CLASP1 and CLASP2 bind to EB1 and regulate microtubule plus-end dynamics at the cell cortex

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 belongs to the Glued, adenomatous polyposis coli (APC), and
CLIP-associating protein (CLASP) 1 and CLASP2
are mammalian microtubule (MT) plus-end binding proteins, which associate with CLIP-170 and CLIP-115. Using RNA interference in HeLa cells, we show that the two CLASPs play redundant roles in regulating the density, length distribution and stability of interphase MTs. In HeLa cells, both CLASPs concentrate on the distal MT ends in a narrow region at the cell margin. CLASPs stabilize MTs by promoting pauses and restricting MT growth and shortening episodes to this peripheral cell region. We demonstrate that the middle part of CLASPs binds directly to EB1 and to MTs. Furthermore, we show that the association of CLASP2 with the cell cortex is MT independent and relies on its COOH-terminal domain. Both EB1- and cortex-binding domains of CLASP are required to promote MT stability. We propose that CLASPs can mediate interactions between MT plus ends and the cell cortex and act as local rescue factors, possibly through forming a complex with EB1 at MT tips.

Introduction

Microtubules (MTs) are polar filaments, which can undergo alternating phases of growth and shortening, a behavior termed dynamic instability (Desai and Mitchison, 1997). In living cells, a large number of protein factors regulate dynamic instability, thus determining the shape of the MT network in different phases of the cell cycle and in different cell regions. An interesting group of MT-associated factors, involved in regulation of MT dynamics, binds specifically to the ends of growing MTs. These factors, named +TIPs (plus-end tracking proteins), include members of structurally unrelated protein families, such as end binding proteins EB1, EB2 (RP1), and EB3, cytoplasmic linker proteins CLIP-170 and CLIP-115, dynactin large subunit p150Gluad, adenomatous polyposis coli (APC), and CLIP-associating proteins (CLASPs; for reviews see Schuyler and Pellman, 2001; Carvalho et al., 2003; Howard and Hyman, 2003; Mimori-Kiyosue and Tsukita, 2003).

CLASPs are a family of MT-associated proteins, conserved in animals and fungi (Inoue et al., 2000; Lemos et al., 2000). A distant member of this family, the budding yeast Stu1p, was isolated as a suppressor of a cold-sensitive tubulin mutation and an essential component of the mitotic spindle (Pasqualone and Huffaker, 1994; Yin et al., 2002). The first member of this family from higher organisms to be characterized in detail, the Drosophila Orbit/MAST, was also shown to be essential for mitosis (Inoue et al., 2000; Lemos et al., 2000), and the same holds true for one of the three C. elegans homologues, cls-2 (R106.7; Gonczy et al., 2000). Mammalian homologues, CLASP1 and 2, were initially characterized through their ability to bind to CLIP-170 and CLIP-115 (Akhmanova et al., 2001). The interaction between the CLASP and CLIP counterparts was also observed in flies (Mathe et al., 2003).

CLASP family members participate in generating polarized MT networks: in migrating fibroblasts, CLASP2 is involved in stabilizing MTs specifically at the leading edge (Akhmanova et al., 2001), whereas during fly oogenesis, Orbit/MAST organizes MTs, which interconnect the oocyte and the nurse cells (Mathe et al., 2003). During mitosis, Orbit/MAST plays a role in maintaining the bipolarity of the mitotic spindle, in the MT-

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kinetochore attachment and chromosome congression (Maiato et al., 2002). Similar functions were ascribed to the human CLASP1 (Maiato et al., 2003a,b). In addition, Orbit/MAST is needed for cytokinesis, because it stabilizes the interior MTs of the central spindle, necessary for the progression of the cleavage furrow (Inoue et al., 2004). Although the MT-stabilizing role of CLASP homologues has been established, the effect of CLASPs on the parameters of MT dynamic instability has not been analyzed, and the functional redundancy of the two mammalian CLASPs has not been addressed.

In this study, we show that CLASP1 and 2 have similar and redundant roles in organizing the interphase MTs in HeLa cells. CLASPs stabilize MTs, by retaining their plus ends in the peripheral cortical region, where they are pausing or undergo short polymerization–depolymerization excursions. To get insight into the mechanism of CLASP action, we have analyzed the targeting domains of the CLASP proteins. We show that a short repetitive region in the middle part of CLASP1 and 2 can bind to EB1 and EB3 and recognize growing MT tips, whereas the COOH-terminal domain of CLASP2 associates with the Golgi apparatus and cell cortex. We propose that the MT-stabilizing activity of CLASPs depends on the interaction with the EB proteins.

**Results**

**Characterization of siRNAs, specific for CLASP1 and CLASP2**

To detect CLASP1 protein, we have raised two new antibodies against CLASP1, #402 and #2292, which strongly react with GFP-CLASP1 and #2358 (CLASP2) and either α-tubulin or the Golgi marker GM130. Bars, 10 μm. (G) HeLa cells, transfected either with the control siRNA or the CLASP1+2B siRNAs, were stained with antibodies #402 or #2358 72 h after transfection. Bars, 10 μm.
were reduced by ~70%, which can be regarded as a hypomorphic state (Fig. 1 B). Control siRNAs had no effect on CLASP levels (Fig. 1 B and not depicted).

The partial down-regulation of CLASPs, separately or together, had no effect on the levels of tubulin or other +TIPs (Fig. 1, C and D). However, when CLASP1 and CLASP2 were knocked down simultaneously, the amount of acetylated tubulin (a marker of stable MTs; Bulinski and Gundersen, 1991) was diminished by ~40%, suggesting that MT stability was reduced (Fig. 1 D), confirming our previous findings in 3T3 fibroblasts (Akmanova et al., 2001).

CLASP1- and CLASP2-directed antibodies #402 and #2358 produced similar staining patterns in interphase HeLa cells: in line with previous observations (Akmanova et al., 2001; Maiato et al., 2003a), they decorated the Golgi apparatus, the centrosome, and MT plus ends (Fig. 1 F and not depicted). The MT tip staining, which nicely colocalized with other +TIPs (EB1, CLIP-170, and p150

Organization of the MT network after CLASP knockdown
Simultaneous knockdown of the two CLASPs caused mitotic defects, which were similar to those observed in the Orbit/MAST mutants (Maiato et al., 2002; Inoue et al., 2004) and will be described elsewhere (unpublished data). In interphase cells, a significant decrease in density of the MT network was observed (Fig. 2, A and B). This effect was particularly obvious in cells, which had no contacts with their neighbors and, therefore, we confined our analysis to such isolated cells.

To get a measure of MT density, we have determined in-cell staining with anti-CLASP antibodies, whereas the specific signals at the MT tips, Golgi, and the centrosome were greatly diminished (Fig. 1 G).

To confirm that the observed effects on the MT network result from the knockdown of CLASP proteins, rather than unspecific effects of the used RNA duplexes, we transfected cells, treated with CLASP1+2 siRNA combinations, with rescue constructs, encoding GFP or monomeric RFP (mRFP) fusions of CLASP1α or CLASP2α, resistant to the siRNAs due to silent substitutions in the siRNA target regions. Both CLASP1α and 2α fusions could restore MT density and the alignment of MT tips along the cell cortex (Fig. 3 A; Video 2, available at http://www.jcb.org/cgi/content/full/jcb.200405094/DC1). After the rescue with mRFP-CLASP2α the number of MT ends in a sector increased from 14.0 ± 4.3 in surrounding CLASP knockdown cells to 27.9 ± 2.6 in cells, expressing the rescue con-
Figure 2. Effect of CLASP knockdown on the interphase MT network. (A and B) HeLa cells 72 h after transfection with the control siRNA (A) or CLASP1 + 2#B siRNAs (B), fixed with methanol, and stained for α-tubulin. Bars, 10 μm. (C) Plots of integrated intensity of fluorescence of α-tubulin staining, which was performed as in A and B, in HeLa cells 72 h after transfection with the indicated siRNAs. The integrated intensity was measured within a 5-μm² box at 5 μm distance from the cell edge, with subtracting the background. Measurements were performed in ~30 cells per siRNA in three different cell regions. Values significantly different from the control (P < 0.001) are indicated by asterisks. (D) Number of MT ends in a trapezoid part of the cell sector (with a base b and side h) in HeLa cells 72 h after transfection with the control siRNA (30 cells, n = 587) or CLASP1 + 2#B siRNAs (30 cells, n = 401). (E) Integrated intensity within a 5-μm² box was measured along the cell radius in the same cells as described in D. MTOC, the MT organizing center, was determined as the site of radial gathering of MTs. (F) HeLa cells, transfected either with the control siRNA or the CLASP1 + 2#B siRNAs, were methanol fixed and stained for CLASPs (using with the mixture of antibodies #402 or #2358) and for tubulin 72 h after transfection. Representative cells were selected to include examples with both high and low levels of CLASP staining. Integrated intensity of CLASP staining was plotted versus integrated intensity of tubulin staining. Each dot represents an average of five independent measurements in different areas at the periphery of one cell. The regression line is shown in red and the 95% confidence interval is indicated by gray lines. (G) Distribution of distances from MT ends to the cell edge within the trapezoid part of the cell sector shown in D. Analyzed images were the same as in D. (H) Distribution of angles of distal segments of MTs to the cell radius. Analyzed images were the same as in D. (I) Confocal microscope images (1-μm optical sections) of HeLa cells, fixed with formaldehyde 72 h after transfection with the control or CLASP1 + 2#B siRNAs and stained for α-tubulin. Control cells were also incubated for 1 h at 4°C before fixation, to depolymerize MTs (“cold treated”). Bar, 10 μm. (J) Plots of the ratio of the intensity of soluble tubulin staining I_s (after subtracting background) to the intensity of polymer staining I_p (after subtracting background and I_b), obtained from the images as shown in H. Measurements were performed in 20 control cells (n = 158), 35 CLASP-knockdown cells (n = 225), and 18 cold-treated cells (n = 130). Error bars represent the SD.
This increase was above the levels in control cells (19.6 ± 4.1), probably due to overexpression of the rescue construct by a factor of 3–4 compared with endogenous CLASPs (unpublished data). Remarkably, the time the MT plus ends stayed in the vicinity (1 μm distance) from the cell edge was significantly reduced after both CLASPs were knocked down and restored after the rescue with mRFP-CLASP2α (Fig. 3 B).

We have also analyzed HeLa cells stably expressing a GFP fusion of EB1, which binds predominantly to the ends of growing MTs (Mimori-Kiyosue et al., 2000). In control cells a large number of MT ends at the cell periphery was strongly positive for EB1, suggesting that these MTs were growing (Fig. 3 C). This accumulation of EB1-positive ends was clearly reduced after the partial CLASP depletion and restored after the knockdown cells were rescued with mRFP-CLASP2α (Fig. 3 C; Video 3, available at http://www.jcb.org/cgi/content/full/jcb.200405094/DC1). In many cases, EB1 signals at the periphery of control cells appeared rather static, while their intensity varied greatly. This could be explained if pausing MTs or MTs, which started to depolymerize, retained small amounts of EB1, whereas the enhancement of EB1 signal would correlate with MT growth.

We conclude that when CLASP levels are normal, MT ends are maintained in the proximity of cell margin, but remain dynamic. This can be most easily accounted for by very short alternating bursts of MT growth and shrinkage. Indeed, the number of MTs, displaying longer (>1 μm) depolymerization episodes, was increased after the partial CLASP knockdown (Fig. 3 D). This effect was reversed by introducing the mRFP-CLASP2α rescue construct.

The analysis of parameters of MT dynamic instability (Table I) revealed that in the cortical 1-μm-wide region MTs spent more time pausing and exhibited less frequent transitions to growth or shortening in control cells, than in cells with partially depleted CLASPs. In addition, shortening and growing MTs underwent more frequent reverse transitions (rescues or catastrophes, respectively); as a result, dynamic MT ends were maintained in the vicinity of the cell margin, accounting for the
HeLa cells, stably expressing GFP-tubulin or EB1-GFP, were imaged with a 2 s interval 72 h after transfection with the control or CLASP1-2 siRNAs. Life history plots of 65 MTs in five control cells and 75 MTs in five CLASP knockdown cells were analyzed as described by Shelden and Wadsworth (1993). In addition, to obtain more accurate values of growth and shortening rates in the internal cytoplasm, subtraction analysis (Vorobjev et al., 1999) and  measurements of displacements of EB1-GFP comets were used. The differences in rates, obtained by different methods, can be explained by a higher contribution of the pausing state to the growth/shortening episodes at the cell periphery, which can be detected by direct observation, as compared with those inside the cells, which can be identified by subtraction analysis. EB1 associates predominantly with growing MTs (Mimori-Kiyosue et al., 2000) and, therefore, MT pauses hardly contribute to the growth episodes, which can be detected by this method. The data are presented in format: mean ± SD. Rates are instantaneous (measured within one interval between successive frames).

Table I. Parameters of dynamic instability

<table>
<thead>
<tr>
<th></th>
<th>Control 1-µm region at the cell edge</th>
<th>Cell interior</th>
<th>CLASP knockdown 1-µm region at the cell edge</th>
<th>Cell interior</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rate of growth, µm/min</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Direct MT observation</td>
<td>9.9 ± 3.5</td>
<td>11.7 ± 4.8</td>
<td>11.1 ± 1.4</td>
<td>12.6 ± 5.1</td>
</tr>
<tr>
<td>Subtraction analysis</td>
<td>14.4 ± 3.9</td>
<td>15.9 ± 6.0</td>
<td>18.2 ± 5.8</td>
<td>22.8 ± 7.3</td>
</tr>
<tr>
<td>EB1 tracks</td>
<td></td>
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<tr>
<td>Rate of shortening, µm/min</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Direct MT observation</td>
<td>−12.0 ± 6.2</td>
<td>−20.7 ± 11.6</td>
<td>−12.6 ± 6.5</td>
<td>−17.8 ± 11.3</td>
</tr>
<tr>
<td>Subtraction analysis</td>
<td>−35.6 ± 11.2</td>
<td></td>
<td>−28.2 ± 9.8</td>
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<tr>
<td>Transition frequencies</td>
<td></td>
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<tr>
<td>Growth–Shortening, s⁻¹</td>
<td>0.078</td>
<td>0.035</td>
<td>0.030</td>
<td>0.045</td>
</tr>
<tr>
<td>Growth–Pause, s⁻¹</td>
<td>0.402</td>
<td>0.238</td>
<td>0.360</td>
<td>0.250</td>
</tr>
<tr>
<td>Shortening–Growth, s⁻¹</td>
<td>0.066</td>
<td>0.085</td>
<td>0.019</td>
<td>0.072</td>
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<tr>
<td>Shortening–Pause, s⁻¹</td>
<td>0.275</td>
<td>0.213</td>
<td>0.217</td>
<td>0.210</td>
</tr>
<tr>
<td>Pause–Growth, s⁻¹</td>
<td>0.016</td>
<td>0.053</td>
<td>0.032</td>
<td>0.046</td>
</tr>
<tr>
<td>Pause–Shortening, s⁻¹</td>
<td>0.026</td>
<td>0.038</td>
<td>0.038</td>
<td>0.040</td>
</tr>
<tr>
<td>Time in growth, %</td>
<td>6.0</td>
<td>16.7</td>
<td>13.3</td>
<td>14.5</td>
</tr>
<tr>
<td>Time in pauses, %</td>
<td>88.9</td>
<td>69.3</td>
<td>80.1</td>
<td>70.7</td>
</tr>
<tr>
<td>Time in shortening, %</td>
<td>5.1</td>
<td>14.0</td>
<td>6.6</td>
<td>14.8</td>
</tr>
</tbody>
</table>

Hela cells, stably expressing GFP-α-tubulin or EB1-GFP, were imaged with a 2 s interval 72 h after transfection with the control or CLASP1-2 siRNAs. Life history plots of 65 MTs in five control cells and 75 MTs in five CLASP knockdown cells were analyzed as described by Shelden and Wadsworth (1993). In addition, to obtain more accurate values of growth and shortening rates in the internal cytoplasm, subtraction analysis (Vorobjev et al., 1999) and  measurements of displacements of EB1-GFP comets were used. The differences in rates, obtained by different methods, can be explained by a higher contribution of the pausing state to the growth/shortening episodes at the cell periphery, which can be detected by direct observation, as compared with those inside the cells, which can be identified by subtraction analysis. EB1 associates predominantly with growing MTs (Mimori-Kiyosue et al., 2000) and, therefore, MT pauses hardly contribute to the growth episodes, which can be detected by this method. The data are presented in format: mean ± SD. Rates are instantaneous (measured within one interval between successive frames).

observed longer time of staying of MT ends at the cell periphery in control cells, compared with CLASP knockdown cells. Treatment with CLASP1+2 siRNAs had no significant effect on transition frequencies in the internal cytoplasm, proving that CLASPs are truly local regulators of MT dynamics. However, MT growth rate was higher, and the MT shrinkage rate lower in CLASP knockdown cells, compared with control cells (Table I). This can be explained by the increased level of soluble tubulin after the partial CLASP depletion (Fig. 2, I and J). It should be noted that the large contribution of the pausing to growth and shortening episodes at the cell margin makes the differences in growth and shrinkage rates at the cell periphery much less apparent, than those in the internal cytoplasm.

CLASP1 and 2 bind to EB1 and EB3

To get a better insight into the mechanism of CLASP action at the MT tip, we have determined domains of CLASP1 and 2, responsible for their plus-end accumulation. Our previous study has shown that the COOH-terminal domains of CLASP1 and 2 bind to CLIP-170 and CLIP-115, but are not essential for MT tip targeting by CLASPs (Akhmanova et al., 2001). Deletion mapping of CLASP2(y) (the shortest splice form of CLASP2, which associates efficiently with the MT tips; Akhmanova et al., 2001) identified a region of ~140 aa in the middle of CLASP2, CLASP2-M, which was sufficient to target GFP to the ends of growing MTs (Fig. 4, A–E). The corresponding region of CLASP1 (CLASP1-M) could also direct GFP to the MT tips (Fig. 4, F and G). This region is extremely rich in serines, arginines, and lysines, and contains a number of short direct repeats of several types, which are conserved between CLASP1, CLASP2, and their Drosophila homologue, Orbit/MAST (Fig. 4 H).

To understand how the CLASP2-M fragment targets MT plus ends, we have purified 6x histidine (HIS)–tagged GFP-CLASP2-M and GFP from bacteria (Fig. 4 I). In vitro pelleting assays demonstrated that GFP-CLASP2-M, but not GFP alone, could bind to taxol-stabilized MTs in vitro (Fig. 4 J). It is possible that the interaction between the MTs and CLASP2-M is at least partly due to the positively charged character of this fragment and the presence of highly acidic COOH termini in α- and β-tubulin. Interestingly, +TIPs, which belong to the EB family, such as EB1 and EB3, also possess highly acidic COOH termini, similar to those of tubulins. Therefore, we tested if CLASP2-M could associate with EB1 and EB3, and found that it was indeed the case. Among the CLASP2 fragments, only CLASP2-M could coprecipitate EB1 and EB3 (Fig. 4 K). The corresponding region of CLASP1 could also coprecipitate EB1 and EB3 (Fig. 4 K), suggesting that this property is conserved between the two CLASPs. In vitro, purified GFP-CLASP2-M bound to purified EB1 and EB3, fused to GST. It associated with the COOH-, but not the NH₂-terminal half of the EB1 protein (Fig. 4, L and M). This binding was highly specific, because a number of other basic proteins tested, such as histone H1, did not bind to EB1 COOH terminus (unpublished data).

Also the full-length CLASP1α and CLASP2γ, expressed in COS-1 cells, associated with the GST fusions of EB1, EB3 and EB1 COOH terminus (Fig. 4 M). In contrast, the CLASP2-ΔM mutant, which contains an in-frame deletion removing the EB1-binding fragment, was no longer pulled down by GST
EB1 or EB3 fusions (Fig. 4 M). Endogenous CLASP1 and CLASP2 coprecipitated with EB1-GFP fusion, as well as with the GFP-EB1 COOH terminus, but only when the acidic tail of this protein (the last 18 aa) was intact (Fig. 4 N). The validity of interaction CLASP1/2–EB1 was further confirmed by co-pelleting of endogenous CLASP1 and 2 proteins with endogenous EB1 (Fig. 4 O). Together, these data establish that CLASP1 and 2 bind to EB1 and EB3. This interaction is likely to be transient and regulated, and may occur preferentially in the context of MT tip binding, which would account for the weak coprecipitation of endogenous proteins.

Although GFP-CLASP2-M associated with MT ends (Fig. 4, B–E), unexpectedly, the in-frame deletion of this fragment did not preclude the binding of the rest of the CLASP2γ protein (CLASP2-ΔM mutant) to the MT tips (Fig. 5, A and B). In fact, the COOH-terminal region of CLASP2, containing the CLIP-binding domain, could also weakly target MT tips, suggesting that CLIPs can recruit CLASP to the growing MT ends. We have used a dominant negative CLIP-170 mutant, overexpression of which removes endogenous CLIPs, but not EB1 and EB3 from the MT tips (Fig. 5 C; Komarova et al., 2002), to determine if the binding of different CLASP mutants to MT ends is CLIP dependent. We found that GFP-CLASP2-ΔM mutant depended on the presence of CLIPs for the association with MT ends, whereas GFP-CLASP2-M did not (Fig. 5, D–I). It is possible that the interaction with EB1/EB3, which occurs in addition to binding directly to MTs, increases the affinity of CLASPs for the growing MT ends and contributes to the specific CLASP accumulation at these sites.

Next, we examined if CLASPs can recruit EB1 to MTs and found that indeed, overexpressed CLASP1 and 2 (Fig. 4, A–G) GFP fusions of CLASP2α, CLASP2γ, different CLASP2γ deletion mutants, CLASP1a and its deletion mutants were transfected in COS-1 or COS-7 cells, fixed and counterstained for EB1, and the MT plus-end accumulation was assessed as: ++, strong; +, clearly visible; +/−, weakly detectable, −, undetectable. B, C, and F, GFP signal; D, E, and G, EB1 staining. Bars, 10 μm. [H] Alignment of the repetitive part of the CLASP2-M fragment with the corresponding regions of CLASP1 and D. melanogaster Orbit/MAST. Repeats of different types are indicated by different arrows. [I] Coomassie-stained gel, showing purified HIS-tagged GFP-CLASP2-M and GFP. [J] MT pelleting assay with purified GFP-CLASP2-M and GFP. Coomassie-stained gel is shown for tubulin and Western blots with anti-GFP antibodies for the GFP fusions. S, supernatant; P, pellet. [K] Immunoprecipitation with anti-GFP antibodies from COS-1 cells, transfected with the indicated GFP-CLASP1 and 2 deletion mutants. [L] Schematic representation of EB1 structure and the deletion mutants used in this study. CH, calponin homology domain; CC, coiled coil; Ac, acidic tail domain. [M] GST pull-down assays with the indicated GST fusions. Purified GFP-CLASP2-M and GFP or extracts of COS-1 cells, overexpressing GFP-CLASP1α, GFP-CLASP2, and GFP-CLASP2-ΔM were used. Coomassie-stained gel is shown for the GST fusions and Western blots with anti-GFP antibodies for the GFP fusions. 10% of the input and 25% of the material, bound to the beads, were loaded on gel. [N] Immunoprecipitation with anti-GFP antibodies from COS-1 cells, transfected with the indicated GFP-EB1 fusions. [O] Immunoprecipitation with a rabbit polyclonal antibody against EB1 or a control rabbit IgG from untransfected COS-1 cells.
CLASP2 induces strong accumulation of EB1 along the MTs (Fig. 5, J and L; not depicted). In these conditions, CLIP-170 was also accumulated on the MT bundles (Akhmanova et al., 2001; Fig. 5 K and not depicted). However, EB1 recruitment was CLIP independent, because it was also observed with the CLASP2-ΔC mutant, which cannot bind to CLIPs and, when present at high levels, actually displaced CLIP-170 from the MTs (Fig. 5, M–O). CLASP2-ΔM mutant did not accumulate efficiently along the MTs and did not influence EB1 localization (unpublished data). Although we cannot exclude that CLASPs cause accumulation of EB1 at MT bundles by modifying the structure of the MT lattice, the EB1 recruitment is most likely due to direct CLASP–EB interaction.

**CLASP2 COOH-terminal domain is responsible for targeting to the cell cortex and the Golgi complex**

The observation that MT ends of control HeLa cells localized in the close proximity of the cell margin raised a possibility that they come in contact with the cell cortex. To support this idea, we used total internal reflection fluorescence (TIRF) microscopy, which visualizes only a very narrow region (~200 nm) at the bottom of the cell. Indeed, in live cells, transfected with either GFP-CLASP1 or GFP-CLASP2γ, the accumulation of these proteins at the distal MT ends at the cell periphery was readily observed by TIRF, indicating that they are very close to the substrate (Fig. 6, A and B). In control cells, expressing GFP-α-tubulin or EB3-GFP (a plus-end marker, similar to EB1-GFP; Stepanova et al., 2003), a large proportion of the MT ends at the cell margin was visualized by TIRF (Fig. 6, C and E; Video 4, available at http://www.jcb.org/cgi/content/full/jcb.200405094/DC1), again suggesting their proximity to the cortical region under the cell. In striking contrast, CLASP1+2 siRNA-treated cells displayed only a few peripheral MT ends, visible by TIRF microscopy (Fig. 6, D and F; Video 4), indicating that most of the MT tips at the cell margin are not touching the underlying cortex. Therefore, we conclude that CLASPs bring peripheral MT segments in close contact with the cortex underneath the cell.
These observations prompted us to investigate which CLASP2 domains are responsible for cortical and/or membrane targeting. Deletion analysis demonstrated that the COOH-terminal portion of CLASP2 alone was sufficient to target GFP to the Golgi, whereas the GFP-CLASP2ΔC mutant, lacking the COOH-terminal 278 aa, was unable to accumulate at the Golgi (Fig. 6, G and H; not depicted). To investigate the cortical localization of CLASP2 deletion mutants, we made use of the fact that the peripheral accumulation of the full-length CLASP1 and 2 was partially maintained after the MTs were depolymerized with nocodazole (Fig. 6 I and not depicted). This MT-independent localization was not observed with GFP-CLASP2ΔC mutant, indicating, that, similar to the Golgi localization, it depends on the COOH-terminal domain of CLASP2. It should be noted that our previous studies identified the COOH-terminal 280 aa of CLASP2 as the CLIP-binding region (Akhmanova et al., 2001). However, CLIPs are not present at the Golgi and do not accumulate at the cell cortex in MT-independent manner (unpublished data), suggesting that association of CLASP2 with these structures is not dependent on CLIPs.

Both the EB1-binding and the COOH-terminal domains of CLASP2 are required to rescue the organization of the MT network after CLASP knockdown

We have established that the middle (EB1-binding) and COOH-terminal domains of CLASP2 target the protein to the MT plus ends and the cell cortex, respectively. To clarify, which CLASP domains are critical to stabilize MTs at the cell edge, we have performed rescue experiments using GFP fusions of the full-length CLASPs or CLASP2 deletion mutants (Fig. 7, A–D, G, and H). We have examined the ratio between the MT and GFP signal, to determine, which proteins can restore MT density at low expression levels (Fig. 7 H). From these experiments, it became clear that both full-length CLASPs could restore MT density more efficiently than the CLASP2 deletion mutants CLASP2ΔC or -ΔM, which lack ei-
Figure 7. Rescue of CLASP knockdown with CLASP2 deletion mutants and a model for CLASP action at the cell cortex. (A–F) HeLa cells, depleted for CLASPs as described in Fig. 3 A, were transfected with different rescue constructs (indicated on the left), fixed with methanol and stained for α-tubulin (A–D) or EB1 (E and F). Cell images were collected with the confocal microscope (1-μm-thick optical sections). GFP (green) and tubulin/EB1 (red) signals are superimposed on the right. Enlarged portions of the boxed areas are shown in the insets. Bars, 10 μm. (G) Average of pixel intensity of α-tubulin staining (10 cells per construct). (H) Ratio of the average pixel intensity of MT fluorescence (I_{MT}) to the integrated intensity of GFP fluorescence (I_{GFP}; ~10 cells per construct). (I) Proposed model of CLASP rescue activity at the cell cortex. CLASPs can interact with the plus end of a growing MT directly and/or through the association with EB1 and the same time make contact with the cell cortex. After the MT undergoes a catastrophe, most of the EB1 proteins are lost from the tip, but CLASPs remain associated with the cortex and the peripheral stretches of the depolymerizing MT. By enhancing the affinity of EB1–MT interaction at these sites, CLASPs help to retain and/or recruit EB1 to such MTs, leading to their rescue. The oscillations of EB1 signals at the cortical MT tips in control cells support this idea (with strong EB1 accumulation corresponding to growth episodes, and weak EB1 accumulation observed during pausing/depolymerization events). Error bars represent the SD.
ther the COOH-terminal or the EB1-binding motif. A shorter deletion mutant, CLASP2-N1, displayed no rescue activity in this assay (Fig. 7 G), and was used as a negative control in this experiment. It should be noted that these deletion mutants associated with MT tips in CLASP1 +2 siRNA-treated cells as efficiently as in control cells, suggesting that they do not depend on heterodimerization with endogenous CLASPs for their plus-end localization (although due to the partial character of the knockdown, we cannot fully rule out this possibility). In addition, full-length CLASPs, but not the CLASP2-ΔC or -ΔM deletion mutants were able to rescue the characteristic enrichment of the EB1-positive tips at the cell margin (Fig. 7, E and F; not depicted). These results indicate that both the EB1-binding and the cortex-binding COOH-terminal domains of CLASP2 are needed to stabilize MT ends at the cell periphery.

**Discussion**

In this study, we have used an RNAi approach to reduce the levels of CLASP1 and CLASP2 in HeLa cells. The specificity of the used siRNAs is supported by the fact that two different oligonucleotides for each CLASP produced similar effects on the MT network, which were not observed with control siRNAs. Moreover, significant defects in MT density and stability were only observed when combinations of siRNAs, specific for the two CLASPs, were transfected into cells simultaneously. The phenotype, caused by CLASP1/2-specific siRNAs could be rescued by introducing into cells CLASP1 or 2 expression constructs, insensitive to the used siRNA duplexes. These data indicate that the observed defects were due to reduction of the CLASP1/2 expression, rather than the off target activities of the used siRNAs.

HeLa cells express CLASP1α and CLASP2α isoforms, which are ~77% similar in their protein sequence. Our data, based on the antibody staining, expression of GFP fusion proteins, RNAi-mediated phenotypes and their rescues, indicate that these two CLASPs display very similar localizations and functions in interphase cells. These observations do not rule out, however, that in the context of the whole organism these proteins may have some unique and specific functions. The depletion of CLASPs in our system was only partial, and it is likely that additional and more severe phenotypes would be observed after a complete loss of both CLASPs. Also, we cannot exclude that a complete inactivation of one of the two CLASPs would have different effects compared with the ~70% inactivation of both, although our results have provided no indication for such a possibility.

Our previous study, based on the analysis of tubulin modifications, has shown that CLASPs are involved in locally stabilizing MTs at the leading edge of motile fibroblasts (Akhmanova et al., 2001). Our new findings fully support the role of CLASPs in stabilizing MTs, by showing that the MT density and the amount of acetylated tubulin are reduced after CLASP knockdown. Detailed analysis of MT behavior shows that CLASPs strongly affect MT plus-end dynamics locally, in the 1-μm-broad region at the cell margin. This is in line with the accumulation of CLASP proteins at the distal segments of MTs specifically in this region of the cell cortex. In the presence of CLASPs, MTs at the cell edge remain dynamic, but keep undergoing alternating frequent rescues and catastrophes. MT pausing, apparently stimulated by CLASPs, might in fact be not a truly nondynamic state, but rather very short alternating growth and shrinking episodes, with amplitudes below the resolution of the fluorescent microscope. We propose that the main activity of CLASPs is MT rescue at the cell edge, which reduces the number of long depolymerization episodes and increases MT longevity. It is possible, that interaction with EB1 is mechanistically important for the CLASP rescue activity by stimulating a switch to the growth phase, because EB proteins promote MT growth (Tirnauer et al., 2002; Busch and Brunner, 2004). The capacity of CLASPs to enhance EB1 accumulation at the MTs is fully in line with this idea, which is also supported by the recent findings showing the essential role of EB1 for generation of stable interphase MTs (Wen et al., 2004). The importance of CLASP–EB interaction is underscored by a recently published report, showing that the *Drosophila* CLASP homologue, Orbit/MAST is present among the proteins, pulled down by GST-EB1 (Rogers et al., 2004). We propose that CLASPs, associated with the distal segments of MTs at the cell cortex, are partially retained on MTs when they start shrinking and can rescue MTs by enhancing their affinity for EB1 (Fig. 7 I).

CLASP-dependent rescues are quickly followed by catastrophes, which we believe to be caused by the cell edge acting as an impermeable boundary (Janson et al., 2003). If our assumption is correct, how can we explain that the MTs, rescued by CLASP, do not make a turn and start to grow along the cell edge, as observed after the CLASP1/2 knockdown? We propose that CLASPs, bound to the growing MT tip and at the same time associated with the cell cortex, bring the MT end in very close proximity to the cell edge. An abrupt MT turning at this point would require a considerable distortion of the MT lattice, which would be energetically unfavorable; therefore, a catastrophe occurs instead. Catastrophes at the cell edge may also be caused by MT-destabilizing factors, such as depolymerizing kinesins or stathmin, and we cannot rule out that there is a functional interplay between CLASPs and such MT-destabilizing proteins.

The effect of CLASPs on MT rescue is local, whereas the transition frequencies in the internal cytoplasm are not significantly affected. Therefore, CLASP function is different from that of CLIPs, which affect MT rescue frequency throughout the cell (Komarova et al., 2002). It is possible that CLIP–CLASP interaction can contribute for the CLASP rescue activity. However, we cannot exclude that the interaction with CLIPs regulates CLASP accumulation at the MT tips, as opposed to other cellular structures, such as the cortex or the Golgi apparatus, whereas the effect of CLASPs on MT dynamics is CLIP independent.

The rescue activity of CLASPs is likely to be relevant to the mitotic function of these proteins. CLASPs/Orbit/MAST are present at the kinetochores, and their inactivation results in monopolar spindles with chromosomes buried in the interior of the aster (Maiato et al., 2002, 2003a). If CLASPs exhibit a local MT rescue activity at the kinetochore, similar to that shown...
by us at the cell cortex, this would explain why in the absence of CLASPs/Orbit/MAST the kinetochore fibers are unable to regrow, once they start shortening, causing the chromosomes to be “reeled in” to the pole (Maiato et al., 2002, 2003a).

We have observed very few depolymerizing minus ends either in control or in CLASP knockdown cells (unpublished data), indicating that in HeLa cells MT renewal is accounted for by plus-end dynamics and that CLASPs do not play a significant role in stabilizing MT minus ends. However, the striking effect of CLASP depletion on MT density indicates, that in addition to regulating plus-end dynamics, CLASPs might play a role in MT nucleation.

Recent studies have identified two other large +TIPS, spectraplakin ACF7 and APC, as MT-stabilizing proteins, which, similar to CLASPs, act the leading edge of motile cells (Kodama et al., 2003; Wen et al., 2004). Similar to CLASPs, APC also binds to EB1 and is negatively regulated by GSK3β (Su et al., 1995; Zumbrunn et al., 2001). ACF7 might be an EB1-binding protein, too, because its Drosophila homologue, Short stop, interacts with the fly homologue of EB1 (Subramanian et al., 2003). CLASPs and APC can bind the cell cortex (this study; Mimori-Kiyosue and Tsukita, 2001), and ACF7 can directly associate with actin filaments (Kodama et al., 2003), suggesting that these proteins can link MTs to these structures. Studies of MT organization at the muscle–tendon junction in fly larvae indicate that Short stop and APC are acting together (Subramanian et al., 2003). No similar data are available for their mammalian counterparts, so the analysis of a potential functional synergism or redundancy between CLASPs, APC, and the spectraplakins presents an important challenge for future studies.

Materials and methods

Cell lines and transfection of plasmids and siRNAs

HeLa, COS-1, and COS-7 cells were grown as described previously (Akmanova et al., 2001). Effectene or Superfect transfection reagents (QIAGEN) were used for plasmid transfection. Mass transfection of COS-1 and COS-7 cells was performed by DEAE-dextran method (Akhmanova et al., 2001). Synthetic siRNAs (Proligo) were transfected, using Oligofectamine (Invitrogen). SiRNAs were directed against the following target sites: CLASP1#A, GCCATTATGCCAACTATCT; CLASP1#B, GGATGATTTA−β-tubulin and acetylated tubulin (Sigma-Aldrich), actin (CHEMICON International, Inc.); rat αβ-tubulin (YLI/2; Abcam), rabbit pAb against GFP (MBL; CHEMICON International Inc., and Abcam), CLASP2 (Akmanova et al., 2001) and CLASP2 siRNA (Dharmacon, Inc.) and Statistica for Windows (StatSoft Inc.). Unless stated differently, images were prepared for publication using Photoshop (Adobe). Statistical analysis was performed using a SigmaPlot (SPSS Inc.) and Statistica for Windows (StatSoft Inc.). Unless stated differently, the statistical significance of the observed differences was evaluated using Kolmogorov-Smirnov two-sample test and in all plots SD is indicated. For FACS analysis, suspended cells were fixed in cold ethanol, labeled with anti-CLASP1/2 antibodies and Cy5-conjugated secondary antibody, and analyzed using FACS Calibur (Becton Dickinson).

Fluorescence microscopy and image analysis

Images of cells were collected with a DeltaVision optical sectioning system using PlanApo 100×/1.40 NA oil, PlanApo 60×/1.40 NA oil ph 3 or PlanApo 20×/0.70 NA dry objectives (Olympus), using a cooled CCD camera (Series300 C1330; Photometrics). Fluorescence signals were visualized using a quad filter set (M6000; Chroma Technology Corp.) for multiple color imaging, or Endow GFP bandpass emission filter set (41017; Chroma Technology Corp.) for GFP imaging. The auto-focus signals were removed using the deconvolution technique with the DeltaVision system. Confocal imaging was performed using LSM510 confocal laser scanning microscope (v. 2.3; Carl Zeiss MicroImaging, Inc.). TIRF microscopy was performed on an Olympus IX70 (PlanApo 100×/1.45 NA, oil TIRF objective), equipped with a 488-nm argon laser line (MELLES GRIOT), an objective-typed TIRF illuminator (Olympus), and an OrcaER cooled CCD camera (Hamamatsu Photonics). The system was controlled by Aquacosmos software (Hamamatsu Photonics). The quantification and analysis of fluorescence signals was performed using MetaMorph software (Universal Imaging Corp.). Images were prepared for publication using Photoshop (Adobe). Statistical analysis was performed using with a SigmaPlot (SPSS Inc.) and Statistica for Windows (StatSoft Inc.). Unless stated differently, the statistical significance of the observed differences was evaluated using Kolmogorov-Smirnov two-sample test and in all plots SD is indicated. For FACS analysis, suspended cells were fixed in cold ethanol, labeled with anti-CLASP1/2 antibodies and Cy5-conjugated secondary antibody, and analyzed using FACS Calibur (Becton Dickinson).
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