Introduction

Cell shape is a result of the interplay between hydrostatic pressure exerting outward force on the plasma membrane, lateral tension of the membrane itself, and contractile forces exerted by the actomyosin cytoskeleton (Sheetz, 2001; Mitchison et al., 2008). Cell shape can largely be explained by varying contributions of these forces. For instance, on the leading edges of crawling cells, protrusion is driven by combinations of densely cross-linked actin and the locally elevated hydrostatic pressure that results from local ion transport and water influx into that region (Stock and Schwab, 2006; Fackler and Grosse, 2008). In contrast, blebs are the result of hydrostatic pressure that causes a segment of plasma membrane to rip away from the cortical actomyosin cytoskeleton, whereas bordering membrane remains firmly anchored by myosin II (Charras et al., 2008). In contrast, blebs are actin poor during their expansion phase, and this expansion is limited both by local surface area and by repolymerization of actin on the bleb surface that is associated with retraction (Charras et al., 2006). It has recently been shown that these same forces of positive hydrostatic pressure and contractile actomyosin control cell rounding during mitosis (Stewart et al., 2011).

Septins are a family of GTP-binding proteins that self-assemble into hexameric or octameric quaternary structures and further into large filaments, rings, and gauzes in vitro and in vivo (Kinoshita et al., 2002; Kinoshita, 2003; Rodal et al., 2005). A variety of functions has been described for septins in mammalian cells. Principle among these are contouring of phosphatidylinositol-containing membranes in vitro, templating of actin fibers, stabilization of microtubule tracks, and interaction with exocytic machinery (Kinoshita et al., 1997; Hsu et al., 1998; Surka et al., 2002; Tanaka-Takiguchi et al., 2009; Gilden and Krummel, 2010). Structurally, they appear to be required to maintain cilia and to assist in the completion of phagocytosis (Huang et al., 2008; Hu et al., 2010). The relative importance of each of these roles remains unclear, but knockout animals lacking specific septin subunits exhibit defects in cytokinesis, neuronal development and function, platelet activation, and spermatozoa motility (Kinoshita et al., 1997; Nguyen et al., 1998; Surka et al., 2002; Tanaka-Takiguchi et al., 2009; Gilden and Krummel, 2010). Structurally, they appear to be required to maintain cilia and to assist in the completion of phagocytosis (Huang et al., 2008; Hu et al., 2010). The relative importance of each of these roles remains unclear, but knockout animals lacking specific septin subunits exhibit defects in cytokinesis, neuronal development and function, platelet activation, and spermatozoa motility (Kinoshita et al., 1997; Nguyen et al., 1998; Surka et al., 2002; Tanaka-Takiguchi et al., 2009; Gilden and Krummel, 2010). Structurally, they appear to be required to maintain cilia and to assist in the completion of phagocytosis (Huang et al., 2008; Hu et al., 2010). The relative importance of each of these roles remains unclear, but knockout animals lacking specific septin subunits exhibit defects in cytokinesis, neuronal development and function, platelet activation, and spermatozoa motility (Kinoshita et al., 1997; Nguyen et al., 1998; Surka et al., 2002; Tanaka-Takiguchi et al., 2009; Gilden and Krummel, 2010).

Increasing evidence supports a critical role for the septin cytoskeleton at the plasma membrane during physiological processes including motility, formation of dendritic spines or cilia, and phagocytosis. We sought to determine how septins regulate the plasma membrane, focusing on this cytoskeletal element’s role during effective amoeboid motility. Surprisingly, septins play a reactive rather than proactive role, as demonstrated during the response to increasing hydrostatic pressure and subsequent regulatory volume decrease. In these settings, septins were required for rapid cortical contraction, and SEPT6-GFP was recruited into filaments and circular patches during global cortical contraction and also specifically during actin filament depletion. Recruitment of septins was also evident during excessive blebbing initiated by blocking membrane trafficking with a dynamin inhibitor, providing further evidence that septins are recruited to facilitate retraction of membranes during dynamic shape change. This function of septins in assembling on an unstable cortex and retracting aberrantly protruding membranes explains the excessive blebbing and protrusion observed in septin-deficient T cells.

The septin cytoskeleton facilitates membrane retraction during motility and blebbing

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Abbreviations used in this paper: DIC, differential interference contrast; GPI, glycosylphosphatidylinositol; NPPB, 5-Nitro-2-(3-phenylpropylamino)benzoic acid; RVD, regulatory volume decrease; shRNA, short hairpin RNA.

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Results

Blebbing and protrusion in septin-deficient cells

We have previously shown that cloned T cell lines expressing short hairpin RNAs (shRNAs) against SEPT7 are largely deficient in all septins and have profound morphological defects that include cortical instability (Fig. S1; Tooley et al., 2009). SEPT7KD cells display periodic blebbing along the cell body (Tooley et al., 2009). The observation that septins may play a role in regulating the shape of the cell cortex led us to the current study, in which we sought to understand the interplay between septins and the forces that regulate cell protrusions. To this end, we manipulated the outward hydrostatic pressure experienced by cells and investigated the roles of septins in response to that force. We found that, in aggregate, septins are required for rapid contraction of the cell cortex, in cooperation with the actomyosin cytoskeleton. Septins dramatically assemble at sites of cortical overextension, a class of events that includes both leading edge protrusions and membrane blebs, where they prove necessary to counteract persistent and extensive blebbing.
pattern (Tooley et al., 2009). For comparison with total plasma membrane, SEPT6-GFP was coexpressed with glycosphatidylglycerol (GPI)-linked mCherry, and cells were imaged while crawling on intercellular adhesion molecule–1 (ICAM-1)-coated coverslips. During motility, crawling cells extended protrusions at their leading edges, and these were sparsely populated by septins. Strikingly, when cells changed directions and retracted leading edges, those retracting protrusions frequently (83%; n = 18) led to a transient increase in membrane SEPT6-GFP. This accumulation rapidly merged backward into the SEPT6-GFP collar (Fig. 1 b and Video 1). This demonstrated that dynamic enrichment of septins normally occurs at sites of retracting membrane near the leading edge and provided a possible explanation for the extra extensions and blebs that are found in cells lacking this cytoskeleton.

Ameoboid motility is mediated by extensive actomyosin contractile forces originating at the rear of the cell that are thought to generate leading edge protrusions. Because septin deficiency seemed to most prominently affect cortical stability at sites on the membrane that were distant from the major region of myosin contraction in crawling cells, we sought to determine whether septin deficiency also regulated cortical stability at other sites distal to myosin contraction. In cytokinesis, septins colocalize prominently with the contractile actin and myosin at the cleavage furrow. To investigate whether septins might also function distal to the cleavage furrow, control and SEPT7KD D10 T cells were treated with nocodazole for 16 h to synchronize their cell cycles and were imaged during the cytokinesis that followed washout. Though SEPT7 is dispensable for successful mitosis in T cells, an unstable membrane during cell division was indeed readily observed and was notable for occurring mostly distal to the cleavage furrow. Thus, whereas control cells had a mostly smooth plasma membrane during cell division with only a few small blebs occasionally appearing (Fig. 1 c and Video 2), SEPT7KD cells had surfaces studded in tiny blebs, which were also more long lived than those seen on control cells, often persisting from the time of their appearance until the end of the time-lapse, up to 30 min (Fig. 1 c and Video 2). When these blebs were counted, SEPT7KD cells produced, on average, 2.5 times as many blebs per surface area and unit time (blebbing index in Fig. 1 c).

Using SEPT6-GFP–expressing cells, we found GFP primarily concentrated at the cytokinetic furrow during the completion of mitosis (Fig. 1 d), consistent with observations of septins in other mammalian cells and in yeast (Gladfelter et al., 2001; Spiliotis et al., 2005). However, confocal imaging during cytokinesis (Fig. 1 e) emphasized that, whereas SEPT6-GFP is predominantly localized to the cleavage furrow, a septin density is associated with the entire cell cortex, perhaps serving to control mitotic blebs at distal sites. Thus, a line scan through a mitotic cell demonstrates peaks of intensity at each cell border in addition to a large peak at the site of the cytokinetic furrow (Fig. 1 e). Together, these data show that septins are enriched at sites of membrane instability, where they seem to control protrusions and blebs.

**Septins are required for cortical retraction in a hydrostatic volume-change assay**

These observations suggested the possibility that rather than preventing the onset of protrusions and blebs, septins might be recruited to sites of nascent extension and play a role in resolving those structures after they arise. Either mechanism would give rise to increased numbers of extensions and blebs. To distinguish between these possibilities, we used osmotic volume change as a model for cortical dynamics. Hydrostatic pressure was manipulated by changing the osmolality of cells’ media, and this led to cellular expansion and subsequent contraction. When cells are exposed to hypotonic medium, they respond in two phases: rapid expansion as water flows into the cell followed by a slower regulatory volume decrease (RVD) phase, during which ion transport–dependent processes return the cytosol to its native tonicity and cell volume shrinks (Grinstein et al., 1984). In such an assay, local hydrostatic pressure inside the cell causes the plasma membrane to separate from cortical actomyosin (Charras et al., 2008), effectively generating a cell-wide bleb.

To monitor the progress of this swelling and contraction, cell volume in isotonic media was measured using the forward scatter parameter of a flow cytometer. Cells were then shifted to isotonic media, and their volumes were monitored over time. Three features of cellular swelling and contraction could then be measured: initial volume, maximum volume, and the rate of contraction (or RVD). This experiment is schematized in Fig. 2 (a and b). SEPT7KD and control cells had indistinguishable initial volumes and expanded to the same extent, exhibiting a total change in diameter of 14% for controls and 18% for SEPT7KD (Fig. 2 c). However, the knockdown samples had smaller slopes in the RVD period (0.320 ± 0.021 [SEM] compared with 0.490 ± 0.025 for controls; P < 0.001; Fig. 2 d) and took longer to return to their original size (149.7 ± 8.205 vs. 206.0 ± 11.81 s; P < 0.01). This indicated that SEPT7KD cells are retarded in their ability to recontract the cortex to its original size and that septins on the whole are required for retraction rather than for preventing initial separation of membrane from the cortical cytoskeleton.

**Septin assembly during cortical retraction**

To better understand what role septins might be playing during cortical contraction, cells were immobilized on anti-CD44–coated coverslips and stained with antibodies to SEPT7 under isotonic conditions and 10 min after shifting to hypotonic conditions. When cells were immobilized, SEPT7 distribution was less strikingly corsetlike than when cells crawl on ICAM-1 or bare glass (Fig. 3 a). Nonetheless, a clear change in septin distribution was observed when cells were moved to hypotonic media. Under isotonic conditions, 74.6% (n = 59) of cells had SEPT7 enriched at the cortex in puncta (Fig. 3 a). After 10 min in hypotonic media, however, only 20% (n = 75) of cells had this distribution. The remaining 80% of cells had SEPT7 redistributed into dramatic superstructures. 60% of cells examined had septin fibers, especially at the bottom surface where the cell contacted the coverslip, and 48% of cells had visible ~1-μm SEPT7 rings on their cortexes (Fig. 3 b). In addition
by septins might participate in the confinement or shaping of dynamic membrane.

Septin aggregation is not sufficient to mediate cortical contraction without ion channel function

An important step in the recovery from osmotic expansion is active transport of K⁺ and Cl⁻ ions that return the cytosol to normal osmolarity, and this has been shown to be necessary for RVD (Grinstein et al., 1984). We used charybdotoxin and 5-Nitro-2-(3-phenylpropylamino)benzoic acid (NPPB), inhibitors of Ca²⁺-gated K⁺ channels and of Cl⁻ channels, respectively, to block ion transport during our flow cytometry volume-change assay. As anticipated, cells treated with the inhibitors had normal volume under isotonic conditions, normal expansion after the shift to hypotonic conditions (Fig. 4 a), and slowed RVD in the presence of both inhibitors (slopes of 0.114 and 0.056 for charybdotoxin and NPPB compared with a slope of 0.323 for untreated control; Fig. 4 b). When SEPT7KD cells were treated with the drugs, neither demonstrated an additive effect on cell shrinkage, though the effect of NPPB was so severe that an additive effect of septin depletion may have been undetectable. These reagents and their profound block of RVD provided a mechanism to determine whether septins were only recruited as a result of cortical contraction or whether they were recruited to the increase in the frequency of cells with SEPT7 rings, the number of rings per cell (Fig. 3 c) was increased under hypotonic conditions (2.1 ± 1.2 rings per cell for isotonic condition vs. 3.4 ± 2.5 for hypotonic condition; only cells with at least one ring were analyzed).

To observe how septin structures formed in relation to the events of swelling and volume decrease, we monitored the process in real time using SEPT6-GFP–expressing cells. To achieve this, transfected cells were immobilized on anti-CD44–coated coverslips, and after 1 min of imaging, water was added to the imaging chamber to reduce the osmolarity of the media by 2/3. Within 1 min after the change in osmolarity, SEPT6-GFP began to aggregate on the cortex of the cells (Fig. 3 d and Video 3). Over time, these aggregates became pronounced rings, which, when measured, were again on the order of 1 µm in diameter. These expanded in number and ultimately studded the whole surface of the cell and persisted for >30 min (Video 4). These rings were apparent in nearly every cell (Video 5), and the dominance of these circular structures compared with the higher proportions of filaments, seen using anti-SEPT7 antibody staining, may derive from SEPT6 overexpression. When SEPT6-GFP rings were examined more closely in cells coexpressing GPI-mCherry to mark the plasma membrane, dimplelike membrane invaginations could be seen beneath the septin rings (Fig. 3 e and Video 6). This suggested that the structures formed...
independently and might then function in subsequent steps. When SEPT6-GFP–expressing cells were shifted to hypotonic media in the presence of charybdotoxin or NPPB, they formed SEPT6-GFP rings indistinguishable from those formed by control cells (Fig. 4 c), indicating that septin rings are not sufficient to mediate cell shrinkage if cells cannot recover their normal osmolality and that formation of septin structures on the plasma membrane does not require cells to effectively shrink.

Septins collaborate with actomyosin to promote cortical retraction and assemble after actin depletion

Rearrangement of the cortical actin cytoskeleton after changes in hydrostatic pressure has been described in several cell types (Cornet et al., 1993; Morán et al., 1996; Pedersen et al., 1999; Carton et al., 2003; Tamma et al., 2007). As interactions of septins with both actin and myosin II have been described, we tested whether the actomyosin cytoskeleton is also required for cell shrinkage after osmotic expansion. None of the inhibitors tested—blebbistatin (a MyoIIA inhibitor), latrunculin B (an actin polymerization inhibitor), or nocodazole (a microtubule inhibitor)—significantly influenced the volume of cells in isotonic media nor their maximum expansion (Fig. 5 a). However, cells treated with latrunculin B shrank with a rate of 0.289, and those treated with blebbistatin to inhibit MyoII activity shrank with a rate of 0.347 compared with 0.490 for vehicle-treated cells. In contrast, depolymerization of microtubules with nocodazole had no impact on the rate of volume change (0.453; Fig. 5 b). Thus, both actin and myosin appear to be required to similar extents as septins in this process. Shrinkage of SEPT7KD cells treated with latrunculin B was not additionally impaired, whereas parallel experiments using ion transport inhibitors could have added effects (Fig. 4), suggesting that septins and the actomyosin cytoskeleton may be functioning together to affect cortical contraction (Fig. 5 c).

Immunofluorescence staining for endogenous SEPT7 demonstrated that cells treated with blebbistatin rearranged septin complexes similarly to control cells when hydrostatic pressure was increased, with 64% of cells assembling septin circles or filaments in control cells and 75% assembling these structures in the presence of blebbistatin (Fig. 5, d and e). This suggests that loss of myosin II activity has relatively little effect on septin distribution in either steady-state or swelling responses. In contrast, latrunculin B treatment induced the spontaneous formation of SEPT6-GFP rings even under isotonic conditions, and these rings persisted under hypotonic conditions (Fig. 5 f). These data suggest that septin assembly after osmotic expansion might be triggered by actin depolymerization, a result consistent with a previous study in a fibroblast cell line (Kinoshita et al., 2002). When cells were treated instead with jasplakinolide to stabilize actin filaments, cells rounded up and lost their characteristic septin band at the midzone and instead had SEPT6-GFP distributed diffusely or in small puncta, resembling the distribution seen in untreated rounded cells (Fig. 5 f and Video 7). Importantly, SEPT6-GFP rings were not formed after jasplakinolide treatment, demonstrating that disruption of normal actin dynamics alone was not sufficient for septin ring formation. In conclusion, septins collaborate with the actomyosin cytoskeleton during cortical retraction. Although MyoII inhibition doesn’t affect septin ring formation, ring formation and/or assembly is enhanced by loss of actin integrity through drug treatment or in the course of normal cellular events.

To investigate in more detail the nature of the collaboration between septins and the actomyosin cytoskeleton during...
that they had similar initial side scatter values, but after the initial drop, SEPT7KD cells recovered more slowly than control cells (slope of 0.029 ± 0.015 for SEPT7KD compared with 0.061 ± 0.014 for controls; Fig. 6, a and b). This indicates that SEPT7KD cells have a defect in recovering membrane complexity concurrent with their slowed shrinkage.

As septins accumulated around actin-rich protrusions and appeared to promote reshaping of excess plasma membrane during RVD, we wanted to test whether they also serve this function in more physiological structures such as blebs. We reasoned that as dynamin activity is required for many endocytic pathways, dynamin blockade would result in excess plasma membrane in need of recapture, possibly by septins. Indeed, when cells were treated with dynasore for 30 min before imaging, they displayed grossly normal morphology, but over a 10-min imaging period, 60% of cells underwent membrane blebbing at the front half of the cell (unpublished data). Interestingly, this blebbing resembled that previously observed in SEPT7KD cells in that it was stochastic and episodic, with cells crawling normally before blebbing began and resuming normal crawling after the blebbing resolved. By treating SEPT6-GFP–expressing cells with dynasore, we were able to use confocal microscopy to directly observe the response of septins to membrane blebs. Before the onset of blebbing, SEPT6-GFP began to accumulate at their bases, persisting there throughout the retraction of each ruffle. These observations demonstrate a spatial relationship between septins and F-actin during RVD and suggest that septins may play a role in regulating retraction through their accumulation at the edges of actin-rich structures.

Septins reshape membrane during cortical contraction

Under many conditions, including swelling, cells respond to cortical stretch initially by smoothing local surface topography such that excess plasma membrane from ruffles and microvilli is recruited into the region under tension (Grinstein et al., 1984). As septins have been implicated in membrane contortion in vitro (Tanaka-Takiguchi et al., 2009) and as we observed topographical changes at the plasma membrane with the assembly of septin rings (Fig. 3 e), we sought to establish whether septins’ role in cell shrinkage might be to recontort excess plasma membrane. To investigate this, we took advantage of the side scatter parameter of a flow cytometer, which measures laser light scattered orthogonal to the direction of incident light and correlates with a cell’s membrane complexity. When we tracked the side scatter of cells subjected to hypotonic stress, we found that during the hydrostatic expansion phase, there was a precipitous drop in the mean side scatter value followed by a gradual rise over the remaining duration of the experiment, coincident with cortical contraction (Fig. 6 a). When we compared control and SEPT7KD cells, we found that they had similar initial side scatter values, but after the initial drop, SEPT7KD cells recovered more slowly than control cells (slope of 0.029 ± 0.015 for SEPT7KD compared with 0.061 ± 0.014 for controls; Fig. 6, a and b). This indicates that SEPT7KD cells have a defect in recovering membrane complexity concurrent with their slowed shrinkage.

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blebbing. High-resolution confocal imaging of individual blebs in these cells indicated that during initial bleb expansion, the bleb membrane was remarkably deficient in SEPT6-GFP. However, septins accumulated at the fully expanded bleb membrane and remained enriched there during bleb retraction (Fig. 6d).

Actin has also been shown to polymerize on blebs during their retraction phase (Charras et al., 2006), so we sought to examine how septins and actin interact on these structures. To this end, we used confocal imaging to visualize SEPT6-GFP and LifeAct-Ruby on dynasore-induced blebs. As demonstrated in Fig. 6e and Video 10, LifeAct-Ruby was recruited to blebs shortly after their appearance, though the structures remained empty of SEPT6-GFP. 10–20 s later, however, SEPT6-GFP was recruited to the retracting bleb, remaining enriched there throughout retraction. This is consistent with our finding that septins were recruited to actin-containing ruffles, apparently participating in their retractile phase (Fig. 5g). In sum, these data demonstrate recruitment of septins after actin polymerization on unstable dynamic membranes in a variety of settings and suggest a function for septins in providing a platform for membrane reabsorption.

**Discussion**

**Mechanisms of septin-mediated membrane recapture**

In this study, we investigated the mechanistic role of the septin cytoskeleton in controlling the T cell cortex. We observed abnormally frequent and persistent membrane blebs during motility and during cell division in septin-deficient cells and found that retracting protrusions in crawling cells usually resolve into the septin-rich midzone of the cell. These findings led us to develop an experimental system in which local blebs were modeled by global hypotonic expansion of the whole cell and bleb retraction was modeled by compensatory RVD. This system
recruitment in our cortical retraction model. This would be consistent with a study of Ehrlich ascites cells that demonstrated a drop in total cellular PIP2 during initial hypotonic swelling and increased PIP2 production during RVD (Nielsen et al., 2007). Our work did not address whether phosphoinositides indeed recruit septins to the plasma membrane during cortical contraction. In fact, our experiments suggest the possibility that septins could be recruited directly in response to particular actin dynamics or cellular protrusions. We found that during cortical contraction, septins assemble on the cortex in rings similar to those observed during latrunculin B–induced actin depolymerization in our system and others (Kinoshita et al., 2002). A feature of the response to hypotonic conditions in several cell types is the loss of actin stress fibers and subsequent repolymerization of actin in peripheral patches (Carton et al., 2003; Tamma et al., 2007). Similarly, we have observed dynamic flarelike actin structures arising during RVD, with aggregated septins appearing at their bases. This supports the idea that septins are recruited to the cortex coordinately with actin structures. Further work will be required to understand the precise nature of the recruitment in our cortical retraction model. This would be consistent with a study of Ehrlich ascites cells that demonstrated a drop in total cellular PIP2 during initial hypotonic swelling and increased PIP2 production during RVD (Nielsen et al., 2007).

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of these structures, how this intersects with septin recruitment by phosphoinositides, and indeed whether that recruitment relies on actin assembly or other cellular components that themselves also dictate actin assembly. Given the results that actin assembly blockade promotes septin ring assembly, it is tempting to speculate that a cellular sensor of depolymerized actin exists and directs both enhanced assembly and septin membrane recruitment for the purposes of membrane control.

Although in vivo evidence of membrane contortion by septins has not yet been shown, observations of septin localization in several concave cortical structures are consistent with our findings. For example, SEPT2 and SEPT11 are recruited to the phagocytic cup in macrophages, concurrent with PIP₂ accumulation, and the presence of septins on this structure is essential for phagocytosis (Huang et al., 2008). Similarly, PIP₂ accumulation at the mammalian cleavage furrow may also play a role in septin recruitment to that site (Field et al., 2005). Further, septins are required for dendritic spine formation by neurons and associate with the plasma membrane at the concave bases of dendritic spines (Tada et al., 2007; Xie et al., 2007).

The physical mechanism by which septins contort the plasma membrane to function in cortical retraction remains to be explored. Of particular interest will be further investigation of the interactions between the septin cytoskeleton and cortical actin in this setting. An important step in bleb retraction is actin polymerization along the inner surface of the bleb and attachment of actin filaments to the plasma membrane by ezrin/radixin/moesin proteins (Charras et al., 2006). We found that septins accumulated on retracting blebs after this actin polymerization. Together with our observation that septins and the actomyosin cytoskeleton function in concert to contract the cortex during RVD, this suggests a model in which septins function together with actin in bleb retraction, perhaps interacting with newly polymerized actin to pull the bleb membrane back to the still intact cortical actomyosin network.

Despite our finding that septins are recruited to sites of dynamic cortical instability, in the steady state, the majority of septins are present in the relatively static midzone of the cell. There are several reasons why this could be the case. First, the midzone and uropod of the cell may be more dynamic than is generally appreciated, and septins assemble there to continuously control separation between the plasma membrane and cortical actin. Alternatively, septins at the midzone may not be stable there but rather may continuously leave the midzone and be replenished as septin-rich retracted membranes (Fig. 1b) flow back into this region. Either of those possibilities could provide clues to the uropodal elongation observed in SEPT7KD cells, as either would lead to accumulation of excess unretracted membrane in that region in the absence of septins. Finally, we cannot exclude the possibility that septins have several distinct functions in T cells and that their presence at the midzone and role in elongating the uropod are unrelated to the mechanisms we have described here. The population of septins in the midzone may be more stable than the population we have studied here, and this variation could perhaps be a result of the diversity in the subunit composition of septin complexes.

**Septin-mediated membrane retraction in motility**

The phenotype of septin-deficient T cells is quite distinct from those observed in other septin-deficient cells. Whereas some cell types, such as neurons and spermatozoa, exhibit morphological changes associated with disruption of septins, in no other system has such dramatic cortical instability been observed. This raises the possibility that septin function in amoeboid cells is different from that in other cell types. Because amoeboid cells such as T cells move very rapidly, their cortices are extremely dynamic, accommodating the near-constant shape change associated with this rapid motility. Amoeboid cells can adopt a variety of motility modes depending on the 3D environment in which they find themselves (Renkawitz and Sixt, 2010). Among these modes is blebbing motility, in which the major leading edge extension is accomplished via polarized blebbing and myosin-driven cytoplasmic flow (Blaser et al., 2006; Langridge and Kay, 2006). Our observation that septins are involved in bleb retraction in T cells suggests that spatial regulation of septins could play a role in restricting blebs to the leading edges of cells exhibiting this kind of motility. Alternatively, septin regulation could be an important component of switching between blebbing motility and other modes.

Understanding whether the composition of septin complexes controls modes of motility may be of particular importance as more is learned about the contributions of septin dysregulation to tumor metastasis. A growing number of studies have found connections between expression of specific septin monomers or splice variants and tumor metastasis, but a mechanism for this relationship remains to be discovered (Osaka et al., 1999; Montagna et al., 2003; Connolly et al., 2011). Overexpression of several SEPT9 isoforms is associated with enhanced motility and invasion in tumor cell lines, and knockdown of Sept9 in metastatic tumor cell lines reduces motility, invasion capacity, and leading edge protrusion (Chacko et al., 2005; Shankar et al., 2010). As tumor cells have been shown to adopt blebbing motility under conditions that are unfavorable to their more common fibroblast-like motility (Friedl and Wolf, 2003; Sahai and Marshall, 2003), it may be informative to investigate whether SEPT9 contributes to metastasis by modulating when and where blebs and protrusions occur or by enforcing particular motility modes.

**Conclusion**

Together, our findings provide mechanistic insight into earlier observations of the phenotypes of septin-deficient T cells and demonstrate the first in vivo evidence of the in vitro phenomenon of membrane contortion by septin complexes. Our data suggest that interactions with remodeling actin at membranes may be the dominant mechanism by which septins regulate the cortex of amoeboid cells. These observations add an important new molecular player to our understanding of how cellular protrusions are controlled and retracted. This mechanism of cortical control may have implications for the diversity of amoeboid motility modes and may inform our understanding of how septin dysregulation contributes to metastasis.
Materials and methods

Cell lines
The D10.G4 cell line was maintained in RPMI-1640 supplemented with 10% FCS, l-glutamine, penicillin, streptomycin, β-mercaptoethanol, and 10 U/ml human interleukin-2. As D10.G4 cells were used only for studies of motility and not activation, they were not periodically restimulated, as has been previously described (Tooley et al., 2009). Cells were maintained at a density <10^5 cells/ml. A subclone of the D10.G4 line was generated by electroporating cells with the SEPT6-GFP plasmid described in the next section, sorting cells using a cell sorter (MoFlo; Dako), and maintaining in the media previously described supplemented with 50 µg/ml Genetin for selection.

Plasmids and transfections
All transfections were performed by electroporation using Gene Pulser (Bio-Rad Laboratories). shRNA against Sept7 and control shRNA plasmids were used as previously described (Tooley et al., 2009). Cells were transfected with knockdown plasmids for Sept7 were cotransfected with EGFFP-N1 (Takara Bio Inc.) as a marker. SEPT6-GFP consists of the mouse Sept6 sequence with a 22-amino acid linker sequence added by PCR and cloned into EGFPC1 (Takara Bio Inc.). LifeAct-Ruby was a gift from G.S. Baron (Rocky Mountain Laboratories, National Institute of Allergy and Infectious Diseases, Hamilton, MT; Speare et al., 2010). LifeAct-Ruby consists of 17 amino acids from Saccharomyces cerevisiae Abp140 (Riedl et al., 2008) subcloned into pCMV-RFP-Ruby-N1 vector (Takara Bio Inc.) via EcoRI-BamHI sites.

Immunofluorescence
For observations of crawling cells, cells were allowed to adhere to poly-L-lysine or 0.2 µg/ml ICAM-1–coated coverslips for 30 min before fixation. For volume-change experiments, chambered coverslips (Thermo Fisher Scientific) were coated with 0.5 µg/ml anti-CD44 antibody (clone IM7) for 60 min and then washed three times. Next, cells were suspended in 150 µl of serum-deficient RPMI-1640 and allowed to adhere to coated coverslips for 30 min. 300 µl of either 37°C deionized water or 37°C RPMI-1640 was added to each chamber (for hypertonic and isotonic conditions, respectively). After these preparations, warmed 16% PFA (Electron Microscopy Sciences) was added to each chamber for fixation at a final concentration of 4% for 10 min. After thorough washing with PBS, fixed cells were blocked with 2% donkey serum and 2% FCS and permeabilized with 0.2% saponin (Sigma-Aldrich) in PBS for 30 min. Cells were incubated with rabbit anti-SEPT7 antibody (IBL-America, Inc.) for 120 min in permeabilization buffer, washed, and then stained with secondary antibodies (Jackson ImmunoResearch Laboratories, Inc.) in permeabilization buffer for 60 min and washed again. Samples were imaged immediately or stored at 4°C overnight for subsequent imaging.

Microscopy
Widefield images were acquired on a modified microscope (Axiovert 200M; Carl Zeiss) with a Plan-Neofluar 63x objective (Carl Zeiss). The microscope was fitted with dual excitation and emission filter wheels and a camera (CoolSNAP HQ; Roper Scientific). Image acquisition was performed using MetaMorph imaging software (Molecular Devices). For motility and mitosis imaging, a 0.25% low–melting point agarose base of actin-containing protrusions during RVD. Video 5 shows septin rings appearing and incubated at 37°C for 30 min before imaging.

Cell sorting
Cell sorting was performed on a cell sorter (MoFlow) or special order products (FACSria II; BD) using 70-µm nozzles.

Analysis and statistics
All flow cytometry analysis was performed in FlowJo (Tree Star), and the FlowJo kinetics platform was used for osmotic volume-change measurements. Images were analyzed in MetaMorph (Molecular Devices) or Imaris (Bitplane) software depending on the type of analysis. MetaMorph was used for 3D reconstructions. The mitotic blebbing index consists of the number of blebs observed divided by the product of the time followed (in seconds) and the circumference of the cell (in micrometers). All statistical analyses were performed in Prism software (GraphPad Software), and individual tests are named in the legends.

Online supplemental material
Fig. S1 shows protein levels of a panel of septins after SEPT7 knockdown. Fig. S2 shows the standard curve used for calibrating cell size in the flow cytometry volume-change assay. Video 1 shows the location of membrane protrusions relative to SEPT6-GFP in a crawling cell. Video 2 demonstrates cortical morphogenesis during mitosis of control and SEPT7KD cells, sequentially. Video 3 shows high-resolution time-lapse images of the formation of septin rings during RVD. Video 4 shows a 3D reconstruction of cells with septin rings. Video 5 shows septin rings appearing during RVD in a field of cells. Video 6 shows invaginations in the plasma membrane during RVD. Video 7 shows SEPT6-GFP distribution in cells treated with jasplakinolide. Video 8 shows septin accumulation at the base of actin-containing protrusions during RVD. Video 9 shows recruitment of SEPT6-GFP to blebs in a crawling dynasore-treated cell. Video 10 shows recruitment of LifeAct-Ruby to a bleb preceding SEPT6-GFP recruitment. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201105127/DC1.

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