Structural centrosome aberrations favor proliferation by abrogating microtubule-dependent tissue integrity of breast epithelial mammospheres

D Schnerch and EA Nigg

A large proportion of human carcinomas exhibit centrosome aberrations already during early stages of carcinogenesis.\(^\text{19–28}\) Centrosome aberrations have traditionally been subdivided into numerical and structural aberrations.\(^\text{24,29}\) Most numerical aberrations are characterized by centrosome amplification, most commonly caused by supernumerary centrioles that result from deregulated centriole duplication or division failure, whereas structural centrosome aberrations refer to altered shapes and compositions of the pericentriolar matrix with normal centriole numbers. Numerical aberrations and their consequences have been studied extensively.\(^\text{26–30}\) In contrast, the origin and role of structural centrosome aberrations has received comparatively little attention.\(^\text{37–39}\)

Here, we have explored the consequences of structural centrosome aberrations for the architecture and function of MCF10A breast epithelial cells grown in two-dimensional (2D) and three-dimensional (3D) tissue culture. We focused on structural centrosome aberrations induced by overexpression of Ninein-like protein (Nlp).\(^\text{40}\) This distant relative of Ninein localizes preferentially to mother centrioles and contributes to MT organization in interphase cells.\(^\text{40}\) Cell cycle regulation of Nlp expression and localization is required for mitotic spindle formation and accurate progression through mitosis.\(^\text{41–45}\) Intriguingly, Nlp is strongly overexpressed in various types of human cancer (including breast cancer),\(^\text{46–49}\) and Nlp overexpression is associated with enhanced proliferation as well as reduced sensitivity to anticancer treatment, which has been proposed to contribute to an adverse clinical course of disease.\(^\text{46,48,49}\) Supporting these findings, transgenic mice overexpressing Nlp spontaneously develop invasive breast cancer.\(^\text{47}\) Moreover, in 2D culture EGF-Nlp was previously shown to form striking centrosome-related bodies (CRBs)\(^\text{50}\) that closely resemble the structurally aberrant centrosomes seen in human cancers.\(^\text{33,36,37}\) Here, we demonstrate
that Nlp overexpression profoundly impacts on the MT organization of epithelial cells. In 3D culture mammospheres, this results in a drastic disruption of epithelial architecture and enhanced responsiveness to growth factor stimulation.

RESULTS
Overexpression of EGFP-Nlp prevents epithelium formation in 3D tissue culture

Structural centrosome aberrations in cancer cells are presumed to result from deregulated expression of centrosomal proteins.\textsuperscript{23,24,51} We therefore applied a retroviral system to study the effects of overexpression of selected centrosomal proteins in MCF10A breast epithelial cells (Supplementary Figure S1). Importantly, this experimental set-up made it possible to induce the overexpression of EGFP-tagged proteins at defined time points, allowing the initial morphogenesis of 3D mammospheres to remain unperturbed (Supplementary Figure S1). Analysis of a number of centrosomal proteins revealed distinct consequences of protein overexpression in 2D culture (Figure 1a). While some proteins localized primarily to centrosomes, others additionally filled up the cytoplasm or formed multiple cytoplasmic aggregates, as illustrated by the fates of Cep192, Cep68 or Cep135, respectively (Figure 1a). Most strikingly, the overexpression of Nlp caused large centrosome-associated patches, confirming and extending previous results.\textsuperscript{40,50} In the following, we refer to these structural centrosome aberrations as CRBs (Figure 1a).

As Nlp is frequently overexpressed in human cancers\textsuperscript{47} and considering that Nlp-induced CRBs closely resemble the structural centrosome aberrations seen in cancer tissue sections,\textsuperscript{37,39} we asked how overexpression of Nlp would affect the morphology of mammospheres. Generation of inducible MCF10A cell lines allowed maximal expression of EGFP-Nlp within 24–48 h of doxycycline induction, and similar kinetics of induction were observed for EGFP-Cep68 analyzed for control (Figure 1b). Importantly, the levels of Nlp achieved in these experiments were in a comparable range with those reported for human cancers.\textsuperscript{47}

Mammospheres were allowed to grow until they had reached maximal sizes with polarized epithelia, before overexpression of centrosomal proteins was induced at day 9 and the consequences of Nlp CRBs were studied 8 days later (Figure 1c and Supplementary Figure S1b). The proficiency of mammospheres to align their nuclei in a spherical layer provided a first convenient readout to distinguish aberrant from normal tissue architecture (Figure 1c). This analysis revealed that the overexpression of EGFP-Nlp abolished the proper basal alignment of nuclei in most mammospheres, whereas the overexpression of EGFP-Cep68 did not detectably disrupt mammosphere architecture (Figure 1c). As expected, the penetration of the Nlp-induced phenotype correlated with the extent of overexpression (Supplementary Figures S1c and d).

To more directly address the impact of Nlp CRBs on epithelial polarity, we also probed mammospheres for the basement membrane component laminin5 and the Golgi protein GM130, markers for basal and apical (luminal) compartments, respectively. As shown in Figure 1d, Nlp CRBs caused a major disruption of epithelial polarity, whereas overexpression of Cep68 did not. The consequences of Nlp CRBs on tissue architecture could also be seen in chimeric clusters of mammospheres where parts expressing little or no transgene were located immediately adjacent to cells expressing high levels of EGFP-Nlp (Figure 1e). While gland-like structures were preserved in the former parts of the chimeras, the latter parts exhibited a striking disruption of tissue architecture, reminiscent of tumors (Figure 1e).

To determine the specificity of the phenotype observed upon overexpression of Nlp in 3D mammospheres, we next studied the consequences of overexpression of other centrosomal proteins. When examined in the exact same assay as used for Nlp, none of the other proteins listed in Figure 1a produced major effects on mammosphere architecture, although minor effects were seen sporadically (Supplementary Figure S2). These results not only attest to the specificity of the Nlp phenotype, but also support the hypothesis that Nlp overexpression in human tumors may causally contribute to carcinogenesis.\textsuperscript{47} In light of a recent study reporting the induction of an invasive phenotype in response to centrosome amplification,\textsuperscript{35} we were surprised that overexpression of the centrosome duplication factors Plk4 or Sas-6 did not detectably induce invasive growth or major perturbations of epithelial cell polarity, although both proteins clearly induced centrosome amplification (Supplementary Figure S2).

Nlp CRBs alter the organization of the MT cytoskeleton
To determine the mechanisms by which excess Nlp interferes with epithelial cell organization, we examined the impact of Nlp CRBs on centriole numbers, MT organization and MT-dependent processes. No evidence was obtained for aberrations in centriole numbers upon overexpression of Nlp (data not shown), in line with previous reports.\textsuperscript{50} However, both γ-tubulin and centrin-2/3 were recruited to Nlp CRBs in both 2D and 3D tissue culture (Figures 2a and b), confirming and extending previous results.\textsuperscript{40} In these respects, Nlp CRBs are highly reminiscent of the large γ-tubulin and centrin staining structures observed in sections of breast adenocarcinoma.\textsuperscript{37,38} To explore the impact of γ-tubulin recruitment on MT organization, we carried out MT regrowth assays in MCF10A cells (Figures 2c and d). When MTs were allowed to regrow for 10 min, Nlp CRB-containing cells generally displayed denser MT asters than controls (Figure 2c), seemingly in line with the increased MT nucleation activity of structurally aberrant centrosomes in permeabilized tumor sections.\textsuperscript{37,39} However, during early phases of MT regrowth, notably after 1 min, we observed that Nlp CRBs caused a reduction in MT nucleation kinetics when compared with Cep68-expressing or control cells (Figure 2d). This surprising result raised the question of how to reconcile initial suppression of MT nucleation activity by excess Nlp with higher density of MT asters at steady state.

The density of centrosomal MT asters at steady state is determined not only by the rate of MT nucleation, but also by the rate of MT release. Thus, we explored the possibility that excess Nlp might conceivably suppress both MT nucleation and release. To this end, we examined the localization of Ninein, one of the key components regulating MT release from centrosomes.\textsuperscript{16,17} Interestingly, Ninein was clearly recruited to Nlp CRBs in both 2D and 3D cultures (Figure 2e). As the centrosomal accumulation of Ninein was previously shown to abrogate MT release from centrosomes,\textsuperscript{18} these results indicate that the concomitant accumulation of Nlp and Ninein reduces the rates of both MT nucleation and release. Assuming that the latter effect prevails over time, this readily explains the high MT density observed around Nlp CRBs (see also Casenghi et al.\textsuperscript{49}).

To corroborate the notion that accumulation of Ninein has a major role in tethering MTs to Nlp-induced CRBs, we sought a way to determine the impact of Ninein mislocalization on CRB structure and MT organization. We took advantage of a previous study demonstrating that the overexpression of a C-terminal Ninein fragment (Supplementary Figure S3a) acts in a dominant-negative manner and causes the mislocalization of endogenous Ninein.\textsuperscript{15} High levels of C-terminal Ninein fragment indeed caused a strong reduction in centrosomal Ninein signals in control cells, confirming the functionality of the expression construct (Supplementary Figure S3b). Moreover, when the same C-terminal Ninein fragment was expressed in cells overexpressing EGFP-Nlp, we observed a marked loss of Ninein from Nlp-induced CRBs (Supplementary Figure S3c). Concomitantly, CRBs underwent fragmentation (Supplementary Figures S3c and d) and MT networks lost their focus on CRBs (Supplementary Figure S3d).
Figure 1. Nlp CRBs interfere with apical basal polarity in 3D tissue culture. (a) Localization of overexpressed EGFP-tagged centrosome proteins in 2D culture, as visualized by fluorescence microscopy. (b) Western blots indicate levels of overexpression of EGFP-tagged centrosome proteins after induction (Dox) over time. (c) Top panel: schematic illustrating experimental design for protein induction in 3D culture; bottom panel: DAPI staining of mammospheres, illustrating proficiency or deficiency in basal alignment of nuclei. See also Supplementary Figure S1. (d) Analysis of mammospheres expressing EGFP-Nlp or EGFP-Cep68 by immunofluorescence microscopy. EGFP-Nlp and EGFP-Cep68 are shown in green, Laminin5 and GM130 in red, and DNA in blue (DAPI). See also Supplementary Figure S2. (e) Chimeric mammosphere cluster, illustrating the impact of Nlp CRBs on 3D architecture. Staining as in b. Asterisks indicate parts of the cluster showing glandular morphology and basal laminin5 staining.
Together, these data support the view that Ninein accumulation contributes to the characteristic shape and properties of Nlp-induced CRBs.

To investigate the functional consequences of the altered dynamics in MT organization induced by Nlp CRBs, we next examined the ability of the MT cytoskeleton to reorganize in...
response to monolayer wounding. This assay induces rapid cell polarization towards the wound edge, providing an opportunity to monitor MT re-organization between leading and lagging cell compartments (for definition, see Figure 2f). In line with the observation that centrosomes of epithelial cells do not generally re-orient towards the wound edge,52 108 out of 120 control MCF10A cells at the wound retained their centrosomes in the lagging compartment (data not shown). Yet, in both control and Cep68-expressing cells, the bulk of MTs redistributed to the leading compartment (Figure 2f). In striking contrast, this reorganization of the MT cytoskeleton in response to wounding was markedly suppressed in cells harboring Nlp CRBs (Figure 2f).

This strongly supports our conclusion that excess Nlp suppresses the release of MTs from centrosomes and demonstrates that Nlp CRBs prevent the reorganization of the MT network in response to cell polarization.

Nlp CRBs affect mammosphere architecture
Having established a profound effect of Nlp CRBs on the dynamics of the MT cytoskeleton in 2D culture, we considered it important to extend these results to polarized single cell layers of epithelium in 3D culture. Mammospheres harboring Nlp CRBs lost the prominent enrichment of MTs near the outer confinement and instead showed higher MT densities near the lumen, demonstrating major reorganization of the MT cytoskeleton also in 3D culture (Figure 3a). However, because mammospheres progressively change their architecture in response to Nlp CRBs, it was difficult to analyze the MT phenotype in individual cells. To circumvent this problem, we induced Nlp expression at different times during 3D culture and compared the kinetics of alterations in the MT cytoskeleton relative to the onset of architectural alterations. We indeed found conditions that induced detectable reorganization of the MT cytoskeleton concomitant with CRB formation but prior to major alterations in tissue architecture (Supplementary Figure S4; shaded panel). This made it possible to study the impact of CRBs on the MT cytoskeleton of individual cells (Figure 3b and Supplementary Figure S4). In line with the data obtained in 2D cultures, cells harboring Nlp CRBs typically exhibited a higher density of α-tubulin in the apical, CRB-containing compartments, adjacent to the lumen of the mammosphere (Figure 3b and Supplementary Figure S4).

Tumors are generally thought to originate from individual founder cells within otherwise healthy tissues. To phenocopy this condition in mammospheres and study the behavior of isolated CRB-harboring cells in an otherwise unperturbed epithelial structure, whole mammospheres were incubated with lentiviral particles to stably integrate expression cassettes encoding EGFP-Nlp into the genome of single cells. This approach led to the generation of mammospheres containing isolated cells harboring Nlp CRBs.

Importantly, these Nlp CRB-containing cells characteristically showed an intense α-tubulin staining adjacent to the CRB in the apical compartment near the lumen, in stark contrast to normal bystander cells (Figure 3c). This indicates that the localized emergence of structural centrosome aberrations is sufficient to trigger a reorganization of the MT cytoskeleton, which in turn exerts a profound impact on epithelial architecture.

Nlp CRBs interfere with the positioning of basal adhesion and lateral junction proteins
Epithelial cell biology is characterized by two key aspects. First, robust adhesion to the basement membrane is essential for line-up of polarized epithelial cells and maintenance of epithelial architecture.1,53 Second, tight intercellular connections between neighboring cells are critical for proper lateral adhesion.1 This prompted us to ask to what extent the reduced density of MTs in the basal compartment of cells harboring Nlp CRBs affects the positioning of basal adhesion and lateral junction proteins. While mammospheres formed from cells expressing excess EGFP-Cep68 exhibited proper α6-integrin localization, this basal adhesion component was partially lost from the basal compartment in mammospheres expressing Nlp CRBs (Figures 4a and c), highly reminiscent of early stage human breast cancer.54 Likewise, the ligand of α6-integrin, laminin5, was largely lost from the basal membrane in Nlp CRBs-expressing mammospheres but normally distributed in Cep68 controls (Figures 4b and c; see also Figures 1d and e). Furthermore, the extent to which these basal adhesion proteins were lost from the basal cortex clearly correlated with the extent to which Nlp CRB formation disrupted tissue architecture (Figure 4c). To determine whether Nlp CRBs also perturb the localization of lateral junction proteins, we probed for E-Cadherin, an important mediator of junction belt initiation.1 While E-Cadherin decorated the lateral boundaries between neighboring cells in control and EGFP-Cep68-overexpressing mammospheres, those harboring Nlp CRBs clearly showed more dispersed and irregular distribution of E-Cadherin (Figure 4d). In addition to being important for junction belt organization, E-Cadherin also recruits the polarity protein and tumor suppressor Scribble to the basolateral membrane.55 As Scribble is frequently lost or mis-localized in breast cancer,55-58 we next examined the consequences of Nlp CRBs on Scribble localization. Consistent with the results obtained for E-Cadherin, mammospheres harboring Nlp CRBs displayed a profoundly dispersed and irregular Scribble staining pattern when compared with controls (Figure 4e). Taken together these results demonstrate that structural centrosome aberrations, as triggered here by excess Nlp, impact not only the MT cytoskeleton but also profoundly disturb key proteins implicated in epithelial adhesion and cell-cell interactions.

Figure 2. Nlp CRBs anchor a surplus of MTs. (a, b) Sequestration of γ-tubulin (a) and centrin (b) to Nlp CRBs in 2D (upper panels; plus/minus Dox) and 3D cultures (lower panels; right hand panels show magnification of boxed area). γ-Tubulin and centrin in red, EGFP-Nlp in green and DNA in blue (DAPI). (c, d) MT regrowth in cells expressing Nlp or Cep68, analyzed for control, before and after protein induction (Dox). Upper panels show merged data in pseudo-color (α-tubulin in red, EGFP-Nlp and EGFP-Cep68 in green, and DNA in blue (DAPI)); arrowheads indicate the location of the centrosome/CRB. Lower panels show α-tubulin staining in grey; dashed ring illustrates MT density in proximity to centrosome/CRB. Histogram to the left shows quantification of mean extension of the MT cytoskeleton. Histogram to the right shows quantification of mean diameter of MT asters; n indicates sample size; data are from three independent experiments (N = 3); error bars indicate standard deviation and P-values were derived from unpaired, two-tailed Student’s t-test. (*) indicates P-value < 0.05, (**) < 0.01 and (***) < 0.001. (e) Sequestration of Ninein to Nlp CRBs in 2D (upper panels; plus/minus Dox) and 3D cultures (lower panels; right hand panels show magnification of boxed area). Ninein in red, EGFP-Nlp in green and DNA in blue (DAPI). (f) Impact of Nlp CRB on MT reorganization in MCF10A cells in response to monolayer wounding-induced cell polarization by immunofluorescence microscopy. Pseudo-colors show α-tubulin in red, γ-tubulin in yellow, EGFP-Nlp in green and DNA in blue (DAPI) (left panel); cells were compartmentalized (white dashed and dotted lines) to monitor MT reorganization, as illustrated schematically in the upper right panel (dashed lines delineate the leading compartment, dotted lines the lagging compartment). Histogram in the lower right panel shows quantities of the ratios of MT fluorescence intensities between the leading and lagging compartments; n indicates sample size; data are from three independent experiments (N = 3); error bars indicate standard deviation and P-values were derived from unpaired, two-tailed Student’s t-test. (*) indicates P-value < 0.05, (**) < 0.01 and (***) < 0.001.
Deregulation of epithelial architecture by Nlp CRBs increases responsiveness to growth stimulation. Loss or aberrant localization of Scribble has been reported to enhance cell proliferation in response to epidermal growth factor (EGF). This prompted us to speculate that Nlp CRBs might disrupt not only MT networks and epithelial organization, but additionally confer a growth advantage to mammospheres. To explore this hypothesis, we examined the activation state of the
MAP kinase pathway, as monitored by phospho-ERK staining (Figure 4f), as well as the proliferative response of mammospheres to EGF (Figure 4g). We found that cells harboring Nlp CRBs displayed a more intense phospho-ERK staining in response to growth factor stimulation than controls expressing Cep68 (Figure 4f). Moreover, the surplus in MAP kinase pathway activity clearly translated into increased proliferation of cells harboring Nlp CRBs. While increasing doses of EGF had only a moderate effect on mammospheres expressing Cep68 (Figure 4g), mammospheres harboring Nlp CRBs showed a clear dose response and grew to progressively larger sizes (Figure 4g, right panel). Remarkably, differential responses to high doses of EGF could be seen even within single mammospheres. Regions with high Nlp expression showed marked expansion, while parts characterized by low expression showed no significant response (Figure 4g). Finally, when grown in the presence of a high dose of EGF, mammospheres harboring Nlp CRBs also showed increased numbers of cells staining positive for the proliferation marker Ki-67 (Figure 4h).

DISCUSSION

Structural centrosome aberrations are common in human tumors, but their impact on the biological properties of tumor cells is poorly understood. To explore the mechanistic consequences of structural centrosome aberrations on epithelial organization, we have developed a 3D culture model based on overexpression of the centrosomal protein Nlp in mammospheres derived from non-transformed MCF10A cells. We show that overexpression of Nlp results in the formation of CRBs, highly reminiscent of the structural centrosome aberrations described in human cancers.37,38 Furthermore, we demonstrate that Nlp CRBs trap MT minus ends through sequestration of the MT release factor Ninein. This in turn affects the organization of the MT cytoskeleton as well as the localization of adhesion and junction proteins, resulting in a profound disruption of apical basal polarity. Finally, we show that the Nlp CRB-induced alterations in epithelial architecture favor sustained proliferation in response to growth factor stimulation. Collectively, these results demonstrate that aberrant expression of a single centrosomal protein in otherwise non-transformed epithelial cells induces profound structural centrosome aberrations. These in turn confer properties to 3D mammospheres that closely resemble typical features of human carcinoma cells.

The Nlp CRB model analyzed here shares several important properties with CRBs that are commonly observed in sections of primary human tumor tissue, notably their altered shape and size and the striking accumulation of both γ-tubulin and centrin.37,38 Moreover, we emphasize that the centrosomal protein Nlp, used here to induce CRBs, is frequently overexpressed in human cancers.47 Although it is difficult to directly compare expression levels between different samples, we emphasize that the extent of Nlp overexpression in MCF10A cells (Figure 1b) appears comparable with that observed in tumor material.47 Previously, CRB formation has been correlated with an increase in the abundance of MTs associated with these structures, both in cultured cells40 and in sections of primary tumor samples.37,39 However, although our data confirm that CRB formation results in increased MT density, presumably reflecting enhanced MT stability at steady state, MT regrowth assays revealed that excess Nlp exerts a suppressive effect on MT nucleation. This apparent paradox could be resolved by showing that Nlp CRBs also sequester Ninein, which then suppresses MT release,18 resulting in a net increase of MT accumulation at CRBs over time. These observations can explain how structural centrosome aberrations, induced here by Nlp, can trap MT minus ends and thereby impair the ability of the MT network to reorganize in polarizing cells. This in turn has profound consequences for the organization of epithelial cell layers.

In a recent study, the overexpression of the centrosome duplication factor Plk460,61 was reported to induce invasive growth of epithelial cells.35 As this work was performed using a 3D culture assay similar to the one used here, we were initially surprised by the apparent lack of phenotype associated with overexpression of either Plk4 or Sas-6 in our mammospheres. Although both of these proteins produced the expected centriole amplification, we saw no evidence for mammosphere deformation indicative of invasive growth. When interpreting these negative results, it is important to consider that mammosphere formation involves different stages.62 While we generally allowed mammospheres to reach maximal size before inducing centrosomal proteins at day 9, Godinho and coworkers35 induced Plk4 overexpression before seeding of cells into matrigel, resulting in cells harboring supernumerary centrosomes prior to 3D expansion and initial polarization.62 Furthermore, we note that Godinho and coworkers cultured mammospheres in the presence of type I collagen, which is known to enhance the stiffness of matrigel preparations and sensitize the assay for detection of invasive growth.63 Thus, we consider it likely that our inability to detect invasive growth in response to centrosome amplification reflects the above differences in experimental design and/or assay conditions.

Interestingly, the proficiency of epithelial cells to generate non-centrosomal MTs correlates with differentiation state.64 While cells residing in stem cell compartments mostly exhibit MT asters emanating from the centrosome, differentiated epithelial cells as well as neurons display an abundance of non-centrosomal MTs.64-66 Release from centrosomes and vectorial transport of MT minus ends depends on Ninein16 and, as we show here, this process is suppressed by Nlp CRBs. We emphasize that structural centrosome aberrations are not the only cause for aberrant MT

Figure 3. Nlp CRBs alter the organization of MTs in polarized epithelia of mammospheres. (a) Impact of Nlp CRBs on MT organization in mammospheres. Micrographs show juxtaposition of two mammospheres, only one of which expresses Nlp in response to Dox. Left panel: merge of pseudo-colors illustrating EGFP-Nlp in green, α-tubulin in red and DNA in blue (DAPI). Middle and right panels: single stainings for α-tubulin highlight disruption of MT organization and concomitant misalignment of nuclei, respectively. (b) Polarization of MT reorganization in mammosphere epithelia at single cell level. Top left panel: schematic illustrating experimental design for protein induction in 3D culture; bottom left panel: micrographs show impact of Nlp CRBs on MT organization (center). Panels show overviews (above) and magnifications of box areas (below). Staining as in a. Right panels: schematic illustrates compartmentalization of polarized mammosphere epithelial cells into apical (luminal) and basal compartments. Histogram shows quantification of ratio of α-tubulin fluorescence intensities in the two compartments (n indicates sample size). Data were obtained from all cells shown in the micrographs of panel (b); virtually identical results were obtained in three independent experiments. See also Supplementary Figure S3. (c) Reorganization of MT cytoskeleton in single cells harboring Nlp CRBs in otherwise unperturbed mammospheres. Viral infections were carried out as indicated in the cartoon to the left. Micrographs to the right illustrate the impact of Nlp CRB on MT organization. Staining in left hand panels as in a, while right hand panels show α-tubulin staining alone (grey).
organization in cancer cells and several important tumor suppressors, including adenomatous polyposis coli, RASSF1A, VHL, E-Cadherin and merlin, were reported to perturb MT networks during early carcinogenesis. However, whereas mutational inactivation of a tumor suppressor generally requires two genetic hits, upregulation of Nlp (or other centrosomal proteins conferring similar structural centrosome aberrations) is expected to exert a dominant effect in response to a single genetic event. The functionality of epithelia depends on their ability to form uniformly polarized layers of differentiated cells. From the perspective of carcino genesis, it is important that the formation of epithelial cell-cell interactions have been recognized as an important barrier against the outgrowth of transformed single cells (reviewed in Fujita, Hogan and Yamauchi and Fujita). Conversely, detachment from the epithelial layer and intraluminal proliferation of cells constitutes a hallmark of early breast cancer. Concomitantly, the loss of polarity that accompanies the evasion from the epithelial sheet abolishes the segregation of growth factor receptors and their activators into distinct membrane domains, allowing promiscuous interactions. Our present study showed that perturbation of the MT network through Nlp CRBs triggers a profound disruption of epithelial integrity, as illustrated by mislocalization of α6-integrin, laminin5 and E-Cadherin. Moreover, the tumor suppressor Scribble, normally associated with basolateral membranes, was found to be dispersed in cells harboring Nlp CRBs. In consideration of data showing mislocalization and downregulation of Scribble in early breast cancer and reports linking Scribble depletion to the MAP kinase pathway, this observation prompted us to explore the response of mammospheres to growth factor stimulation. Strikingly, when mammospheres were treated with EGF, we observed that structures harboring Nlp CRBs grew to larger sizes than controls. Moreover, the extent of proliferative response was proportional to EGF concentration. These results support the view that structural centrosome aberrations disrupt not only epithelial architecture but also favor cell expansion in response to proliferative stimuli.

In conclusion, our results demonstrate that structural centrosome aberrations exert a profound impact on the biological properties of developing tumor cells, indicating that they deserve as much attention in future studies as numerical centrosome aberrations.

MATERIALS AND METHODS

Cell culture

MCF10A ecor cells, authenticated by Short Tandem Repeat profiling, were a kind gift from Tilman Brummer (University of Freiburg) and grown in Dulbecco’s modified Eagle’s medium supplemented with 10% fetal calf serum, sodium pyruvate and Pen/Strep. Tissue cultures were routinely tested for mycoplasma contamination by PCR using growth medium from high density cultures as a template. Doxycycline-inducible MCF10A ecor cell lines allowing expression of selected EGFP-tagged centrosomal proteins were generated as illustrated in Supplementary Figure S1a. For antibiotic selection, hygromycin and puromycin were used at concentrations of 100 μg/ml and 2 μg/ml, respectively. Doxycycline was used at a final concentration of 2.5 μg/ml. For generation of mammospheres in 3D culture, MCF10A ecor cells were plated and grown in matrigel using the overlay method as described previously.

Generation of retroviral and lentiviral expression constructs

The retroviral vector pBabe rtTA Hygro was kindly provided by Cheuk Leung (University of Minnesota). Coding sequences of EGFP-Nlp, EGFP-Cep68, EGFP-α-tubulin, EGFP-Cep135, EGFP-Cep192, EGFP-PIK4 and EGFP-Sas-6 were PCR-amplified and ligated into pPMD2.GutPur (Clontech, Mountain View, CA, USA). The lentiviral vector pLEGO-iC2 was obtained from Boris Fehse via Addgene (Addgene plasmid #27345; Cambridge, MA, USA). For the generation of pLEOG EGF-Nlp-iC2, the sequence encoding EGF-Nlp was PCR-amplified and ligated into LeGO-iC2. For the generation of pLEOG EGF-Nlp-iC2, the sequence encoding Nininein (1885-2135) was PCR-amplified and ligated into LeGO-iC2. For the generation of pLEOG Nininein, the sequence encoding Nininein (1885-2135) was PCR-amplified and ligated into LeGO-iC2. For the generation of pLEOG Nininein, the sequence encoding Nininein (1885-2135) was PCR-amplified and ligated into LeGO-iC2.

Fluorescence microscopy

Immunolabeling of 2D cultures was performed using standard fixation and labeling protocols, using ice-cold methanol. Procedures for staining of 3D mammospheres were adapted from previously published protocols. For detection of γ-tubulin, centrin and phospho-p44/42 MAPK mammospheres were fixed using ice-cold methanol (15 min). Primary antibodies used in this study were anti-α-tubulin (T9026, Sigma, St Louis, MO, USA), anti-γ-tubulin (for 2D: R60; 80 for 3D: T-6557, Sigma), anti-α-integrin (RJ60, Sigma), anti-λ-integrin (ab27076, Abcam, Cambridge, MA, USA), anti-Sas-6 Cy5, anti-PI-110-Cys, anti-Ninein (R154), anti-Centrin2/3, anti-Laminin V (MAB19562) and anti-E-Cadherin (610181) (both from BD, Franklin Lakes, NJ, USA), anti-Scribble (sc-11049, Santa Cruz Biotechnology, Santa Cruz, CA, USA), anti-Ki-67 (18-0191Z, Invitrogen, Germany), anti-GM130 (610822) and anti-E-Cadherin (610181) (both from BD, Franklin Lakes, NJ, USA), anti-Scribble (sc-11049, Santa Cruz Biotechnology, Santa Cruz, CA, USA), anti-Ki-67 (18-0191Z, Invitrogen, Germany), anti-GM130 (610822) and anti-E-Cadherin (610181) (both from BD, Franklin Lakes, NJ, USA), anti-Scribble (sc-11049, Santa Cruz Biotechnology, Santa Cruz, CA, USA), anti-Ki-67 (18-0191Z, Invitrogen, Germany), anti-GM130 (610822) and anti-E-Cadherin (610181) (both from BD, Franklin Lakes, NJ, USA), anti-Scribble (sc-11049, Santa Cruz Biotechnology, Santa Cruz, CA, USA), anti-Ki-67 (18-0191Z, Invitrogen, Germany), anti-GM130 (610822) and anti-E-Cadherin (610181) (both from BD, Franklin Lakes, NJ, USA).

Figure 4. Nlp CRBs impair localization of basal and lateral adhesion proteins and confer enhanced responsiveness to EGF stimulation. (a, b) Impact of Nlp CRBs on basal localization of α6-integrin (a) and laminin5 (b) by immunofluorescence (IF) microscopy. Pseudocolors show EGFP-Nlp- and EGFP-Cep68 in green, α6-integrin and laminin5 in red, and DNA in blue (DAPI). (c) Panels to the left show merge of pseudocolors, panels to the right show single stainings in grey for α6-integrin (above) and laminin5 (below). Overviews and blow-ups illustrate correlation between Nlp CRBs and displacement of basal adhesion proteins. (d, e) Impact of Nlp CRBs on basal localization of α6- and β1-integrin (d) and Scribble (e) by IF microscopy. E-Cadherin and Scribble in red, EGFP-Nlp and EGFP-Cep68 in green, and DNA in blue (DAPI). (f, g, h) Nlp CRBs confer enhanced responsiveness to EGF. EGF treatment of mammospheres, as illustrated schematically in f, enhances p-ERK1/2 staining (f) and, concomitantly, cause mammospheres to grow to larger size (g). Representative micrographs in g show effect of EGF on the proliferative expansion of mammospheres harboring Nlp CRBs; EGFP-Nlp in green, DNA in blue (DAPI); arrowhead points to control mammosphere showing minimal Nlp expression and asterisk marks part of chimeric mammosphere that fails to express EGFP-Nlp. Histogram to the right shows quantification of dose-dependent response to EGF by measurement of equatorial sections (illuminated in red; clashed plane in green). Representative micrographs in h show Ki-67-positive cells in mammosphere harboring Nlp CRBs next to a non-expressing mammosphere; upper micrograph shows merge of EGFP-Nlp in green, DNA in blue (DAPI) and Ki-67 in red. Lower micrograph shows Ki-67 in greyscale. n indicates sample size; data are from three independent experiments (N = 3); error bars indicate standard deviation and P-values were derived from unpaired, two-tailed Student’s t-test. (*) indicates P-value < 0.05, (**) < 0.01 and (***) < 0.001.
anti-mouse (A21236), AlexaFluor647 donkey anti-goat (A21447) and AlexaFluor647 goat anti-rat (A21247). Immunofluorescence microscopy on 2D cultures was performed using an Applied Precision DeltaVision system (GE Healthcare, Little Chalfont, UK) equipped with a ×20/0.75 U Plan Apo objective as well as ×60/1.2 Plan Apo N and ×100/1.4 UPLS Apo oil immersion objectives. Alternatively, analyses were performed using a FEI company wide-field system (FEI Munich, Graefelfing, Germany) equipped with a ×40/0.95 U Plan Apo objective and a ×60/1.49 TIRF APON oil immersion objective. Confocal images were acquired using a Leica SPS-IL-MATRIX point-scanning confocal microscope (Leica Microsystems, Wetzlar, Germany) equipped with ×10/0.30 HC PL Fluotar and ×20/0.70 HCX Plan Apo CS air objectives and a ×63/1.40-1.60 HCX Plan Apo lambda blue oil immersion objective. A 488-nm diode laser light was applied for DAPI staining, 488-nm Argon laser light for visualization of EGFP and 633 nm HeNe laser light for visualization of AlexaFluor647 stainings. Image analyses and final adjustments were carried out in Omero 5.0.2.

Biochemical and cell biological techniques
For western blotting, cell pellets were lysed in Tris lysis buffer (20 mM Tris-HCl (pH 7.4), 150 mM NaCl, 0.5% IGEPAL CA-630) with protease and phosphatase inhibitors, before 20 or 50 μg of total protein were separated by sodium dodecyl sulfate–polyacrylamide gel electrophoresis and transferred onto polyvinylidene fluoride membranes. These were incubated with primary antibodies diluted in phosphate-buffered saline/5% skim milk powder (w/v). Primary antibodies were anti-human Nlp (R110), anti-GFP (ab290, Abcam) and anti-α-tubulin (DM1A (T9026), Sigma). Horse radish peroxidase-conjugated secondary antibodies were obtained from Bio-Rad (170-6515 and 170-6516, Hercules, CA, USA).

MT regrowth assays were performed essentially as described previously. In brief, MCF10A eoor cells allowing inducible overexpression of centrosomal proteins were seeded on ibidi eight-well microscopy slides (ibidi GmbH, Planegg-Martinsried, Germany) and grown in the absence or presence of doxycycline. MTs were depolymerized by cold treatment (30 min), before MT regrowth was triggered by addition of warm (37°C) medium for 1 to 10 min, and cells were fixed using ice-cold methanol. Monolayer woundng assays were performed essentially as described. In brief, 100,000 MCF10A cells were cultured to confluence on ibidi eight-well microscopy slides, before wounding using a plastic pipette tip. After 24 h, cells were fixed with ice-cold methanol and examined by immunofluorescence microscopy.

Statistical analyses
Statistical analyses were carried out using Microsoft Excel 2010. Sample sizes (n) were chosen to allow the detection of statistically significant differences between subgroups within biological replicates (N). Inclusion and exclusion of samples was carried out based on viability and typical appearance of the negative control sample, which was run side by side with the test sample. Standard deviation and variance were calculated and served as estimates of variation within each group of data. For statistical comparison of groups with similar variance, a homoscedastic t-test was performed. For statistical comparison of groups with diverging variance, a heteroscedastic t-test was applied. A statistically significant effect was assumed for P-values < 0.05.

CONFLICT OF INTEREST
The authors declare no conflict of interest.

ACKNOWLEDGEMENTS
We thank Tilman Brummer (University of Freiburg), Stefan Zimmermann and Ralph Wasch (University Medical Center Freiburg) as well as Heike Brinkhaus and Mohamed Bentires-Alj (FMI Basel) for cell lines and technical advice, and Cheuk Leung (University of Minneapolis) as well as Yi-Ren Hong (Kaohsiung Medical University, Taiwan) for viral vectors. We also thank Alecay Lovinton-Ferrand and other members of the Imaging Core Facility (IMCF) for help and Elena Nigg-Martignoni for technical assistance. Finally, we are grateful to Felix B. Engell and the ‘Emerging fields initiative’ (EFI) for ‘Cell Cycle in Disease and Regeneration (CYDER)’ from the Friedrich-Alexander-Universität Erlangen-Nürnberg (FAU) for constructive comments and support, as well as to all members of the Nigg lab for helpful discussions. This work was supported by the University of Basel and the Swiss National Science Foundation (310030B_149641).

AUTHOR CONTRIBUTIONS
DS performed all experiments. DS and EAN conceived experiments, analyzed the data and wrote the manuscript.

REFERENCES
1 Rodríguez-Boulan E, Macara IG. Organization and execution of the epithelial polarity programme. Nat Rev Mol Cell Biol 2014; 15: 225–242.
2 Jemal A, Siegel R, Ward E, Hao Y, Xu J, Murray T et al. Cancer statistics, 2008. CA Cancer J Clin 2008; 58: 71–96.
3 Lee M, Vasioukhin V. Cell polarity and cancer–cell and tissue polarity as a non-canonical tumor suppressor. J Cell Sci 2008; 121: 1141–1150.
4 Musch A. Microtubule organization and function in epithelial cells. Traffic 2004; 5: 1–9.
5 Meng W, Mushika Y, Ichii T, Takeichi M. Anchorage of microtubule minus ends to adherens junctions regulates epithelial cell-cell contacts. Cell 2008; 135: 948–959.
6 Hotta A, Kawakatsu T, Nakatani T, Sato T, Matsui C, Sukenzane T et al. Laminin-based cell adhesion anchors microtubule plus ends to the epithelial basal cell cortex through LSLalpha/beta. J Cell Biol 2010; 189: 901–917.
7 Azimzadeh J, Bornens M. Structure and duplication of the centrosome. J Cell Sci 2007; 120: 2139–2142.
8 Nigg EA, Stearns T. The centrosome cycle: Centriole biogenesis, duplication and inherent asymmetries. Nat Cell Biol 2011; 13: 1154–1160.
9 Brito DA, Gouveia SM, Bettencourt-Dias M. Deconstructing the centrole: structure and number control. Curr Opin Cell Biol 2012; 24: 4–13.
10 Goczcy P. Towards a molecular architecture of centriole assembly. Nat Rev Mol Cell Biol 2012; 13: 425–435.
11 Luders J, Stearns T. Microtubule-organizing centres: a re-evaluation. Nat Rev Mol Cell Biol 2007; 8: 161–167.
12 Andersen JS, Wilkinson CJ, Mayor T, Mortensen P, Nigg EA, Mann M. Proteomic characterization of the human centrosome by protein correlation profiling. Nature 2003; 426: 570–574.
13 Jakobsen L, Vanselow K, Skogs M, Toyota Y, Lundberg E, Poser I et al. Novel asymetrically localizing components of human centrosomes identified by complementary proteomics methods. EMBO J 2011; 30: 1520–1533.
14 Kollman JM, Merdes A, Mourey L, Agard DA. Microtubule nucleation by gamma-tubulin complexes. Nat Rev Mol Cell Biol 2011; 12: 709–721.
15 Delgehyr N, Sillibourne J, Bornens M. Microtubule nucleation and anchoring at the centrosome are independent processes linked by ninine function. J Cell Sci 2005; 118: 1565–1575.
16 Mogensen MM, Malik A, Piel M, Bouckson-Castraing V, Bornens M. Microtubule minus-end anchorage at centrosomal and non-centrosomal sites: the role of ninine. J Cell Sci 2000; 113: 3013–3023.
17 Moss DK, Bellett G, Carter JM, Liovic M, Keynton J, Prescott AR et al. Ninine is released from the centrosome and moves bi-directionally along microtubules. J Cell Sci 2007; 120: 3064–3074.
18 Abal M, Piel M, Bouckson-Castraing V, Mogensen M, Sibartia JB, Bornens M. Microtubule release from the centrosome in migrating cells. J Cell Biol 2002; 159: 711–737.
19 Guo HQ, Gao M, Ma J, Xiao T, Zhao LL, Gao Y et al. Analysis of the cellular centrosome in fine-needle aspirations of the breast. Breast Cancer Res 2007; 9: R48.
20 Jiang F, Caraway NP, Sabichi AL, Zhang HZ, Sabichi AL, Zhang HZ, Ruitrok A, Grossman HB et al. Centrosomal abnormality is common in and a potential biomarker for bladder cancer. Int J Cancer 2003; 106: 661–665.
21 Kayser G, Gerlach U, Walch A, Nitschke R, Haxelmann S, Kayser K et al. Numerical and structural centrosome aberrations are an early and stable event in the adenoma-carcinoma sequence of colorectal carcinomas. Virchows Arch 2005; 447: 61–65.
22 Kronenwett U, Huwendiek S, Castro J, Ried T, Auer G. Characterisation of breast fine-needle aspiration biopsies by centrosome aberrations and genomic instability. Br J Cancer 2005; 92: 389–395.
23 Nigg EA. Centrosome aberrations: cause or consequence of cancer progression? Nat Rev Cancer 2002; 2: 815–825.
24 Nigg EA. Origins and consequences of centrosome aberrations in human cancers. Int J Cancer 2006; 119: 2717–2723.
25 Pihan GA, Wallace J, Zhou Y, Dowsey SJ. Centrosome abnormalities and chromosome instability occur together in pre-invasive carcinomas. Cancer Res 2003; 63: 1398–1404.
26 Sato N, Mizumoto K, Nakamura M, Nakamura K, Kusumoto M, Niyama H et al. Centrosome abnormalities in pancreatic ductal carcinoma. Clin Cancer Res 1999; 5: 963–970.
3288

55. Navarro C, Nola S, Audebert S, Santoni MJ, Arsanto JP, Ginestier C et al. Functional recruitment of mammalian Scribble relies on E-cadherin engagement. Oncogene 2005; 24: 4330–4339.

56. Feigin ME, Akshintala SD, Arabi K, Rosenberg AZ, Muthuswamy LB, Martin B et al. Mislocalization of the cell polarity protein Scribble promotes mammary tumorigenesis and is associated with basal breast cancer. Cancer Res 2014; 74: 3180–3194.

57. Royer C, Lu X. Epithelial cell polarity: a major gatekeeper against cancer? Cell Death Differ 2011; 18: 1470–1477.

58. Zhan L, Rosenberg A, Bergami KC, Yu M, Xuan Z, Jaffe AB et al. Deregulation of cell polarity promotes mammary tumorigenesis and reveals a role for cell polarity in cancer. Cell 2008; 135: 865–878.

59. Dow LE, Elsum IA, King CL, Kinross KM, Richardson HE, Humbert PO. Loss of human Scribble cooperates with H-Ras to promote cell invasion through deregulation of MAPK signalling. Oncogene 2008; 27: 5988–6001.

60. Bettencourt-Dias M, Rodrigues-Martins A, Carpenter L, Riparbella M, Lehmann L, Gatt MK et al. SAK/PLK4 is required for centriole duplication and flagella development. Curr Biol 2005; 15: 2199–2207.

61. Habadanck R, Stierhof YD, Willkinson CJ, Nigg EA. The Polo kinase Plk4 functions in centriole duplication. Nat Cell Biol 2005; 7: 1140–1146.

62. Debnath J, Muthuswamy SK, Brugge JS. Morphogenesis and oncogenesis of MCF-10A mammary epithelial acini grown in three-dimensional basement membrane cultures. Methods 2003; 30: 256–268.

63. Xiang B, Muthuswamy SK. Using three-dimensional acinar structures for molecular and cell biological assays. Methods Enzymol 2006; 406: 692–701.

64. Bartolini F, Gundersen GG. Generation of noncentrosomal microtubule arrays. J Cell Biol 2006; 171: 4155–4163.

65. Lechler T, Fuchs E. Desmoplakin: an unexpected regulator of microtubule organization in the epidermis. J Cell Biol 2007; 176: 147–154.

66. Mogenen MM. Microtubule organizing centers in polarized epithelial cells. In: Nigg EA (ed). Centrosomes in Development and Disease vol. 1. WILEY-VCH Verlag GmbH & Co. KGaA: Weinheim, 2004, pp 299–320.

67. Hernandez P, Timauer JS. Tumor suppressor interactions with microtubules: keeping cell polarity and cell division on track. Dis Model Mech 2010; 3: 304–315.

68. Leung CT, Brugge JS. Outgrowth of single oncogene-expressing cells from suppressive epithelial environments. Nature 2012; 482: 410–413.

69. Partanen JI, Nieminen AI, Makela TP, Klefstrom J. Suppression of oncogenic properties of c-Myc by LBK1-controlled epithelial organization. Proc Natl Acad Sci USA 2007; 104: 14694–14699.

70. Simpson DR, Yu M, Zheng S, Zhao Z, Muthuswamy SK, Tansey WP. Epithelial cell organization suppresses Myc function by attenuating Myc expression. Cancer Res 2011; 71: 3822–3830.

71. Fujita Y. Interface between normal and transformed epithelial cells: a road to a novel type of cancer prevention and treatment. Cancer Sci 2011; 102: 1749–1755.

72. Hogan C. Impact of interactions between normal and transformed epithelial cells and the relevance to cancer. Cell Mol Life Sci 2012; 69: 203–213.

73. Yamauchi H, Fujita Y. Epithelial self-defense against cancer. Cell Res 2012; 22: 1527–1529.

74. Muthuswamy SK, Li D, Lelievre S, Bissell MJ, Brugge JS. ErbB2, but not ErbB1, reorganizes proliferation and induces luminal repopulation in epithelial acini. Nat Cell Biol 2003; 1: 785–792.

75. Carraway CA, Carraway KL. Sequestration and segregation of receptor kinases in epithelial cells: implications for ErbB2 oncogenesis. Sci STKE 2007; 2007: re3.

76. Guo W, Pylayeva Y, Pepe A, Yoshioka T, Muller WJ, Inghirami G et al. Beta 4 integrin amplifies ErbB2 signaling to promote mammary tumorigenesis. Cell 2006; 126: 489–502.

77. Nagasaka K, Pim D, Massimi P, Thomas M, Tomaic V, Subbiah VK et al. The cell polarity regulator hScrb controls ERK activation through a KIM site-dependent interaction. Oncogene 2010; 29: 5311–5321.

78. Weber K, Bartsch U, Stocking C, Fehse B. A multicolor panel of novel lentiviral vectors for functional gene analysis. Mol Ther 2008; 16: 619–626.

79. Zufferey R, Nagy D, Mandel RJ, Naldini L, Trono D. Multiple attenuated lentiviral vector achieves efficient gene delivery in vivo. Nat Biotechnol 1997; 15: 871–875.

80. Fry AM, Meraldi P, Nigg EA. A centrosomal function for the human Nek2 protein kinase, a member of the NIKA family of cell cycle regulators. EMBO J 1998; 17: 470–481.

81. Kleyenstein S, Westendorf J, Le Clech M, Habedanc R, Stierhof YD, Nigg EA. Plk4-induced centrosome biogenesis in human cells. Dev Cell 2007; 13: 190–202.

82. Schmidt TI, Kleyenstein S, Westendorf J, Le Clech M, Lavoie SB, Stierhof YD et al. Control of centriole length by CPAP and CP110. Curr Biol 2009; 19: 1005–1011.

© 2016 Macmillan Publishers Limited

Oncogene (2016) 2711 – 2722
83 Yan X, Hagedanck R, Nigg EA. A complex of two centrosomal proteins, CAP350 and FOP, cooperates with EB1 in microtubule anchoring. Mol Biol Cell 2006; 17: 634–644.

84 Thein KH, Kleylein-Sohn J, Nigg EA, Gruneberg U. Astrin is required for the maintenance of sister chromatid cohesion and centrosome integrity. J Cell Biol 2007; 178: 345–354.

85 Liang CC, Park AY, Guan JL. In vitro scratch assay: a convenient and inexpensive method for analysis of cell migration in vitro. Nat Protoc 2007; 2: 329–333.

Supplementary Information accompanies this paper on the Oncogene website (http://www.nature.com/onc)