The nucleus measures shape changes for cellular proprioception to control dynamic cell behavior

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Abstract: The physical microenvironment regulates cell behavior during tissue development and homeostasis. How single cells decode information about their geometrical shape under mechanical stress and physical space constraints within tissues remains largely unknown. Here we show that the nucleus, the biggest cellular organelle, functions as an elastic deformation gauge that enables cells to measure cell shape deformations. Inner nuclear membrane unfolding upon nucleus stretching provides physical information on cellular shape changes and adaptively activates a calcium-dependent mechano-transduction pathway, controlling actomyosin contractility and migration plasticity. Our data support that the nucleus establishes a functional module for cellular proprioception that enables cells to sense shape variations for adapting cellular behavior to their microenvironment.

One Sentence Summary: The nucleus functions as an active deformation sensor that enables cells to adapt their behavior to the tissue microenvironment.

Short Title: The nucleus is a cellular mechano-controller

Main Text:

Introduction

The 3D shape of an organism is built by active force-generating processes at the cellular level and the spatio-temporal coordination of morphodynamic cell behavior. Contractility of the actomyosin cell cortex represents a major cellular force production mechanism underlying cellular shape change (1), cell polarization (2) and active cell migration dynamics (3). Contractility levels are
regulated by the activity of non-muscle myosin II motor proteins (4) and are precisely controlled
to tune single cell and tissue morphodynamics during development (5, 6) and tissue homeostasis
and disease in the adult organism (7, 8). Still, mechanisms that regulate the set point level of
cortical contractility on the single cell level remain poorly understood.

To adjust cortical contractility levels, cells need to make quantitative measures of their mechano-
chemical 3D tissue microenvironment and translate this information into a defined morphodynamic output response. During embryogenesis, morphogens that act as chemical information carriers have attracted major attention (9), modulating cytoskeletal and cellular dynamics via receptor signaling pathways that tune protein activities (such as phosphorylation states) and/or protein expression levels. In contrast, physical parameters of the 3D tissue niche and mechanical forces gain importance as regulators of cellular morphodynamics and myosin II-
dependent cortical contractility levels (10, 11). In vivo, mechanical cell deformation and cellular
packing density in crowded tissue regions has been shown to influence major morphodynamic
processes such as cortical actomyosin contractility (12, 13), cell division (14-17), cell extrusion
and invasion (18). Ex vivo studies further provided evidence on the single-cell level that physical
cell deformation is sufficient to modulate cortical myosin II localization and motor protein activity
(19, 20) and influence morphodynamic cell behavior (21, 22).

A recent example has been the identification of a fast amöeboid migration mode, termed stab le-
blep migration, that is triggered by an increase in cortical contractility via genetic or physical cell
perturbation (23). This morphodynamic migration switch was shown to be present in both
undifferentiated and lineage committed embryonic progenitor cells and was further identified in
various other cell types (24-29). This suggests that a conserved, yet unknown, mechanosensitive
cellular signaling module regulates myosin II-based cortical contractility and motile cell
transformation depending on cellular shape deformations in constrained tissue microenvironments.

**Cell contractility levels increase upon cell confinement and regulate migration plasticity**

To approach the question of how cells can measure and adaptively respond to physical cell shape
changes within their 3D tissue microenvironments, we established a synthetic approach that
enables to mimic mechanical cell deformations in controlled 3D microconfinement assays (30). Primary progenitor stem cells were isolated from blastula stage zebrafish embryos and cultured in
planar confinement assays of defined height to mimic various cell deformation amplitudes
(Fig. S1A). Lowering confinement height in discrete steps increased cell deformation, which
scaled non-linearly with a pronounced enrichment of myosin II at the cell cortex relative to cortical
actin accumulation (Fig. 1A,B and Fig. S1B-D; Movie 1). Cortical accumulation of myosin II was
accompanied by an increase in cellular bleb size (Fig. S1E, Movie 1), indicative of an active
increase in cortical contractility levels depending on confinement height. Myosin II re-localization
to the cell cortex in confined cells was rapid (t\textsubscript{1/2} < 1 min, Fig. 1C,D) and temporally stable under
confinement. Of note, distinct plateaus of cortical myosin II enrichment were evident, with
myosin II re-localization increasing for larger cell deformations (Fig. 1C). A cell confinement
height below 7 µm caused a pronounced increase in cell lysis during compression, defining a
maximal threshold deformation of ~30% of the initial cell diameter, given a blastula cell size of
d~25 µm (Fig. S3H). Overall, these data support that the physical microenvironment defines a
specific set point level of cortical contractility as a function of cell deformation.
We have previously shown that an increase in myosin II-mediated cortical contractility induced a stochastic motility switch into a highly motile amoeboid migration phenotype termed stable-bleb mode (23). In accordance with these results, rapid cortical myosin II enrichment in confinement resulted in spontaneous cell polarization which initiated amoeboid cell migration (Fig. 1E,F and Fig. S1F,G, Movie 2,3). Polarized cells revealed characteristic actomyosin density gradients from the cell front towards the rear accompanied with fast retrograde cortical flows (Fig. S1H, Movie 2,3). These cortical flows have been shown to power fast amoeboid migration in polarized cells (23, 31), while unpolarized cells showed random tumbling with minimal net translocation (Fig. S1F,G, S2A). These data support that physical cell deformation in confinement is sufficient to increase actomyosin network contractility and trigger rapid amoeboid cell migration.

Release of cell compression induced a rapid re-localization of cortical myosin to the cytoplasm (Fig. 1G and S1I), followed by a rapid loss of cell polarization and related migratory capacity (Fig. S1G, Movie 4). Interfering with myosin II activity via Blebbistatin inhibited cell polarization and associated cell motility in confinement (Fig. 1E and S1J), in accordance with a necessary role of myosin II-based contractility in cell polarization and migration induced by mechanical cell shape deformation. Cortical myosin II enrichment and cell polarization occurred independently of caspase activation (Fig. S1K), supporting that morphodynamic changes are not caused by the activation of pro-apoptotic signaling programs. Furthermore, transcriptional inhibition did not block cortical myosin II re-localization and cell polarization (Fig. S1L), indicating that a non-genetic program is regulating cellular morphodynamics under cell compression.

During gastrulation, blastoderm embryonic progenitor stem cells specify into different lineages (ectoderm, mesoderm, endoderm) and acquire distinct biomechanical and morphodynamic characteristics, driving germ layer positioning and shape formation of the embryo (32, 33) (Fig. 2A,B). To test the mechanosensitive response to cell deformation at later developmental stages, we obtained different progenitor cell types from embryos via genetic induction or using endogenous reporter lines. Under confinement, non-motile ectodermal cells rapidly polarized and started to migrate in an amoeboid stable-bleb mode. Similarly, mesendodermal cells underwent a fast mesenchymal-to-amoeboid transition in confinement (Fig. 2B-D and Fig. S2A, Movie 3,5). The fraction of polarized stable-bleb cells and their cell speed was comparable for different progenitor cell types in confinement (Fig. S2B,C). Together, these results support that physical cell shape deformation in confined tissue microenvironments activates a mechanosensitive signaling pathway regulating adaptive cortical contractility levels and morphodynamic migration plasticity in pluripotent and lineage committed embryonic stem cells.

The cell nucleus is a mechanosensor of large cell shape deformation

We next sought to identify potential mechanisms that control cellular shape deformation sensing and adaptive morphodynamic behavior. Cortical myosin II re-localization and amoeboid cell transformation occurred on passivated confinement surfaces independently of adhesive substrate coating (Fig. S2A,C and S3A) and cell-cell contact formation (Fig. S3B). These observations support that the activation of cortical contractility in confinement occurs independently of adhesion-dependent mechano-transduction pathways (34). The temporal characteristics of myosin II re-localization dynamics in confined cells showing a fast, stable and reversible accumulation of cortical myosin II (Fig. 1C,D), suggested that shape deformation is sensed by a
non-dissipative cellular element that can rapidly measure and convert gradual cellular shape changes into stable contractility response levels.

The actomyosin cytoskeleton itself has been implicated to act as a mechanosensitive network (20), but generally limits deformation sensing due to rapid turnover of the cell cortex (35). To test for the activation of mechanosensitive ion channels we used Gadolinium and GsMTx4, inhibitors of stretch activated channels, with GsMTx4 having been shown to block the tension dependent Piezo1 channel which is activated following confinement of human cancer cells (36). Treatment with both inhibitors showed no significant reduction in cortical myosin II accumulation under cell deformation (Fig. S3C), despite the presence of functional Piezo1 channels in these cells as validated with the Piezo1 specific agonist Yoda (Fig. S3D).

Interestingly, we observed that cortical myosin II enrichment only started to occur below a threshold confinement height (~13 µm) that correlated with the spatial dimension of the nucleus (Fig. 3A and S3H). Analyzing nuclear shape change versus cortical myosin accumulation revealed a bi-phasic behavior, with a first phase in which the nucleus diameter remained nearly constant and no myosin II accumulation was observed, and a second phase in which the relative myosin accumulation linearly increased with the relative change in nucleus diameter (Fig. 3A,B and S3E). In accordance with this observation, we expected a proportional change of nuclear surface ruffling upon deformation of an initially spherical nucleus. Measuring of nuclear surface folding by the expression of the inner nuclear membrane (INM) protein Lap2b-eGFP revealed that membrane ruffling was continuously reduced when nucleus deformation started to occur at a threshold deformation of ~13 µm (Fig. 3C-E and S3F, Movie 6). In addition, analysis of nucleus membrane curvature for confined versus control cells in suspension indicated INM surface unfolding (Fig. 3F,G, Movie 6), with no significant difference in total nuclear volume and surface (Fig. S3G). Nucleus deformation further correlated with cortical myosin II accumulation in the endogenous in vivo context during the blastula to gastrula transition, when a gradient of cellular packing density appears from the animal pole towards the lateral margin (37) (Fig. S2D,E).

To further probe the dependence of cortical myosin II accumulation on nucleus size, we dissociated primary embryonic stem cells from early and late blastula stages as cells reduce their size in consecutive rounds of early cleavage divisions (Fig. S3H). Deforming cells of different sizes under similar confinement heights revealed that myosin II accumulation is correlated with relative changes in nucleus deformation but not cell deformation (Fig. 4A,B). To test a functional role of the nucleus in regulating cortical contractility levels during cellular shape deformation, we analyzed cortical myosin II accumulation in mitotic cells that present a disassembled nuclear envelope. To arrest cells in mitosis and further increase the percentage of mitotic cells, we used Nocodazole, a microtubule disrupting drug. Confinement of mitotic cells (either spontaneous or Nocodazole-induced) did not trigger a cortical myosin II accumulation at 7 µm confinement height compared to interphase blastula cells (Fig. 4C) or cell polarization (Fig. S3I), although they accumulated myosin II (Fig. 4D) and polarized (Fig. S3I) in response to LPA, a potent activator of Rho/Rock signaling that has previously been shown to induce rapid cortical myosin II enrichment and amoeboid migration in zebrafish embryonic progenitor stem cells (23). During entry into mitosis, cells further gradually lost cortical myosin II accumulations which temporally correlated with the start of nuclear envelope breakdown (Fig. 4E). Altogether, these data show that myosin II enrichment is associated with nuclear shape deformation and stable INM membrane unfolding. This suggests that the nucleus functions as a continuous non-dissipative sensor element
of cell deformation involved in the mechano-sensitive regulation of cortical contractility levels and cellular morphodynamics.

To directly test biophysical characteristics of the nucleus, we developed an assay to probe intracellular nucleus mechanics by optical tweezer measurements. For this purpose, latex beads of 1 µm size were injected into 1-cell stage embryos that dispersed across embryonic cells during early cleavage cycles and acted as intracellular force probes to measure rheological properties of the nucleus (Fig. S4A). Trapezoidal loads were measured for cells in suspension and under 10 µm confinement (Fig. S4B-E). The recorded force followed the fast-initial indentation to reach a peak force before it relaxed to a non-zero constant force-plateau. The relaxation time remained unchanged between suspension ($t = 6.08s \pm 1.1s$) and confined cells ($t = 4.00s \pm 0.6s$) (Fig. S4D-H), suggesting a passive but rapid (second scale) relaxation of a viscous component. The force-plateau on long timescale corresponds to an elastic component of the nucleus (Fig. S4I), in line with previous measurements that identified an elastic behavior of the nucleus (38) that can act as a cellular strain-gauge. In addition, we observed that INM unfolding was stable over a measurement period of 60 min (Fig. S4J) under mechanical cell deformation, supporting that INM stretch does not relax over extended time periods.

**Nuclear deformation activates a calcium-dependent mechanotransduction pathway regulating myosin II activity**

We next aimed at identifying nucleus deformation-dependent signaling pathways that link the spatio-temporal correlation of nuclear shape changes with fast myosin II activation and changes in morphodynamic cell behavior. Our previous observations suggested that nucleus deformation and associated mechano-sensitive processes at the INM interface are involved in the regulation of myosin II activity and cortical contractility. Among a set of molecules tested under confinement conditions (Table S1), we identified cytosolic phospholipase A2 (cPLA2) as a key molecular target mediating the activation of cortical myosin II enrichment (Fig. 5A-B) and amoeboid cell transformation under cell compression (Fig. 5C). Inhibition of cPLA2 by pharmacological interference using Pyrrophenone robustly blocked cortical myosin II re-localization under varying confinement heights (Fig. S5A). Furthermore, we observed a significant reduction of cortical myosin II levels in confined cells by morpholino (MO) interference with cPLA2, while overexpression of cPLA2 mRNA rescued the morphant phenotype and led to a comparable myosin II accumulation as in control cells (Fig. 5A,B). Residual myosin II activation in cells obtained from morphant embryos suggests a maternal contribution of cPLA2 protein in the early embryo which cannot be targeted by MO interference. To exclude that other mechanisms such as structural changes in the actin network prevent cortical myosin II re-localization under cPLA2 inhibition, we added LPA as an exogenous myosin II activator to cPLA2 inhibited cells. Under this condition, myosin II was strongly accumulated at the cell cortex (Fig. S5B,C) and induced cell polarization associated with amoeboid motility (Fig. 5C). These data support that myosin II can be activated by extrinsic pathways when cPLA2 signaling is inhibited and remains competent to bind to the cell cortex.

Recent work identified that the activation of pro-inflammatory signaling during leucocyte recruitment to wounding sites is regulated by tension-sensitive binding of cPLA2 to the INM (39). We thus tested a role of cPLA2 in the nucleus by generating a modified cPLA2 construct containing a nuclear export sequence (NES). Using Leptomycin B as a blocker of nuclear export, an
accumulation of cPLA2-NES-GFP within the nucleus was observed, showing a concomitant increase of cortical myosin II levels in confined cells (Fig. 6A,B). These data support, that cPLA2 localization in the nucleus is required for myosin II enrichment at the cortex.

We further validated that cortical myosin II enrichment in cells of different sizes (early versus late blastula cells) and different embryonic cell lineages (mesendoderm/ectoderm cells) depends on the activation of cPLA2 signaling. Pharmacological inhibition of cPLA2 activity blocked cortical myosin re-localization in confined cells (Fig. 6C) and strongly reduced cell polarization and associated migration competence (Fig. S2B), supporting a consistent role of cPLA2 activation under physical cell deformation across early to late developmental stages. These data support that activation of cPLA2 signaling in the nucleus mediates adaptive cytoskeletal and morphodynamic behavior under cell deformation.

Arachidonic acid (AA) is the primary cleavage product generated by cPLA2 activity (40). To directly validate whether nucleus deformation in confinement triggers cPLA2 activity, we measured the release of AA by Raman spectroscopy. The analysis of Raman spectra confirmed the specific production of AA in confined cells (Fig. 6D and Fig. S5E), with the increase in AA production in confined versus control cells being specifically blocked in the presence of cPLA2 inhibitor (Fig. 6E). We further observed that AA was exclusively detected in the cytoplasm of confined cells, arguing that AA is directly released from nuclear membranes into the cytoplasm. These data support that cell confinement leads to enhanced cPLA2 activity and production of arachidonic acid associated with INM unfolding and stretching of the nucleus surface.

AA has been implicated to regulate myosin II activity both directly (41) and indirectly via protein phosphorylation (42). We tested the involvement of Rho/Rho associated protein kinase (ROCK) and Calcium/Myosin light chain kinase (MLCK) signaling that act as key regulatory pathways of myosin II activity (4). MLCK inhibition showed no significant effect on myosin II enrichment in confined cells, while a pronounced reduction of cortical myosin II recruitment was observed when using Staurosporin, a general kinase inhibitor, and under specific inhibition of Rho activity (Fig. 6F). Using a RhoA-Fret sensor further indicated an increased RhoA activity in confined cells versus control cells in suspension which was significantly reduced in the presence of cPLA2 inhibitor in confined cells (Fig. S5F). These data support that AA production by cPLA2 activity engages upon nuclear envelope unfolding, regulating phosphorylation-dependent myosin II activity at the cell cortex. AA and its metabolic products have been widely implicated in para- and autocrine signaling functions involving GPCRs (43). We assessed the role of AA product release by compressing cells directly under micropillars at an approximate height of 7 µm, leading to confined and non-confined cells in close proximity (Fig. S5G). Interestingly, while confined cells showed high levels of cortical myosin II and amoeboid cell transformation, non-confined cells in direct contact revealed no alterations in cortical myosin II and cellular morphodynamics (Fig. S5H). These results support that cPLA2-dependent myosin II activation does not depend on the release of diffusible signals and regulates cellular morphodynamics via a cell autonomous increase of cortical contractility under cell deformation in confinement.

Interference with intracellular calcium levels by addition of BAPTA-AM or in combination with cPLA2 inhibitor blocked myosin II enrichment in confined cells, without altering cortical myosin II levels in unconfined control cells (Fig. S5I). LPA stimulation of BAPTA-AM treated cells confirmed that myosin II can be activated by the Rho-ROCK signaling pathway in the absence of intracellular calcium and remains competent to bind the cell cortex (Fig. S5C). Similarly, chelating extracellular calcium reduced cortical myosin II re-localization, while depletion of internal
calcium stores using Thapsigargin led to a slight increase in myosin II enrichment in confinement (Fig. S5I). The addition of ionomycin showed that high intracellular calcium levels, in the absence of cellular shape deformation, were not sufficient to evoke AA production (Fig. S5J) and cortical myosin II enrichment (Fig. S5B,D). This suggests that intracellular calcium has a permissive function for cPLA$_2$ association with the INM and increasing cortical contractility under cell confinement. Our findings are in line with the observation that cPLA$_2$ contains a calcium-dependent C2 domain that modulates protein binding to the INM (44-46), which has been shown to get further enhanced and stabilized by mechanical stretching of the nucleus (39). Consistently, cPLA$_2$ showed a transient localization to the INM in the presence of ionomycin, while additional mechanical stretch due to hypotonic swelling or cell confinement induced a stable association with the INM (Fig. S3J-K).

**INM unfolding and nucleus positioning enable to decode different types of cell shape deformations**

To study if INM unfolding under cell confinement was sufficient to trigger cPLA$_2$ activity, we measured cortical myosin II levels and amoeboid migration competence under hypotonic swelling of cells. Quantification of nuclear shape parameters (size, volume, surface) revealed that hypotonic swelling induced comparable nuclear surface expansion and INM unfolding as nucleus deformation under a confinement height of 7 µm (Fig. S6A-C). Interestingly, cortical myosin II levels in hypotonic conditions (Fig. 7A, Movie 7) and associated changes in bleb size (Fig. S6D) and cell polarization rate (Fig. 7C) were significantly lower compared to cells deformed at 7 µm confinement height. These observations suggest that nuclear envelope unfolding alone is not sufficient to trigger high levels of cortical myosin II enrichment under isotropic cell stretching in hypotonic conditions versus anisotropic cell deformation in confinement.

Comparing intracellular calcium levels between deformed cells in confinement and under hypotonic conditions showed a pronounced increase in intracellular calcium concentrations in confined cells, with a specific calcium increase in the cell nucleus (Fig. 7B and Fig. S6E). Of note, ectopically increasing intracellular calcium levels under hypotonic conditions via the addition of ionomycin led to a pronounced and rapid increase in cortical myosin II enrichment in a cPLA$_2$-dependent manner (Fig. S6F, Movie 7) that triggered spontaneous cell polarization (Fig. 7C,D and S6G). Similar to cells in confinement, cell polarization under this condition was associated with a rapid transformation of non-motile cells into a highly motile stable-bleb amoeboid mode with fast migration speed under confinement ex vivo and in vivo (Fig. 7E and S6H,I,M, Movie 7,8). Raman spectroscopy to directly measure cPLA$_2$ activity confirmed that hypotonic stress increased AA levels (Fig. 5K) in a cPLA$_2$-dependent manner (S5L), with addition of ionomycin in hypotonic conditions further increasing AA production (S5M). Relative measured AA levels further directly correlated with cortical myosin II levels (Fig. S5N). Together, these data reveal that different mechanical shape deformations regulate intracellular calcium levels and modulate cPLA$_2$ activity under similar INM stretch: uniaxial compression in confinement induces high intracellular calcium levels specifically in the nucleus, while isotropic radial stretch in hypotonic stress conditions leads to a lower intracellular calcium levels. Independently modulating nucleus deformation and calcium levels under different shape deformations confirmed that both parameters engage synergistically to regulate cortical contractility (Fig. 8A) and cellular dynamics under mechanical stress (Fig. 7E), thereby enabling a cell to distinguish between different types of shape deformation and to acquire a specific morphodynamic response.
Intracellular nucleus positioning appeared a promising candidate to differentially modulate calcium levels. ER-plasma membrane (ER-PM) proximity has been implicated as an important regulator of cellular calcium signaling (47). Visualization of membrane proximal ER structures showed that the ER was highly dynamic under conditions of low confinement but was increasingly immobilized between the nucleus-PM interface for larger cell deformations in confinement (Movie 9). In addition, the expanding nucleus contact area close to the plasma membrane closely correlated with an intracellular calcium increase (Fig. S6J). We speculated that STIM-Orai, with STIM proteins located at the endoplasmic reticulum (ER) and Orai proteins representing calcium-selective plasma membrane (PM) calcium channels, could be involved in cellular calcium regulation in confined cells. STIM-Orai proteins have an established function in store operated calcium entry (SOCE) upon depletion of calcium from the ER which are mediated via ER-PM proximity (48, 49). Analysis of STIM-Orai protein localization revealed that both proteins accumulate at the interface between the nucleus and plasma membrane in confined cells at 7 µm (Fig. S6K,L). In contrast, we observed a homogenous distribution of both channels in cells under 13 µm confinement height when the ER was not spatially confined between the nucleus and PM. These data support that ER immobilization is associated with the specific enrichment of STIM-Orai in the ER-PM contact region where mechanical ER confinement occurs. Inhibition of the STIM-Orai complex by using 2APB further blocked myosin II enrichment under cell deformation in confinement (Fig. S5I). Our observations support that mechanical compression of the cell nucleus induces a tight connectivity between ER-PM structures and STIM-Orai localization at the ER-PM contact interface involved in the upregulation of intracellular calcium levels in confined cells.

Conclusions

Our data support that the nucleus establishes a core element to measure cellular shape deformation via two key physical parameters: 1) nuclear shape deformation leading to INM unfolding and 2) intracellular spatial positioning of the nucleus. In this model, INM unfolding under nuclear shape change allows for the deformation-dependent activation of cPLA$_2$ signaling, whereby cPLA$_2$ activity is modulated by intracellular calcium levels set by nucleus-PM proximity (Fig. 8B and S7A). The parameter space of these two variables (INM unfolding and calcium levels) provides a unique identifier for a cell to decode distinct shape deformations as exemplified on anisotropic cell deformation in confinement versus isotropic hypotonic cell stretching, allowing cells to acquire a unique adaptive response depending on the type of physical shape deformation (Fig. S7B).

Biochemical, physical and mechanical cues in the surrounding of a cell create manifold information for cells which is continuously sensed, integrated and transduced to allow for complex cellular functioning. Here we show that the cell nucleus functions as a cellular mechano-gauge for precisely decoding cellular shape changes, allowing cells to adaptively and rapidly tune cytoskeletal network properties and morphodynamic behavior within their 3D tissue microenvironment during development. This mechanism lays a foundation for functional principles underlying cellular proprioception that, comparable to the sensing of spatio-temporal changes in body posture and movement (50), enable a precise interpretation of shape changes on the single cell level.

The nucleus, being the largest organelle within the cell, represents a prominent structure to transmit and modulate mechanosensitive processes (51-55), and nucleus deformation has been shown to
influence nuclear transport and cell differentiation (56-58), chromatin organization (59-61), migration (62-66) and pathfinding in constrained environments (67). Our findings support that nucleus deformation and its intracellular positioning establish a cellular sensing module that equips cells to rapidly and reversibly adapt their dynamic response to shape fluctuations. This “nuclear ruler pathway” was also identified in an accompanying study (68), supporting its conservation between embryonic and differentiated cells in the adult organism. The observation of a rapid contractile cell response upon cell squeezing in confinement which is followed by fast amoeboid motility is reminiscent of a “cellular escape reflex” that enables cells to cope with physical constraints and acute deformations of their shape. Such a mechanism might be relevant for various biological processes associated with migration plasticity of cancer and immune cells in constrained 3D tissue niches (69-72) mecano-chemical feedback processes during morphogenesis (73) and homeostatic tissue functions such as cell density regulation (74), which require accurate mechanisms to detect variations in cellular size and shape and multicellular packing density in crowded 3D tissues.

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RNA preparations; F.C-C. and V.V. performed optical tweezers experiments and F.C. analyzed the data; M.M. and V.V. performed Raman experiments and M.M. analyzed the data; M.C-R. analyzed the Lap2B-GFP data; H-M.M. and Q.T.-R. performed in vivo experiments and H.-M.H. performed injections and helped with mesendoderm-ectoderm experiments; S.J-D. cloned plasmids, synthetized mRNA and performed mRNA and bead injections; S.P-L. performed calcium imaging related to the role of Piezo channels; M.A.V. supervised S.P-L. and contributed with discussions and support to calcium imaging experiments; J.W. provided biological reagents and supported molecular cloning. M.K. supervised F.C.-C. and designed tweezer experiments. S.W. and V.R. supervised the project, contributed to data analysis and wrote the manuscript.

**Competing interests:** The authors declare no competing financial interests. **Data and materials availability:** All data relevant for the conclusions of this work are available in the main text or the supplementary materials.

**Supplementary Materials:**

Supplementary Methods
Figures S1-S7
Supplementary Table S1
Supplementary Movies S1-S9
Supplementary References (80-86)
Figure 1. Cell deformation in confined environments defines cell contractility, polarization and fast amoeboid cell migration. (A) Relative cortical myosin II enrichment for decreasing confinement height in un-polarized progenitor cells (N=477 (suspension, unconfined); N=56 (18 µm); N=35 (16 µm); N=103 (13 µm); N=131 (10 µm); N=49 (8.5 µm); N=348 (7 µm)). Significance values are with respect to the suspension condition. Black line shows a mono-exponential fit with offset to the data. (B) Exemplary confocal fluorescence images of control progenitor stem cells in suspension (Susp.) and indicated confinement heights expressing Myl12.1-eGFP (myosin II). White arrows point at cellular blebs. (C) Temporal dynamics of cortical myosin II recruitment upon mechanical confinement at time t=0 at indicated heights. Thick lines correspond to the mean and areas to the standard error of the mean (sem). N>50 cells for all conditions. (D) Exemplary cross-sectional time-lapse images of myosin II-eGFP expressing cells under 7 µm confinement. (E) Percentage of polarized motile stable-bleb cells in suspension at
indicated confinement heights and myosin II inhibition (blebbistatin) at 7 µm (each N>500). (F) Representative time-lapse images of a myosin II-eGFP expressing cell undergoing spontaneous stable-bleb cell polarization and migration initiation. Dashed line shows the stable-bleb cell front and red arrow points in the direction of movement. (G) Relative cortical myosin II enrichment during reversible cell confinement. Cells were confined for 15 min before confinement was released and cortical myosin II levels were measured at $t_0$ (0-5 min) and at $t_0+\Delta t$ (30-60 min) after release. ***p<0.0001, **p<0.001, *p<0.01, not significant (n.s.). All scale bars 10 µm.
Figure 2. Physical confinement triggers amoeboid migration in different cell lineages. (A) Sketch of the developing zebrafish embryo at sphere (4 hpf), shield (6 hpf) and 75% epiboly (8 hpf) stage. (B) Exemplary confocal and bright field images of mesodermal cells in vivo expressing Lyn-Tomato (magenta, membrane) and GFP (green) under the mezzo promoter (left), induced mesendodermal cells in vitro plated on a 2D fibronectin-coated surface (middle) and under 7 μm confinement (right), asterisk points at stable-bleb cell front. (C) Relative cortical myosin II intensity for mesendodermal and ectodermal progenitor cells in control suspension and confinement conditions. (D) Exemplary confocal images of stable-bleb polarized (top) and non-polarized (bottom) progenitor cell types expressing myosin II-eGFP under 7 μm confinement. From left to right: ectoderm, endoderm and mesoderm cells. Dashed line and yellow asterisks indicate the stable-bleb cell front and red arrows the direction of cell migration. ***p<0.0001, **p<0.001, *p<0.01, not significant (n.s.). All scale bars 10 μm.
**Figure 3.** Nuclear envelope unfolding is associated with increasing cortical contractility. (A) Double boxplot of relative cortical myosin II enrichment (left axis, grey) and relative nuclear diameter (right axis, green) for decreasing confinement height. Statistical test for relative nuclear diameter performed with respect to suspension (N=144 (suspension); N=44 (18 µm); N=32 (16 µm); N=37 (13 µm); N=45 (10 µm); N=37 (8.5 µm); N=50 (7 µm)). Statistical tests for cortical myosin II levels and experimental N related to Fig.1A. (B) Exemplary confocal top views (x-y) and lateral views (y-z) of progenitor stem cells expressing myosin II-eGFP stained with DNA-Hoechst and ER-TrackerRed in 13 µm and 7 µm confinement. (C) Illustration showing the unfolding of the inner nuclear membrane (INM) with increasing cell confinement. (D) Nuclear area to perimeter ratio (top) and nuclear invagination ratio (bottom) for increasing confinement. (E) Exemplary confocal images of cells expressing Lap2B-eGFP under varying cell deformation. (F) Curvature analysis of nuclear shape for 20 consecutive frames (t_{lag}=10 s) for unconfined (suspension, top) and 7 µm confined nuclei (bottom). (G) Histogram of nuclear curvature for
unconfined and 7 µm confined nuclei related to panel F with N=10 cells for each condition (p<10^{-12}, ***). ***p<0.0001, **p<0.001, * p<0.01, not significant (n.s.). All scale bars 10 µm despite 5 µm in (B).
Figure 4. Nucleus size and integrity determine the morphodynamic cell response to confinement. (A) Relative cortical myosin II intensity with respect to cell deformation as defined by the ratio of cell size/confinment height (left) and nucleus deformation (right) measured via relative nucleus diameter increase (right) for cells dissociated from embryos at high-oblong (3.3 hpf) and sphere stage (4 hpf) and cultured under similar confinement heights as indicated. Data points and error bars indicate mean and standard error of the mean (sem); (3.3 hpf: 16 µm N=58, 13 µm N=61; 4 hpf: 16 µm N=44, 13 µm N=60, 10 µm N=45). The Pearson correlation coefficient in between myosin II intensity and nuclear deformation is 0.48. (B) Exemplary confocal images of progenitor cells expressing myosin II-eGFP under 16/13 µm confinement dissociated from 4 hpf (top) and 3.3 hpf (bottom) embryos. Dashed lines outline cell nuclei. (C) Relative cortical myosin II enrichment for interphase and mitotic cells under 7 µm confinement cultured in suspension (control) in the presence of 1 µM Nocodazole. Exemplary confocal images of progenitor cells expressing myosin-II-eGFP in interphase (yellow asterisk) or during mitosis (cyan
asterisk) under 7 µm confinement. (D) Exemplary confocal images of progenitor unpolarized (top) and polarized (bottom) cells expressing myosin-II-eGFP during mitosis treated with 1 µM Nocodazole and 50 µM LPA. Dashed line shows the stable-bleb cell front. (E) Cortical myosin II intensity of N=7 cells undergoing mitosis and representative confocal time lapse images of cells expressing myosin-II-eGFP and H2A-mCherry at indicated time points (t=0: time of nuclear envelope breakdown). The arrows point at the cell cortex with cortical myosin enrichment (t=-1min) or loss of cortical myosin levels (t=7.5min). Data points and error bars indicate mean and standard error of the mean. ***p<0.0001, **p<0.001, * p<0.01, not significant (n.s.). All scale bars 10 µm.
Figure 5. Nucleus deformation activates a mechanosensitive lipase signaling pathway regulating myosin II activity. (A) Relative cortical myosin II intensity for progenitor cells cultured in suspension versus 7 µm confinement conditions for control cells (DMEM), with cPLA2 inhibitor, or injected with cPLA2 MO and cPLA2 MO + cPLA2 mRNA. (B) Exemplary confocal images of progenitor cells expressing myosin II-eGFP related to (A) for indicated conditions. (C) Percentage of stable-bleb polarized cells for control cells under 7 µm confinement and in the presence of cPLA2 inhibitor or unconfined (suspension) cells stimulated with 50 µM LPA. For all conditions N>200. ***p<0.0001, **p<0.001, *p<0.01, not significant (n.s.). All scale bars 10 µm.
Figure 6. cPLA2 activity at the inner nuclear membrane generates AA as a metabolite regulating cortical contractility in confinement. (A) Relative cortical myosin II fluorescence intensity for cells dissociated from controls (un-injected) embryos or embryos injected with cPLA2 MO, cPLA2 MO + cPLA2-NES-GFP mRNA (RNA) and with or without addition of Leptomycin B (Lb). (B) Exemplary confocal fluorescence images of cell expressing myosin II-mCherry (right) and cPLA2-NES-GFP (left) under 7 µm confinement with (top) or without (bottom) the addition of Lb. (C) Relative cortical myosin II fluorescence intensity upon cPLA2 inhibition for cells dissociated at 3.3 hpf, induced mesendoderm or ectoderm cells in suspension and upon confinement at indicated height. (D, E) Scores of Raman component associated to AA in suspension (unconfined, N=24) and confined cells (10 µm, N=28) and (H) in control confinement condition (Ctrl, N=52) or treated with cPLA2 inhibitor (N=22). Red lines indicate mean and sem. (F) Relative cortical myosin II intensity for control cells and different chemical (Y-27637, M-L7,
Staurosporine) or genetic interference (dnRhoA) with myosin II regulators. ***p<0.0001, **p<0.001, *p<0.01, not significant (n.s.). All scale bars 10 µm.
Figure 7. Nucleus stretch and intracellular positioning enable an adaptive cellular response to different types of physical cell deformation. (A) Relative cortical myosin II enrichment for progenitor cells cultured under different osmolarity conditions. (B) Normalized cell (left) and nuclear (right) Ca\(^{2+}\) levels (Calbryte520) for control (Ctrl) and hypotonic (0.5 x) conditions and mechanical confinement (13 µm, 7 µm). (C) Percentage of stable-bleb polarized cells in isotonic and hypotonic (0.5 x) conditions for cells cultured in DMEM (Ctrl) or supplemented with 1 µM ionomycin. N>1000 cells for all conditions. (D) Exemplary confocal images of cells expressing myosin II-eGFP in isotonic (ctrl, top-left), hypotonic (bottom-left) and hypotonic conditions supplemented with ionomycin treatment (right): non-polarized cell (top) and stable-bleb polarized cell (bottom). (E) Sketch of cell polarization and motile cell behavior in 2D (top) versus 3D confined environments (bottom) and for control conditions (isotonic media; first column) versus cPLA\(_2\) interference (second column) and hypotonic condition alone (third column) or in the presence of ionomycin (fourth column). ***p<0.0001, **p<0.001, *p<0.01, not significant (n.s.). Scale bars 10 µm.
Figure 8. INM unfolding and intracellular calcium levels enable cells to decode isotropic stretch versus cell squeezing in confinement. (A) Normalized relative cortical myosin II fluorescence intensity as a function of nucleus unfolding and normalized Ca$^{2+}$ (Calbryte) intensity for different physical cell deformations (dark blue: 7 µm; light blue: 13 µm confinement; yellow: 7 µm confinement + Bapta-AM; red: hypotonic (0.5x) condition; magenta: hypotonic condition (0.5x) + ionomycin). Data indicate mean and sem. The gray area sketches the relation between cortical myosin II and nuclear deformation versus intracellular calcium levels. (N: calcium data related to Fig. 4B, except for hypo + ionomycin (N=41) and Bapta-AM + 7 µm N=9; for nuclear unfolding related to Fig. 2D, Fig S6B and for cortical myosin II levels to Fig. 1A, S5I, 4A, S6F). (B) Sketch depicting how nucleus deformation and intracellular nucleus positioning correlate with INM unfolding and intracellular calcium levels, which differentially regulate cortical contractility and cellular morphodynamics. ***p<0.0001, **p<0.001, *p<0.01, not significant (n.s.). Scale bars 10 µm.
Materials and Methods

Zebrafish Maintenance Zebrafish (Danio rerio) were maintained as previously described (75). Embryos were kept in E3 medium at 25°C–31°C prior to experiments and staged based on morphological criteria (76) and hours post fertilization (hpf). Wild type embryos were obtained from the AB strain background. All protocols used have been approved by the Institutional Animal Care and Use Ethic Committee (PRBB–IACUEC) and implemented according to national and European regulations. All experiments were carried out in accordance with the principles of the 3Rs.

Transgenic fish lines The following transgenic lines were used: Tg(actb2:Lifeact-GFP) (77), Tg(actb2:Myl12.1-eGFP) (78), Tg(actb2:Myl12.1-mcherry) (4), Tg(zebbio:eGFP) (23), Tg(actb2:Lyn-TdTomato) (79). All progenitor cells expressing Myl12.1-eGFP (Myosin II) and Lifeact-GFP (Actin) were obtained from Tg(actb2:Myl12.1-eGFP) and Tg(actb2:Lifeact-GFP).

Cell culture To culture progenitor cells, embryos were manually dechorionated in E3 buffer at sphere stage (4 hpf) or different stages if indicated. Five to twenty embryos were transferred to DMEM-F12 (with L- Glut and 15mM HEPES, without sodium bicarbonate and phenol red) culture medium (Sigma) and mechanically dissociated by manual tapping followed by centrifugation at 200 g for 3 min.

Sample preparation and surface coatings The following products for surface coatings at the indicated concentration have been used: 0.5 mg/ml PLL(20)-g[3,5]-PEG(2) (Susos) and 0.2 mg/ml fibronectin (Sigma-Aldrich). Prior to PLL-PEG coating, both coverslips and dishes were plasma cleaned. Uncoated or PLL-coated glass dishes #1.5 were purchased from MatTek (MatTek Corporation).

Cell confiner Cells were confined using a dynamic confiner (4DCell) similar to previously established planar microconfinement methods (30). In order to confine cells at different heights, multiple Si molds were produced by photolithography in a clean room (Nanofabrication laboratory, ICFO) by depositing a SU-8 resin on a silicon wafer. In brief, a photomask with the desired geometry was created. Confinement coverslips were prepared with polydimethylsiloxane (PDMS) with the following heights: 18, 16, 13, 10, 8.5 and 7 µm. Coverslips were always plasma cleaned, coated with PLL-PEG if not indicated otherwise and equilibrated in DMEM prior to each experiment. A pressure pump (AF1 – Microfluidic pressure pump, Elveflow) together with the ESI software was used to change the pressure for tuning the confinement heights. For Raman measurements and optical tweezers two coverslips separated with microbeads or with a PDMS membrane have been used (height h=10 µm).

Reagents and inhibitor treatments Pharmacological inhibitors were used at the following concentrations: 1 µM cPLA2 inhibitor (Pyrropheneone, Merck-Millipore), 10 µM Bapta-AM (Cayman), 10 µM Blebbistatin (+) (Tocris Bioscience), 10 µM Y-27632 (Tocris Bioscience), 10 µM ML-7 hydrochloride (Tocris Bioscience), 1 µM Nocodazole (Sigma), 50 nM LeptomycinB (Sigma-Aldrich), 1 µM Ionomycin (Sigma-Aldrich), 1 µM Thapsigargin (Thermofisher), 10 µM GsMTx4 (Tocris), 50 µM 2-APB (Biogen-Santa Cruz), 10 µM Gadolinium chloride (Tocris), 2 µM Actinomycin D (PanReac), 100 µM Z-VAD(OMe)-FMK (Abcam), 1 µM Staurosporine (Abcam), 1-oleoyl lysophosphatidic acid (LPA, Tocris Bioscience) was used at the indicated concentrations. Measurements were done directly after exposure to MLCK inhibitor, GsMTx4, Ionomycin, LPA; all the other inhibitors have been pre-incubated for 30 min and 60 min for Y-27632 prior to experiments.
Fluorescence staining Calbryte520 (AAT BIOQUEST) was used to study calcium dynamics. The staining kit-Red Fluorescence-Cytopainter (ER Tracker, Abcam) or ER-Tracker™ Green (BODIPY™ FL Glibenclamide) were used to visualize the endoplasmic reticulum respectively for confocal 3D colors imaging and for TIRF microscopy experiment. DNA-Hoechst (Thermofisher) was used to stain the cell nucleus. Cells were incubated with 5 µM Calbryte520 for 20 min, with 1 µM ER-tracker for 30 min and with 1 µg/ml DNA-Hoechst for 7-10 min as reported in the corresponding protocols. After incubation, cells were washed, centrifuged at 200 g for 3 min and re-suspended in DMEM media.

Variable osmotic culture conditions D-Mannitol (Sigma) was diluted in DMEM in order to obtain a culture medium with an osmolarity of approx. 450 mOsm/l (corresponding to a 1.5x media). Milli-Q water was added to DMEM for hypotonic conditions.
References

49. X. Qin et al., Increased Confinement and Polydispersity of STIM1 and Orai1 after Ca(2+)+ Store Depletion. Biophys J 118, 70-84 (2020).


Supplementary Materials for

The nucleus measures shape changes for cellular proprioception to control dynamic cell behavior

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This PDF file includes:

Supplementary Methods
Figures. S1 to S7
Supplementary Table S1.
Captions for Movies S1 to S9
Supplementary References

Other Supplementary Materials for this manuscript include the following:

Movies S1 to S9
Supplementary Methods

**Plasmid cloning** The following constructs were subcloned in a pCS2+ or pCS2+-eGFP vector, linearized with BamHI-EcoRI restriction enzymes, using the Gibson cloning system: LAP2b-eGFP was amplified from a pME 18S-FL3 vector (clone #2643665, Dharmacon); cPLA2a (pla2g4aa) plasmid was amplified from a pCR4-TOPO vector (clone #9037889, Dharmacon). Human STIM and Orai were subcloned from mCFP-N1 and mYFP-N1 plasmids (80), respectively, into a pCS2+ vector and linearized with EcoRI-XbaI restriction enzymes. All cDNAs were amplified using Phusion HF DNA Polymerase (Thermofisher F530S), see primers below. NES-cPLA2a-eGFP was amplified from the pCS2+-cPLA2a-eGFP vector using primers encoding for an N-terminal NES sequence (LPPLERLTL). The following constructs were provided from different labs: pCS2-DNRhoA N19 (81); pCS2+_cyclops (82); pCS2+ lefty and Casanova (courtesy Carl-Philipp Heisenberg); pCS2+ Lyn-TdTomato (courtesy Berta Alsina); pTriEx-RhoA FLARE.sc Biosensor WT was a gift from Klaus Hahn (Addgene plasmid #12150; RRID:Addgene_12150).

Oligonucleotides used for cloning:

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**Zebrafish mRNA injections** mRNA was synthesized using the mMessage mMachine Kit SP6 Kit (Ambion AM1340M). All mRNA injections were done in 1-cell stage embryos. To visualize the inner nuclear membrane (INM) 80 pg of Lap2B-eGFP were injected into wild type AB or Tg(actb2:Lyn-TdTomato) embryos. To interfere with myosin II regulators 100 pg of dn-RhoA mRNA (81) have been injected in Tg(actb2:MyI12.1-eGFP) embryos together with 100 pg LynTomato mRNA (to visualize plasma membrane). For RhoA-Fret imaging 400 pg of RhoA-Biosensor (83) were injected in wild type embryos. To induce mesendoderm, endoderm, mesoderm or ectoderm cells, 1-cell stage wild type or Tg(actb2:MyI12.1-eGFP) embryos were injected respectively with: 100 pg cyclops mRNA, 50 pg Casanova mRNA, 100 pg cyclops mRNA and 2 ng Casanova morpholino (GCATCCGGTCGAGATACATGCTGTT), 100 pg Lefty mRNA, all supplemented with 100 pg LynTomato mRNA. To visualize cPLA₂ localization 50 pg of cPLA₂-eGFP mRNA were injected into 1-cell stage wild type or Tg(actb2:MyI12.1-mCherry) embryos. To visualize the STIM-Orai complex 80 pg of STIM-CFP together with 50 pg of Orai-YFP mRNA were injected into 1-cell stage wild type embryos.

**Morpholino interference and rescue experiments** For inhibiting cPLA₂ activity, 2.7 ng of cPLA₂ morpholino (AAGCGTCACTTATAATGTTGGA) were injected in 1-cell stage Tg(actb2:MyI12.1-eGFP) or Tg(actb2:MyI12.1-mCherry) embryos. To rescue the activity, cPLA₂ morpholino was co-injected with 200 pg of cPLA₂ mRNA and 100 pg of LynTomato mRNA or 200 pg of NES-cPLA₂-GFP mRNA. The same concentrations of mRNA were used for control experiments.

**Zebrafish blastula injections** For injection of hypotonic media, Zebrafish embryos at sphere stage were dechorionated and placed in single embryo agarose wells (Adaptive Science Tools). Injections were applied into the extracellular space at the animal pole. Each embryo was injected with an average of 16 nl of injection mix containing hypotonic media (D3:MilliQ 1:1) with Dextran and Alexa Fluor™ 546 (10 kMW, Anionic, ThermoFisher Scientific) to label the interstitial fluid and together with 10 μM Ionomycin calcium salt (Sigma-Aldrich).

**Embryo mounting:** Embryos were mounted in 2% low melting point agarose prepared in Danieu’s solution (58 mM NaCl, 0.7 mM KCl, 0.4 mM MgSO4, 0.6 mM Ca(NO3)2 and 5 mM HEPES) on a Mattek dish and covered with Danieu’s solution.

**Fluorescence imaging** Confocal fluorescence images were acquired using a commercial Leica TCS SP5 STED CW or Leica TCS SP8 STED 3X microscope equipped with a white light laser source (Leica Microsystems, Wetzlar, Germany). In both cases we used a 63x oil objective.
For myosin II-eGFP, LifeACT-eGFP, Lap2B-eGFP and Calbryte520 AM, ePLA2-NES-GFP imaging, samples were excited with a 488 nm Argon laser using the SP5 microscope. For co-staining with the Lyn-Tomato membrane reporter or for myosin II-mCherry experiments, a HeNe laser at 543 nm has been used for excitation and consecutives images have been acquired. In both color channels fluorescence was collected using single molecule detectors (SMD-HyD) in photon counting mode and transmission light was collected using a forward PMT. The Leica SP8 confocal microscope has been used for three color imaging. Myosin II-GFP and ER-tracker Red have been excited respectively at 488 nm and 587 nm using a tuneable white light laser. The fluorescence light was collected using two backwards HyD detectors in photon counting mode while the DNA-Hoechst has been excited with a 405 nm semiconductor laser and fluorescence recorded using a PMT. The same microscope was used for FRET imaging: the CFP was excited with the 405nm laser and the YFP at 512 nm using the white light laser and the emitted photons have been collected using the HyD detector in photon counting modes. Transmission light was collected using a forward PMT. For time-lapse movies related to Fig. S2A and Supplementary Movie S3, a 40X oil objective (NA=1.25, HCX PL APO) was used and myosin II-eGFP samples were excited with a 488 nm Argon laser and fluorescent intensity recorded with a SMD-HyD detector in standard mode using the SP5 microscope. A temperature controller set at T=28.5°C was used for all experiments.

In vivo confocal imaging Embryos were imaged using a Leica SP5 confocal microscope equipped with a Leica 20X NA 0.7 immersion objective using a 488 nm laser and the emitted fluorescence was detected with a HyD detector. The temperature during imaging was kept constant at 28.5°C using a temperature chamber.

Bright field imaging Bright field movies were acquired using a Leica DMI-LED microscope equipped with IDS-CMOS cameras (UI-3880LE-M-GL) and a Leica 0.4x C-mount. Air objectives 10x (NA=0.25) or 20x (NA=0.40) were used to image cell dynamics. Acquisition was controlled using μManager (84).

TIRF imaging (related to Supplementary Movie 9) Images were acquired using a custom-built TIRF microscope equipped with an Andor Zyla 4.2 cMOS camera and a UPLAP0100XOHR objective. ER-tracker Green was excited with a 473 nm GEM laser by Laser Quantum using an exposure time of 0.5 s.

Optical trapping experiments The optical tweezer (OT) platform (SensoCell, Impetux Optics, Spain) consists of a continuous wave laser (λ=1064 nm, 5 W nominal output power, Azur Light) steered with a pair of acousto-optic deflectors (AOD) and a force detection unit that captures the forward-scattered light from the optical traps. This is mounted around an inverted research microscope (Nikon Eclipse Ti2) equipped with a spinning disk confocal microscope (Andor DragonFly 502) on top of an active isolation table (Newport). The laser is directed onto a microscope objective (MO, 60x/NA=1.2, water immersion, Nikon) after being expanded with a telescope to fill the MO entrance pupil through the epi-fluorescence port. A short-pass dichroic mirror reflects the IR trapping beam and transmits both the excitation and emission light for fluorescence microscopy, as well as bright-field.

The force detection unit of our optical tweezers platform operates by detecting light-momentum changes, after capturing the scattered trapping beam through an NA = 1.4, oil immersion lens, with a position-sensitive detector (PSD) placed at the back focal plane (BFP) (85). This allowed us to measure forces beyond the linear trapping regime, thus covering the full spectrum until ~300 pN and enabling to work with lower laser powers during intracellular force measurements, as
compared to standard BFP interferometry \((85)\). To perform intracellular trapping, we injected a 0.5 nL drop of diluted (1:5) 1 μm polystyrene beads (Sigma-Aldrich) into 1-cell stage embryos. Cells were seeded in homemade trapping microchambers consisting of a bottom-dish (Wilco Glass, #1.5) and a 1 x 1 inch cover glass spaced with a double scotch, 90 μm high tape. The bottom dish was coated with Concanavalin A (0.05 mg/ml, 30 min, Sigma-Aldrich) to partially adhere the cells and avoid movement due to cellular blebbing. For the optical trapping experiments in confinement conditions, we used two cover glasses (60 x 24 cm, # 1.5, Ted Pella) to enclose a few μl of cells and 10 μm PS beads used as spacers. Only non-polarized cells containing one bead were used for each measurement. The bead was trapped by manually directing the focused laser beam on it, at 300 mW power (at the sample plane; 2.5 W output power) and was then brought against the nuclear membrane. Trap positioning and force measurements were carried out by addressing the AODs in synchronization with the PSD with the OT platform software (LightAce, Impetux Optics, Spain), based on LabView (National Instruments). This software allows us to perform custom trap trajectories for our nucleus indentation tests while detecting force (Fig. S4 D,E).

After fast indentation perpendicular to the nuclear membrane (~2-3 μm), the trap position was kept constant for 10 s. The same was applied with no bead to account for initial momentum changes arising from dynamic trap location. Upon indentation, an elastic increase in force was followed by a force relaxation that was fitted by the following expression: \( f(t) = A + (B - A)t^{-p}e^{-t/\tau}; \) where A [pN] is the static force given by the residual stress applied onto the nuclear membrane; B [pN] is the peak force; p is the exponent for the initial, short time scale power law decay; and \( \tau \) [s] is the characteristic time for the exponential decay at longer time scale. \( \tau \) agreed to that obtained from a linear fit onto the data plot at semi-logarithmic scale, as \( \tau = -1/m \), where m is the slope of the fitted line. Normalized force relaxation profiles (Suppl. Fig. S4 F,G) were obtained by scaling the force as \( (f(t) - A)/(B - A) \). \( \tau \) and B were obtained for \( N = 21 \) cells in suspension and \( N = 15 \) cells under mechanical confinement (Fig. S4 H,I). Data were processed with custom-written Matlab scripts.

Nuclear deformation was imaged during the trapping routine using a Nipkow spinning-disk confocal imaging platform (Andor DragonFly 502). To block reflections from the trapping laser light, an IR filter was placed in the beam path of the microscope. Nuclear DNA was stained after incubation for 5-10 min in 5 μg/μl Hoechst dye and cells were excited with \( \lambda = 405 \) nm (DNA-Hoechst staining) and \( \lambda = 488 \) nm (myosin II-eGFP). The two laser lines were transmitted through a multi-band dichroic (405-488-561-637 nm, AHF) to simultaneously excite the two fluorophores in the sample. After emission, light is directed into a long-pass dichroic (500 nm), which enables parallel imaging of the two channels using two back-illuminated sCMOS cameras (Sona, Andor) after passing through emission filters 445/46 nm and 521/38 nm, respectively. Image acquisition was performed using Fusion Software and post-processed in Fiji. Tracking of nuclear morphodynamics was carried out by fitting a double sigmoid function over the segment along the indentation direction with a custom written analysis script.

**Raman spectroscopy (RS)** For RS experiments, progenitor cells dissociated from wild type embryos at sphere stage were used and measurements were carried out using re-suspending cells in DMEM media solution. For confinement conditions, a drop of cells was placed between two quartz coverslips (ESCO products, Oak Ridge, NJ). For non-confined cells, a separation of around 100 μm was left between the two coverslips. A total of minimum 25 spectra were obtained from different non-polarized cells for each condition in the cytoplasm or nucleus. The Raman system
(inVia Renishaw, Apply Innovation, Gloucestershire, U.K.) comprises a 532 nm laser (~10 mW) which is focused onto the sample plane using a 60X water immersion Nikon objective (backscattered configuration). The laser spot size was set to 0.8 μm allowing for localized Raman measurements. Raman spectrum were recorded on a deep depletion charge coupled device (CCD) detector (Renishaw RenCam). The recorded Raman spectrum was digitalized and displayed on a PC using Renishaw WiRE software. The spectra were background subtracted with a custom-written Matlab code (see methods in (86)). First, an exploration of the spectral data set was performed using Principal Component Analysis (PCA). Second Multivariate Curve Resolution (MCR) algorithm were performed to extract molecular components from the Raman spectral dataset (spectral profile and abundance in each measured sample). For PCA and MCR analysis, the PLS toolbox in Matlab was used.

**Calcium imaging (related to Suppl. Fig. 3D)** Progenitor cells were added to a Concanavalin A-coated glass bottom dish (0.05 mg/mL by incubation at 31°C for 1.5h) and loaded for 20 min at 28°C with 5 μM of FURA-2 plus a nonionic surfactant (0.02% pluronic F-127) dissolved in DMSO. The cells were then washed before initiating the experiment. The fluorescence signal was measured in a standard bath solution containing 140 mM NaCl, 2.5 mM KCl, 1.2 mM CaCl₂, 0.5 mM MgCl₂, 10 mM HEPES, and 5 mM glucose, pH 7.4, adjusted with NaOH (~320-340 mOsm/l)). The following concentrations of reagents were used: 20μM of YODA1 (to activate Piezo1, Tocris Bioscience), 10 μM GsMTx4 (Tocris). Fluorescence time lapse measurements of intracellular calcium concentrations were obtained using an Olympus IX70 inverted microscope (Hamburg, Germany) with a 40x oil-immersion objective (Olympus). A Polychrome IV monochromator (Till Photonics, Martinsried, Germany) supplied the excitation light (340 and 380 nm), which was directed towards the cells in the field of view by a 505DR dichromatic mirror (Omega Optical, Brattleboro, VT). Fluorescence images were collected by a digital charge-coupled device camera (Hamamatsu Photonics, Hamamatsu City, Japan), after their passage through a 535DF emission filter (Omega Optical), using the AquaCosmos software program (Hamamatsu Photonics). Cytosolic calcium concentration was presented as the ratio of emitted fluorescence after excitation at 340 and 380 nm relative to baseline.

**Data Analysis**

**Myosin II/actin accumulation and relative enrichment at the cortex** Myosin and actin relative cortical accumulation values were quantified from confocal images of non-polarized progenitor cells acquired as described before, using a custom-written script in Matlab (2017b, Mathworks). Quantification was performed in non-polarized cells given that polarized cells present an inhomogeneous cell cortex with an increase in cortical density from the cell front towards the rear. The relative cortical accumulation is defined as: \( \frac{(I_{cortex} - I_{bleb})}{I_{bleb}} \). Cortical intensities were quantified by manually selecting the cortical region where the fluorescence signals are homogenous. The statistics over many cells was used to calculate the median of the mean peak intensities of the cortical regions. The bleb intensity in a single cell is calculated by computing the mean intensity in a manually selected rectangular region in the bleb. Relative myosin II accumulation in vivo was quantified on raw data using Fiji (plot profile tool, Fiji) and for each ratio (peak to averaged cytosolic intensity fluorescence) the distance of the corresponding cell to the yolk margin was measured. Nuclear aspect ratio was calculated in Fiji (measurement tool, Fiji) and for each nucleus the normalized distance to the yolk margin was measured.
Cell size, bleb size and nuclear size estimation

Cell and nuclear diameter measurements were calculated with Fiji (Measure tool, Fiji) from myosin II-GFP confocal images of non-polarized progenitor cells; transmission images were used as control. Bleb sizes were quantified from bright fields movies by segmenting single non-polarized cells and manually selecting the bleb and cell regions using Fiji (Measure tool, Fiji).

Myosin accumulation during mitosis (related to figure 2L)

Myosin II-GFP time lapse movies of cells undergoing spontaneous mitosis under 7 µm confinement were analyzed. For each movie the time point t=0 was defined as the first frame in which the myosin II-GFP signal diffused into the nuclear region (from which it is excluded in interphase cells), indicative of nuclear envelope breakdown. The movies were temporally aligned according to this time frame and successive time points defined according to it. Therefore t=-1min refers to cells in interphase and t>0 to cells undergoing mitosis.

Lap2B-eGFP nuclear analysis

Inner nuclear membrane properties were analyzed using confocal images of unpolarized progenitor cells expressing LapB2-eGFP images in Python using the Scikit-Image library (https://scikit-image.org/). We first determined the inner and outer outlines using the contour detection function and used a median filter for smoothing. From that we determined area, perimeter and the convex image. The ratio between the inner area (\(A_{in}\)) and the associated convex area (\(C_{in}\)) is defined as the invagination ratio, computed as \(1 - A_{in}/C_{in}\).

Lap2B-eGFP curvature analysis

Nuclear envelope fluctuations and bending analysis was performed on Lap2B-eGFP cross-sectional confocal images of unpolarized progenitor cells. The nuclear envelope was manually tracked in Fiji and the discrete x-y positions were further post-processed using a custom made Matlab script. Between each pair of discrete x-y positions a cardinal spline function was interpolated (tension=0) passing through all x-y positions. The resulting spline vector was overlaid to the fluorescence image to manually control the match with the nuclear envelope circumference. The spline was further used to calculate the curvature along the line using 2D bending vectors. The histogram of bending vectors was compared between confined cells (7 µm) and cells in suspension.

Calcium imaging

Progenitor cells derived from WT embryos were stained with Calbryte520 AM, a calcium activity reporter. The cell perimeter was segmented using either myosin II-mCherry, LynTomato or a transmission image and the nucleoplasm manually selected using these same channels. The mean intensity in the Calbryte channel has been measured in Fiji.

Percentage of polarized cells

Percentages of polarized cells were computed from bright field movies as the number of polarized cells/total cells. Stable-bleb polarized cells can be easily distinguished from non-polarized cells due to their different morphology (pear shaped versus rounded blebbing) and thanks to their different morphodynamics behavior (motile versus non-motile).

FRET

RhoA-Fret images of non-polarized progenitor stem cells have been analyzed with the FRET analyzer plugin in Fiji. Donor, acceptor and FRET images are used by the plugin to compute the FRET index and to obtain the FRET image.

Orai-YFP basal accumulation

Orai-YFP intensity line profiles (basal membrane) were exported from Fiji and the intensity level changes were automatically detected in Matlab using the findchangepts function (N=2). For cells cultured in 7 µm confinement a region with a mean intensity increase in the central region of the intensity profile was automatically detected; for cells cultured at 13 µm confinement no change in the intensity could be detected.

cPLA2 translocation ratio (related to Fig S3J)

cPLA2 translocation ratio was quantified only in progenitor stem cells that showed visible translocation via differential intensity of cPLA2-GFP in
between the INM and the nucleoplasm (indicated in the plot by the percentage of translocation). In these cells, the cPLA$_2$-INM translocation ratio was defined as the ratio in between the fluorescence intensity at the INM divided by the intensity in the nucleoplasm.

**Statistical tests** Statistical significance tests were performed with either (1) the two-sample t-test using the ttest2 function in Matlab when the data followed a normal distribution or (2) a non-parametric Kruskal-Wallis test (Matlab) when the data were not normally distributed. In the case of multiple t-tests we corrected for multiple comparisons using the family wise error rate (FWER). Single pairwise datasets were considered non-significant (n.s.) if $p>0.01$. The following significance symbols have been used for the corresponding $p$-values: * $p<0.01$, ** $p<0.001$ and *** $p<0.0001$, while in multiple comparisons we adapted the $p$-values using the FWER correction for each range of significance. All $p$-value indicators and sample sizes (N) were provided in figure legends. Data were obtained from at least three independent experiments. Unless indicated otherwise, points and bars represent mean and standard error of the mean. For the boxplots, the central horizontal line represents the median and the box the interquartile range (with the lower limit the first quartile or 25$^{\text{th}}$ percentile and the upper limit the third quartile or 75$^{\text{th}}$ percentile).
Figure S1. (A) Sketch of plan-parallel cell microconfinement using coverslips with PDMS micro-pillars of desired heights, with plate distance regulated by a pressure controller. (B) Top: representative confocal image of a single progenitor cell expressing myosin II-eGFP under 10 µm confinement. White arrow points at cellular cortex. Bottom: myosin II-eGFP intensity plot along orange and magenta lines showing cortical enrichment and the intensity in the bleb protrusion,
respectively (baseline). The relative cortical intensity was defined as the difference in between the cortical peak intensity and the bleb baseline intensity (as indicated by the black arrow) and was then normalized to the bleb intensity. Scale bar 10 µm. (C) Relative cortical myosin II-eGFP and LifeAct-GFP intensity normalized to the relative cortical suspension value for increasing confinement in un-polarized progenitor stem cells. (D) Exemplary confocal images of progenitor cells expressing Lifeact-GFP under different confinement heights. Scale bars 10 µm. (E) Normalized bleb to cell area for cells cultured in control conditions (Suspension, Susp.), different confinement heights and for 7 µm confinement supplemented with 10 µM Blebbistatin. For all conditions N=35. (F, G) Cell tracks (F), and mean cell velocity (G) for stable-bleb polarized (blue) and blebbing non-polarized (orange) progenitor stem cells cultured in 7 µm confinement. Red lines represent mean and standard error of the mean. (H) Exemplary confocal image of the basal cortex of a stable-bleb cell expressing myosin II-eGFP under 7 µm confinement and associated kymograph along yellow line. Scale bars: left 20 µm, kymograph 10 µm (x) and 10 s (t). Magenta arrow indicates the direction of the cortical retrograde flow and orange arrow the direction of cell migration in both panels. (I) Representative fluorescence image of a progenitor stem cell at 7 µm confinement before release of cell confinement (-30 s) and after confinement release (+30 s). Scale bars 10 µm. (J) Percentage of polarized migratory stable-bleb cells over time after applying 7 µm confinement (t=0) for progenitor stem cells cultured in control conditions (DMEM, blue, N=620) or with Blebbistatin (green, N=486); t_{1/2}~4 min. (K) Relative cortical myosin II intensity for control cells or cells treated with caspase inhibitor in suspension and 7 µm confinement. (L) Relative cortical myosin II intensity for control cells or cells treated with Actinomycin D (transcription inhibitor) in suspension and 7 µm confinement. ***p<0.0001, **p<0.001, * p<0.01, not significant (n.s.).
Figure S2. (A) Representative confocal time lapse images of (from top to bottom) blastula cells (4 hpf) and induced ectoderm, mesendoderm cultured on non-adhesive surfaces and mesoderm cells cultured on fibronectin in 2D versus 3D confinement. Blastula, mesendoderm and ectoderm are non-motile when plated on a non-adhesive 2D substrate and transform to stable-bleb cells if cultured in confinement (7 µm). Mesoderm cells are motile when cultured on fibronectin (mesenchymal migration mode) but transform to stable-bleb cells when confined (7 µm). Dashed lines delimitate the front of stable bleb cells. Scale bars 20 µm. (B) Percentage of stable-bleb cells for induced mesendoderm and ectoderm cells on passivated surfaces and mesoderm cells on fibronectin-coated surfaces for suspension (2D) and 7 µm confinement (3D). N>350 for all
conditions. (C) Cell velocity for mesendoderm and ectoderm cells on passivated surfaces and mesoderm cells on fibronectin-coated surfaces in suspension (2D) and 7 µm confinement (3D). Red lines represent mean and standard error of the mean (sem). N: mesoderm on fibronectin in 2D N=14; mesendoderm 3D N=17; ectoderm 3D N=19; mesoderm on fibronectin 3D N=25. (D) Exemplary in vivo image of embryonic myosin II-eGFP distribution at the lateral margin at 4.5 hpf. The yolk interface (orange dashed line) and embryonic tissue surface (white dashed line) are highlighted. Scale bar 50 µm. (E) Relative cortical myosin II intensity and nuclear aspect ratio as a function of normalized distance from the yolk margin. N=58 cells from 2 embryos. Data represent mean and standard error of the mean. *** p<0.0001, not significant (n.s.).
**Figure S3.** (A) Relative cortical myosin II enrichment under 7 µm confinement for cells cultured on passivated (PLL-PEG) or adhesive surfaces (PLL, fibronectin). (B) Relative cortical myosin II intensity for cells cultured in DMEM (suspension) or under 7 µm confinement for isolated cells (no cell-cell contact, low density) or contacting cells under high cell density and exemplary confocal images of progenitor cells expressing myosin II-eGFP under the indicated conditions (right). (C) Relative cortical myosin II intensity and representative confocal image for cells expressing myosin II-eGFP in control conditions or treated with the cationic mechanosensitive channel inhibitor GsMTx4 and with Gadolinium (inhibitor of stretch-activated ion channels) in suspension and indicated confinement heights. (D) Normalized calcium signal (left) and representative images (right) in isolated progenitor cells under stimulation with the Piezo1 activator YODA (gray, N=39) and in the presence of YODA supplemented with GsMTx4 (yellow, N=46). Data represent mean and standard error of the mean. (E) Representative top view (top) and lateral view (bottom) confocal images of cells expressing myosin II-eGFP and H2A-mCherry cultured in suspension (DMEM). (F) Exemplary Lap2B-eGFP image analysis protocol: (i) raw image; (ii) detection of inner and outer contour (magenta and yellow line); (iii) inner surface area (yellow); (iv) detection of convex area of (iii), (blue). The invagination ratio (IR) is defined as 1 – the ratio in between the yellow and blue area. (G) Nuclear surface and volume for cells in suspension and variable confinement heights. (H) Cell size (left) and nuclear size (right) during embryo development between 3.3 hours post fertilization (hpf) and 10 hpf for undifferentiated progenitor stem cells and induced mesoderm, ectoderm and endoderm progenitor cells (4.7, 8, 10 hpf). In order: N=63 (blastula high); N=60 (blastula sphere); N=39, 71, 38 (mesoderm, endoderm, ectoderm - 4.7 hpf); N=41, 49, 43 (mesoderm, endoderm, ectoderm - 8 hpf); N=43, 65, 63 (mesoderm, endoderm, ectoderm - 10 hpf). (I) Percentage of stable-bleb polarized cells after treatment with 1 µM nocodazole in suspension (unconfined, N>100), 7 µm confined (N>100) and in suspension with the addition of 50 µM LPA (N=40). (J) cPLA2 nucleoplasm to inner nuclear membrane (INM) translocation ratio for cells cultured in DMEM (isotonic), hypotonic condition or under 7 µm confinement with or without the addition of ionomycin. The grey-scaled bar indicates the percentage of cells in which cPLA2-INM translocation is observed and the colormap shows the dynamics of the translocation ratio, defined as the ratio in between the fluorescence intensity of cPLA2-GFP at the inner nuclear membrane and in the nucleoplasm. (K) Representative confocal images of unpolarized progenitor cells expressing cPLA2-GFP cultured in isotonic, hypotonic conditions or for 7 µm confinement with or without the addition of ionomycin. Nuclei with or without cPLA2-INM translocation are shown in columns as indicated. All scale bars 10 µm except (D) 20 µm. ***p<0.0001, **p<0.001, * p<0.01, not significant (n.s.).
Figure S4. Mechanical characterization of cell nuclei in live cells with optical micro-manipulation and spinning-disk confocal microscopy. (A) Schematic showing the trapping beam focused through a microscope objective and captured by a condenser lens (CL) to carry out light-momentum force measurements on an optical trap deforming the nuclear membrane. $F_{\text{mem}}$ and $F_{\text{trap}}$ are the nuclear membrane and trapping forces exerted onto the trapped 1 µm latex bead,
respectively. (B,C) Exemplary fluorescence image of a cell in suspension (B) or confinement (C) expressing myosin II-eGFP and DNA-Hoechst (top) and snapshot prior to (middle) and during (bottom) indentation on the nucleus. The magenta arrow indicates the position of the trapped micro-bead. (D,E) Trap trajectory (left) and nuclear force profile (right, normalized to nuclear indentation) for exemplary force relaxation experiments in cells cultured in DMEM in suspension (D) or under 10 µm confinement (E). (F,G) Normalized and time-shifted nuclear force-relaxations tracks (as in panel D,E) for cells in suspension (F, N=21) and under 10 µm confinement (G, N=15). (H) Boxplot of characteristic relaxation time (τ) obtained from fitting an exponential decay function with plateau offset to the force-relaxation curves shown in (F,G). (I) Boxplot of static force component obtained from fitting an exponential decay to the force-relaxation curves. (J) Exemplary confocal images of cells expressing Lap2B-eGFP (inner nuclear membrane marker) for different time points (time indicates hour after confinement) at 7 µm confinement. Not significant (n.s.).
Figure S5. (A) Relative cortical myosin II enrichment for cells cultured in suspension (DMEM) or supplemented with cPLA$_2$ inhibitor for decreasing confinement heights. (B) Exemplary confocal images of progenitor cells expressing myosin II-eGFP cultured in suspension with cPLA$_2$ inhibitor + 50 µM LPA, Bapta-AM + 50 µM LPA, 50 µM LPA alone or ionomycin (iono). Scale bars 10 µm. (C) Relative cortical myosin II intensity for cells cultured in suspension (DMEM) and upon 50 µM LPA addition in control conditions and for cells treated with cPLA$_2$ inhibitor or Bapta-AM. (D) Relative cortical myosin II enrichment for cells cultured in suspension (DMEM) or supplemented with ionomycin. (E) Components of the Raman spectra associated with arachidonic
acid (AA) used for the quantification of AA production (related to Fig. 2). Raman peaks indicated in red are specific for AA. (F) Scatter plot of Fret index for unconfined (suspension) and confined cells cultured in control (DMEM) conditions or supplemented with cPLA₂ inhibitor. Red lines represent mean and (sem). From left to right: N=20, N=26, N=19, N=21. Right: Representative images showing the Fret index of cells in suspension (Susp., unconfined) and 7 µm confinement in control conditions or supplemented with cPLA₂ inhibitor. Scale bars 10 µm. (G) Representative stitched confocal view of cells expressing myosin II-eGFP cultured in DMEM and confined below a micropillar (approx. height 7 µm) resulting in confined cells and non-confined cells outside the micropillar in close proximity. The magenta line shows the edge of the pillar; the dashed lines indicate the front of stable-bleb polarized/migrating cells and the magenta asterisk a mitotic cell. Scale bar 50 µm. The zoomed view and sketch show adjacent confined and unconfined cells. Scale bar 20 µm. (H) Relative cortical myosin II intensity for confined/non-confined pairs of cells in contact at the micropillar edge, as shown in panel G of this figure. N=27 pairs. Myosin accumulation levels in non-confined cells of this experiment show no significant difference from control cells cultured in suspension. (I) Relative cortical myosin II intensity for cells cultured in control (DMEM) or under different conditions, Thapsigargin (Thapsi.), Calcium free media (no Ca²⁺), 2APB, Bapta-AM and Bapta-AM+cPLA₂ inhibitor. (J-M) Scores of Raman component associated with intracellular AA levels in control DMEM suspension (Iso, N=26) and in the presence of 1 µM ionomycin (Iono, N=30), (K) in control DMEM suspension (Iso, N=24) and in hypotonic condition (Hypo, N=32), (L) in hypotonic condition (Hypo, N=32) and with the presence of cPLA₂ inhibitor (Hypo + cPLA₂-, N=31), and (M) in hypotonic condition (Hypo, N=27) and in hypotonic condition supplemented with ionomycin (Hypo + Iono, N=25). Red lines represent mean and sem. (N) Myosin II cortical accumulation as a function of relative AA Raman score for all tested conditions. Both values are normalized to suspension (isotonic, control). The line indicates a linear regression, R²=0.68 (considering only three last point: R²=0.99). Data points indicate mean and bars standard error of the mean (sem). Scale bars 10 µm except for panel G. ***p<0.0001, **p<0.001, *p<0.01, not significant (n.s.).
Figure S6. (A) Exemplary confocal images of progenitor stem cells expressing Lap2B-eGFP cultured in isotonic (top) and hypotonic (bottom) conditions. Scale bars 5 µm. (B) Boxplot of nuclear invagination ratio (top), nuclear volume (bottom) for cells cultured in isotonic and hypotonic media. (C) Outer nuclear surface for cells cultured at decreasing confinement height or at different osmolarities. (D) Relative bleb to cell size under different osmolarity conditions. N=25 for each condition. (E) Representative confocal images of cells stained with Calbryte520 AM in isotonic condition (suspension), under 0.5x hypotonic shock or confinement at 13/7 µm height. Dashed lines outline the cell and nucleus perimeter. (F) Relative cortical myosin II intensity for progenitor cells cultured in isotonic, hypotonic and hypotonic + ionomycin conditions supplemented with cPLA₂ inhibitor alone or in combination with Bapta-AM. (G) Percentage of polarized stable-bleb cells under reference hypotonic conditions with 1 µM ionomycin (magenta), depletion of extracellular calcium (orange), cPLA₂ inhibition (green) and cPLA₂ + Bapta-AM (blue). Statistical tests with respect to control. N>300 for all conditions. (H) Left: Cell tracks for stable-bleb polarized (blue, motile) and blebbing non-polarized (orange, non-motile) progenitor stem cells cultured in hypotonic media supplemented with 1 µM ionomycin under 16 µm confinement (related to Fig. 4F). Right: Instantaneous cell speed of blebbing non-polarized cells versus polarized stable-bleb cells migrating in 16 µm confinement for cells cultured in hypotonic media with ionomycin. Red lines represent mean and standard error of the mean (sem). (I) Mean square displacement (MSD) analysis of cell tracks related to (H). A persistent random walk model is fit to the data (Fürth formula) for 7 µm confined cells (blue points, velocity and tracks shown in Fig. S1F,G) and hypo+ionomycin treated cells under 16 µm confinement (magenta points, velocity shown in Fig. 4F), with persistence time of $P_t=1.9$ min (2.8 min), respectively. (J) Mean calcium intensity with respect to the normalized nuclear contact area for isotonic condition (control yellow, N=100), hypotonic shock (red, N=39), 13 µm confinement (light blue, N=44) and 7 µm confinement (dark blue, N=33). The nuclear contact area is normalized to the cross-sectional area for each cell. Points represent mean and bars the standard error of the mean. (K) Representative confocal images of cells expressing STIM-CFP and Orai-YFP at 13/7 µm confinement. Cross-sections and basal membrane sections are shown along with the transversal view for 7 µm confinement. Magenta asterisks indicate the nucleus and the arrow points at the region of the basal membrane for 7 µm confinement where Orai-YFP accumulates (ER-PM contacts). (L) Orai-YFP intensity at the basal membrane for cells at 13/7 µm confinement. Each line corresponds to one cell (13 µm: N=6, 7 µm: N=14). Black lines represent mean values. (M) Representative in vivo image of a motile stable-bleb cell (dashed red line) in a zebrafish embryo at blastula stage (4 hpf) injected with hypotonic media supplemented with ionomycin (10 µM). Asterisk denotes cell front. Scale bar 10 µm in all panels apart from panel transversal views of panel K and panel A 5 µm. ***p<0.0001, **p<0.001, *p<0.01, not significant (n.s.)
Figure S7. (A) Illustration of the mechano-transduction pathway that translates nucleus deformation into myosin II activation and re-localization to the cortex resulting in an increase in cortical contractility and regulating morphodynamic migration plasticity. (B) Schematic representation of inner nuclear membrane (INM) unfolding (top to bottom) and intracellular calcium levels (left to right) depending on the type of physical cell deformation (anisotropic cell compression and hypotonic swelling leading to isotropic cell stretching) and ectopic calcium regulation via ionomycin.
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**Supplementary Table S1.** Table summarizing different inhibitors and methods used in order to identify and characterize the mechanotransduction pathway responsible for mechanosensitive cortical actomyosin regulation under cell shape deformation.
Captions for Movies S1 to S9

**Movie S1.** Adaptive myosin II dynamics upon cell confinement and role of cPLA$_2$ in cell mechanotransduction (related to Fig. 1 and Fig. 3). Time lapse confocal fluorescence movies of progenitor cells expressing Myl12.1-eGFP (myosin II) on non-adhesive substrates (PLL-PEG) cultured in DMEM media in suspension (left), under 7 µm confinement in control condition (middle) or supplemented with 1 µM cPLA$_2$ inhibitor. Upon confinement, myosin II accumulates at the cortex in control condition and the accumulation is blocked by the inhibition of cPLA$_2$.

**Movie S2.** Cortex remodeling during amoeboid stable-bleb cell transformation and cortical retrograde flow in stable-bleb polarized cells (related to Fig. 1 and Suppl. Fig. 1). (I) Time lapse confocal fluorescence movie of Myl12.1-eGFP (myosin II) localization during stable-bleb transformation of a progenitor cells under 7 µm mechanical confinement: a non-polarized and non-motile blebbing cell spontaneously polarizes into stable-bleb cell upon mechanical confinement and initiates cell migration. (II) Time lapse confocal fluorescence movie of the basal cortex of a stable-bleb polarized cell expressing Myl12.1-eGFP (myosin II) under 7 µm mechanical confinement. Myosin II shows a cortical density gradient and retrograde flow (magenta arrow, opposite to the direction of migration indicated by the orange arrow).

**Movie S3.** Adaptive myosin II morphodynamics under confinement (related to Fig. 1-2 and Suppl. Fig 2). Confocal fluorescence time lapse movie of pluripotent progenitor cells (I, sphere stage) on non-adhesive substrate (PLL-PEG); ectoderm cells (II) on non-adhesive substrate (PLL-PEG); and mesoderm cells on fibronectin (III) for 2D substrate (left) and 3D mechanical confinement (7 µm, right). Pluripotent and ectoderm cells are non-motile when plated on 2D substrate and transform into a stable-bleb polarized and motile phenotype upon confinement. Mesoderm cells plated on a 2D fibronectin substrate migrate with a mesenchymal phenotype and transform to stable-bleb amoeboid migration upon mechanical confinement.

**Movie S4.** Myosin II accumulation is reversible (related to Fig. 1). Confocal fluorescence time lapse movie of progenitor cells expressing Myl12.1-eGFP (myosin II) during confinement (10 µm) and subsequent release (i) and for confined cells squeezing out from underneath a micropillar (“escape reflex”, ii). Myosin II is rapidly re-localized from the cortex to the cytoplasm upon confinement release.

**Movie S5.** Mesenchymal-to-amoeboid transition: mesoderm cells transform to stable-bleb cells upon mechanical confinement (related to Fig. 2). Time lapse confocal fluorescence movie of mesendoderm-induced progenitor cells expressing Myl12.1-eGFP (myosin II) plated on fibronectin substrate showing mesenchymal migration (white asterisk indicates migrating cell,
white arrow points at cell protrusions). Upon confinement (7 µm), cells rapidly accumulate myosin II at the cortex and transform to stable-bleb polarized cell (magenta asterisks point at cell front).

**Movie S6.** The inner nuclear membrane is unfolded under mechanical shape deformation in confinement and the unfolding is stable over time (related to Fig. 3). Time lapse confocal movies of progenitor cells stained with Lap2b-eGFP (i) in suspension, (ii) during confinement and (iii) under 7 µm confinement. The time lapse in (iii) was acquired 1h after mechanical confinement.

**Movie S7.** Adaptive Myosin II dynamics under hypotonic conditions or ionomycin treatment (related to Fig. 7). Confocal fluorescence time lapse movie of progenitor cells expressing Myl12.1-eGFP (myosin II) on non-adhesive substrate (PLL-PEG) in isotonic media (control, DMEM), upon hypotonic shock (adding milliQ water, 0.5x shock) and upon the addition of 1 µM ionomycin. Hypotonic treatment is followed by a myosin II accumulation at the cortex; further addition of ionomycin leads to a pronounced myosin II accumulation and triggers cell polarization.

**Movie S8.** Cell dynamics under hypotonic conditions and ionomycin treatment (related to Fig. 7). Bright field time lapse movie of progenitor cells cultured in isotonic condition (top-left), hypotonic media (top-right), hypotonic media supplemented with 1 µM ionomycin in suspension (bottom-left) or under 16 µm confinement (bottom-right). Hypotonic conditions supplemented with 1 µM ionomycin lead to rapid cell polarization and induce a stable-bleb cell transformation (bottom-left), but cells can migrate only under confined conditions (bottom-right).

**Movie S9.** The endoplasmic reticulum (ER) is immobilized under the nucleus in deformed cells under high confinement (related to Fig. 7 and Suppl. Fig. 6). Time lapse TIRF movie of progenitor cells stained with ER tracker green under increasing mechanical confinement. At low mechanical confinement, the ER is mobile in the nucleus-plasma membrane contact area, while for increasing confinement the ER is immobilized in the region underneath the nucleus. The yellow line marks the nuclear area (visualized from bright field imaging).
Supplementary References