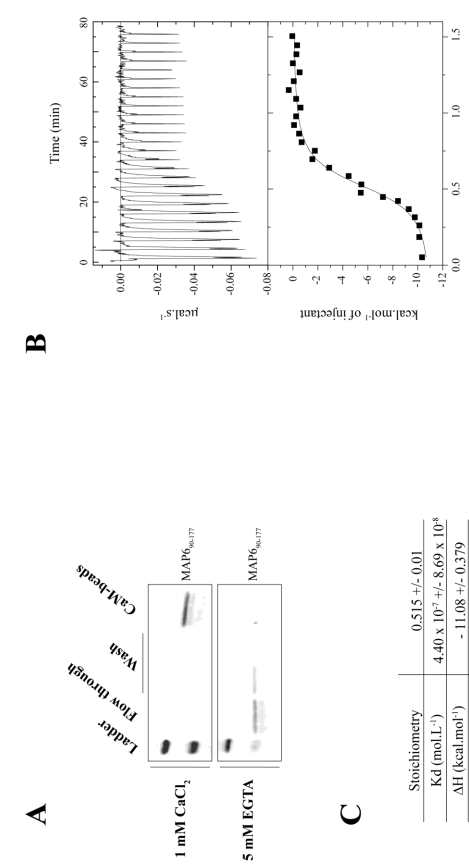


Molar ratio

- 8.71

 ΔS (cal.mol⁻¹.K⁻¹)

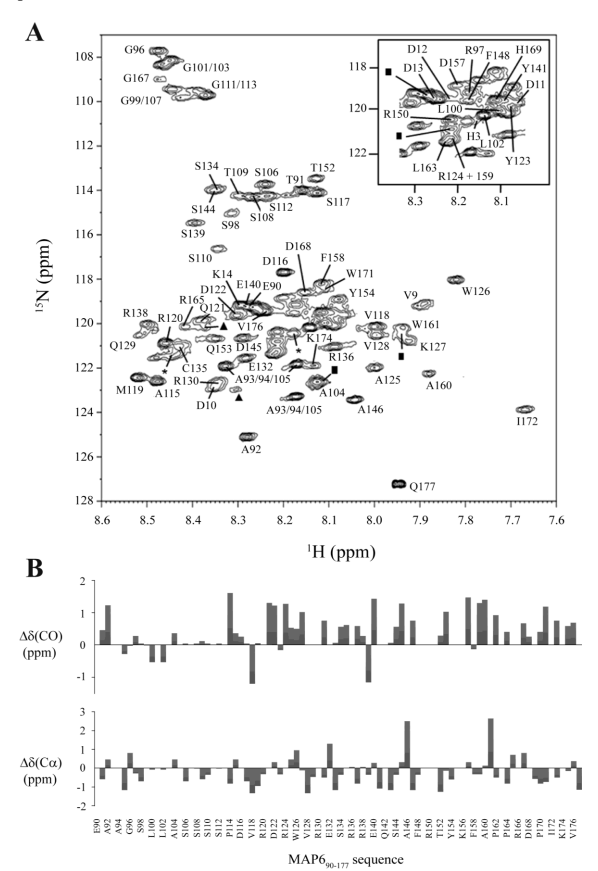


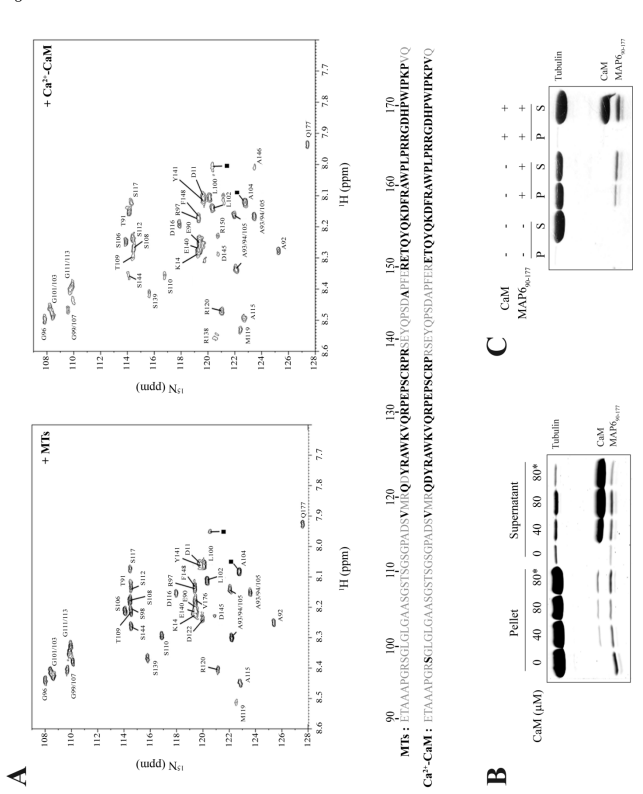




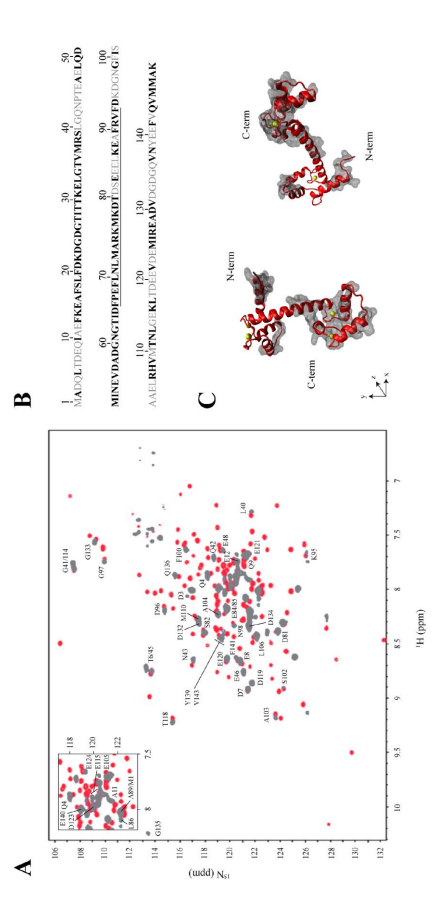
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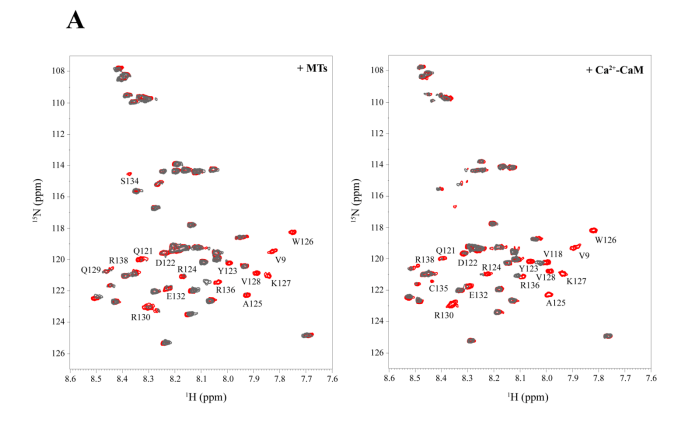












B

90 100 110 120 130 140 MTs : ETAAAPGRSGLGLGAASGSTSGSGPADSVMRQDYRAWKVQRPEPSCRPRSEYQ Ca²⁺-CaM : ETAAAPGRSGLGLGAASGSTSGSGPADSVMRQDYRAWKVQRPEPSCRPRSEYQ

Figure 9

	1	5	10
	ᡟ	ᡟ	♥
1-5-10 motif	[Φ] -	[Φ] -	0
<i>CaMKII/101-115</i>			TTMLAT
ΡΙ3Κγ/679-693	RFLLK	RGLR	NKRIGH
Cathelicidin-4/129-143	S V I L P	WK W P	WWP W RR
Synapsin I/123-137	H T D'W'A	K Y F K	G K K I H G
MAP6-N/116-130	D S VMR	QDYR	AWK VQ R
MAP6-N/151-165	E T Q Y	K D <u>F</u> R	AWPLPR