1 Self-organization of Tissue Growth by Interfacial Mechanical

2 Interactions in Multi-layered Systems

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- 4 Tailin Chen^{1,4,#}, Yan Zhao^{3,#}, Xinbin Zhao¹, Shukai Li¹, Jialing Cao¹, Jing Du^{1,4,6*},
- 5 Yanping Cao^{2,*}, and Yubo Fan^{1,4,5*}
- 6

7 ¹ Key Laboratory for Biomechanics and Mechanobiology of Chinese Education

- 8 Ministry, School of Biological Science and Medical Engineering, Beihang University,
- 9 Beijing 100191, China
- 10 ² Institute of Biomechanics and Medical Engineering, Department of Mechanical
- 11 Engineering, School of Aerospace, Tsinghua University, Beijing 100084, China
- ³ State Key Laboratory of Advanced Design and Manufacturing for Vehicle Body,
- 13 College of Mechanical and Vehicle Engineering, Hunan University, Changsha 410082,
- 14 China
- ⁴ Beijing Advanced Innovation Centre for Biomedical Engineering, Beihang University,
- 16 Beijing 100191, China
- ⁵ Key Laboratory of Human Motion Analysis and Rehabilitation Technology of the
- 18 Ministry of Civil Affairs, National Research Center for Rehabilitation Technical Aids,
- 19 Beijing 100176, China
- 20 ⁶ Lead Contact
- 21 [#] These authors contributed equally
- 22
- 23 * Correspondence: dujing@buaa.edu.cn (J.D.), caoyanping@tsinghua.edu.cn (Y.P.C),
- 24 yubofan@buaa.edu.cn (Y.B.F.)
- 25

26 Abstract

27 Morphogenesis is a spatially and temporally regulated process involved in various 28 physiological and pathological transformations. In addition to the associated 29 biochemical factors, the physical regulation of morphogenesis has attracted increasing 30 attention. However, the driving force of morphogenesis initiation remains elusive. Here, 31 we show that during the growth of multi-layered tissues, morphogenetic process can be 32 self-organized by the progression of compression gradient stemmed from the interfacial 33 mechanical interactions between layers. In tissues with low fluidity, the compression gradient is progressively strengthened during growth and induces stratification by 34 35 triggering symmetric-to-asymmetric cell division reorientation at the critical tissue size. 36 In tissues with high fluidity, compression gradient is dynamic and induces cell junction 37 remodelling regulated cell rearrangement leading to 2D in-plane morphogenesis instead 38 of 3D deformation. Morphogenesis can be tuned by manipulating tissue fluidity, cell 39 adhesion forces and mechanical properties to influence the progression of compression 40 gradient during the development of cultured cell sheets and chicken embryos. Together, the dynamics of compression gradient arised from interfacial mechanical interaction 41 42 provides a conserved mechanism underlying morphogenesis initiation and size control 43 during tissue growth.

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47 Key words: compression gradient, interfacial interaction, morphogenesis,48 biomechanics, self-organization, tissue fluidity

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50 Introduction

51 Morphogenesis is a common process that occurs widely in embryonic development, 52 tissue regeneration and cancer progression. The orchestration of morphogenetic 53 processes is complex and involves spatial and temporal regulation by biochemical 54 factors (e.g., cell polarity signals and morphogen gradient) and physical factors (Gallet, 55 2011; Heisenberg and Bellaiche, 2013; Nishimura and Takeichi, 2008). Emerging 56 studies have revealed that proper morphogenesis relies on the mechanical force of the 57 cells and their environment. For example, apical constriction of cells caused by the 58 contractility of myosin (Martin et al., 2009), cell junction remodelling (e.g., cell 59 intercalation) (Rauzi et al., 2010) and tissue stiffness-dependent cell migration (Barriga 60 et al., 2018) have been reported as important mechanisms in tissue shaping during 61 embryonic development. These studies indicate the essential functions of cellular and 62 molecular mechanics in the progression of morphogenesis. However, the upstream 63 events, especially the initial driving forces of morphogenesis, remain unknown.

64 Most biological tissues have multi-layered structures. The interactions between 65 layers are essential in tissue homeostasis maintenance and morphogenesis during embryonic development and pathological progression (Bhowmick and Moses, 2005; 66 67 Carvalho and Heisenberg, 2010; Lilly, 2014). For example, the interaction between 68 cancer cells and their adjacent stroma plays a key role in the progression of the diseases, 69 including tumour invasion (Gupta and Massague, 2006; Mueller and Fusenig, 2004). In addition to biochemical communications, increasing evidence has shown the 70 71 essential role of physical interactions between adjacent layers in the regulation of 72 morphogenesis (Bailles et al., 2019; Barriga et al., 2018; Carvalho et al., 2009; Munster 73 et al., 2019). For instance, follicle formation in chicken embryos is initiated by

74 mechanical forces transduced from the dermal layer to the epidermal layer (Shyer et al., 75 2017). The villi of human and chicken guts are formed by the compressive stresses 76 generated by smooth muscle layers on the endoderm and mesenchyme layers (Shyer et 77 al., 2013). These studies reveal that morphogenesis processes are dependent on the 78 mutual collaboration and mechanical compatibility of multiple layers. In this sense, it 79 is necessary and important to address the general mechanism underlying the initiation 80 of morphogenesis during the growth of various multi-layered tissues. 81 Here, by combining *in vitro* and *in vivo* biological experiments, theoretical analysis 82 and numerical simulations, we report a general mechanism underlying the initiation of

83 morphogenesis driven by the progression of compression gradient stemmed from the

84 interfacial mechanical interactions between growing tissue layers.

85

87 **Results**

88 **Progressive compression gradient is strengthened in epidermal layer during**

89 chicken feather follicle morphogenesis

90 During the development of avian skin, the feather follicles develop by the 91 stratification of single-layered epidermis (Mayerson and Fallon, 1985). In the in vitro 92 culture of chicken skins, we found that the feather primordia emerged at Day 2 in 93 cultured epidermis combined with dermis after isolation from HH30 stage embryos 94 (Figure 1a-c). However, when epidermal cell sheet was isolated and cultured alone 95 (without dermal cell layer), it failed to stratify and no primordium was formed (Figure 96 1b and c), indicating that the interaction between epidermal and dermal layers is 97 essential for the morphogenetic process of epidermal cell sheet. Moreover, during the 98 evolution of epidermis from monolayer to multilayer, the shapes of epidermal cells 99 showed significant alteration from flat to columnar and correlated with the stages in 100 embryo development and with the increased epidermal cell layer number (Figure 1d 101 and e). The deformation degree of epidermal cells was gradually declined with the 102 increased distance to the primordium center (Figure 1f and g). These experimental 103 observations indicate that in the beginning of follicle morphogenesis, a local 104 compression gradient was progressively strengthened in the epidermal cell sheet 105 accompanied with the 3D deformation of epidermis. According to our previous studies 106 about the surface wrinkling pattern formation in a non-living chemical film/substrate 107 composite soft material, the progression of compression gradient could be generated by 108 mismatch deformation between adjacent layers through interfacial mechanical 109 interactions, which leads to intriguing morphogenesis (Han et al., 2015; Zhao et al., 110 2015a) (Figure S1). Indeed, it has been reported that during the development of chicken skins, dermal cells form aggregations and compress the adjacent epidermal cells to form follicle primordium (Ho et al., 2019; Shyer et al., 2017). Thus, we hypothesize that the mismatch deformation between epidermal and dermal layers by dermal cell aggregation generates compression gradient through interfacial mechanical interactions and causes stratification of epidermis. However, whether tissue stratification could be triggered by the mechanical interactions between adjacent layers during tissue growth remains elusive.

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3D tissue morphogenesis could be self-organized at a critical size in multi-layered system

121 To study the role of interfacial mechanical interactions between layers in the 122 initiation of tissue morphogenesis, we developed a simple film/substrate system 123 composed of a freely growing monoclonal cell sheet and extracellular matrix (ECM) (Figure 2a). To ensure the occurrence of stratification, cells without contact inhibitory 124 125 properties were studied. During the continuous live imaging, the emergence of a 2D 126 monolayer-to-3D multilayer transition was stably observed at a critical tissue size in a 127 wide variety of cell types including skin-derived cells (B16F10). The critical size was 128 relatively constant for a given cell type, indicating a self-organized mechanism of 129 morphogenesis during cell sheet growth (Figure 2a-e). In addition, similar phenomenon 130 was observed in different types of ECM as well as altered substrate stiffness (Table S1). 131 The behaviours of individual cells during cell sheet growth prior to the 3D 132 morphogenesis transition were examined using a holographic imaging cytometer. 133 Although the cell proliferation rate was negligibly altered during cell sheet expansion 134 (Supplementary Figure S2a), the average area of individual cells significantly decreased, and the average cell thickness concurrently increased during cell sheet 135

136 growth, indicating significant cell deformation (Figure 2f-h). Moreover, the largest cell 137 deformation was observed at the central region of the cell sheet (Figure 2i-l). Single 138 cell tracing of cell deformation also suggested significant compression of cells in the 139 central region of cell sheet (Supplementary Figure S3). This cell deformation behaviour 140 indicates that during cell sheet growth, a compression gradient within the cell sheet 141 emerges. To further verify the compression gradient, a scratching experiment was 142 performed crossing the center to the edge of the cell sheet. After scratching, cells in the 143 central region showed much faster expansion and migration speed compared with 144 peripheral cells, indicating the release of compressive strain (Figure S4). Similar 145 mechanical gradients have also been obtained by Traction Force Microscope (TFM) 146 and Monolayer Stress Microscopy (MSM) in previous studies (Perez-Gonzalez et al., 147 2019; Puliafito et al., 2012; Trepat et al., 2009).

148 To investigate the generation mechanism of compression gradient in the cell sheet 149 during growth, we performed theoretical analysis. As illustrated in Figure 3a, the cell 150 sheet is considered as continuum material. During cell sheet expansion caused by cell 151 proliferation, interfacial shear stress (ISS) would be generated between the cell sheet 152 and substrate layers. Since the direction of ISS is contrary to the relative motion 153 between the adjacent layers, the cell sheet is subjected to ISS directed toward the center, 154 which is consistent with the previous observations by TFM (Trepat et al., 2009). Thus, 155 cells in the central region of the cell sheet would sustain higher level of compression 156 than those in other regions (Figure 3a and Supplementary Mechanical Modelling), 157 which is confirmed by the experimental observations (Figure 2i-l). The elastic strain 158 energy stored in the cell would increase with the expansion of cell sheet. When the elastic strain energy in the cell is small, interfacial normal adhesion would impose 159 160 restriction on delamination, making the cell monolayer grow in plane. Thus, higher

161 compressive strain would be generated further. When the compressive strain reaches a 162 critical value, elastic strain energy stored in the cell may be greater than the energy of 163 interfacial normal adhesion and led to the occurrence of interfacial delamination. In this 164 critical condition, stratification may happen, and the elastic strain energy can be 165 released (Supplementary Mechanical Modelling). Indeed, we found that when cell 166 sheet grew beyond the critical size, amounts of cells in the central region were 167 delaminated from the substrate (Figure 3b). Theoretical analysis can also give valid 168 quantitative predictions of experiments. First, theoretical results of the distribution of 169 cell areas agree well with the experimental results during cell sheet expansion (Figure 170 3b). According to the theoretical analysis, the critical size of the cell sheet for 3D 171 morphogenesis depends on the mechanical properties of cell sheet and substrate and 172 interactions between them. This is confirmed by finite element simulations which well 173 resembled the morphogenesis at the cell sheet center and also indicated that, when the 174 compressive strain exceeded the critical value, alterations in the cell-substrate 175 interactions could significantly affect the compression gradient during cell sheet growth 176 (Figure 3d and Supplementary Mechanical Modelling). Moreover, experimentally 177 manipulating cell-substrate interactions using the integrin inhibitor RGD significantly 178 attenuated the compression gradient and morphogenesis (Figure 3e-g). In addition, 179 inhibition of the contractility of the cell sheet using the cytoskeleton inhibitor 180 blebbistatin or myosin shRNA significantly disrupted the compression gradient and 181 morphogenesis (Figure 3e-g and Supplementary Figure S5). Based on the theoretical 182 analysis, adhesion between cells may also affect the compression gradient. Consistently, 183 inhibition of the cell-cell adhesion force by an E-cadherin neutralizing antibody also 184 reduced the maximum compressive strain in the cell monolayer and inhibited the 185 morphogenetic process (Figure 3e-g). These results show that the emergent 3D

186 morphogenesis of a cell sheet/substrate system could be physically triggered by
187 interfacial mechanical interactions between adjacent layers during growth.

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189 Critical Compression triggers symmetric-to-asymmetric reorientation of cell

190 division

191 We proceeded to investigate the biological mechanism of the cell layer number 192 increase in the central region of the cell sheet induced by cell layer/substrate interfacial 193 interaction. We found that, in contrast to symmetric (parallel) cell division in the 194 peripheral region of large cell colonies, abundant asymmetric (oblique or perpendicular) 195 cell division was observed in the central cells which bore significant compressive 196 strains (Figure 4a-f). The orientation of the division plane was closely correlated with 197 cell thickness (Figure 4f). Moreover, after release of compressive strain by cell 198 scratching, the reorientation of cell division was diminished (Figure S6), indicating the 199 effect of compression gradient on cell division orientation. This asymmetric cell 200 division ultimately led at least one daughter cell to locate at the top layer of the cell 201 sheet (Figure 4g and Supplementary Movies 1 and 2). The cell division reorientation 202 induced by compression gradient was also confirmed by the subcellular localization of 203 the nuclear-mitotic apparatus protein (NuMA), which is involved in the orchestration 204 of mitotic spindle positioning (Pirovano et al., 2019) (Figure 4h). Moreover, spindle-205 rocking experiments indicate that cells in the central region showed a significantly 206 higher level of metaphase plate oscillations which is always associated with asymmetric 207 cell division (Haydar et al., 2003) (Figure 4i). These results suggest that compression 208 gradient induced by interfacial interaction between layers triggers cell division 209 reorientation from symmetric to asymmetric leading to tissue stratification.

211 Tissue fluidity controls 2D/3D morphogenetic transition by regulating

212 compression gradient progression

213 Our experimental results show large variations in the critical sizes of cell sheets for 214 3D morphogenesis among different cell types with critical diameter ranging from 356 215 mm in HeLa to 1997 mm in U2OS cells (Figure 2e). Notably, some cell types, such as 216 Madin-Darby canine kidney (MDCK) cells, did not display stratification during our 217 observations. To further investigate the intrinsic properties of tissue layers that affect 218 the emergent morphogenesis, we compared cell behaviours during cell sheet growth 219 between HeLa and MDCK cells. First, in order to exclude the influence of contact 220 inhibition on MDCK cell behaviour, we measured cell proliferation rate and cell density 221 alterations during cell sheet growth. As shown in Supplemental Figure S2b and d, 222 MDCK cells did not show contact inhibition during our observation. Moreover, 223 according to the previous report that MDCK cell sheet did not display contact inhibition 224 until a critical size of approximately $2x10^6 \ \mu m^2$ (Puliafito et al., 2012). In our experiment, the cell sheet size of MDCK colony was below $1 \times 10^5 \ \mu m^2$. Thus, the 225 226 differential behavior of MDCK cell sheet was not caused by contact inhibition. 227 Interestingly, we found that, instead of 3D morphogenesis, MDCK cell sheet showed 228 dramatic alterations in its in-plane shape during growth, with dynamically organized 229 protrusions at cell sheet edges, indicating an emergent 2D morphogenesis 230 (Supplementary Figure S7). While few differences in the proliferation rate were 231 observed between these two cell types (Supplementary Figure S2a and b), expansion 232 rate was significantly higher in MDCK cell sheet than in HeLa cell sheet during growth 233 (Supplementary Figure S2c and d). Importantly, in contrast with the HeLa cell sheet 234 behaviour, the MDCK cell sheet did not show progressive compression gradient 235 strengthen during growth (Figure 5). Moreover, while the individual cell area in the 236 HeLa cell sheet was relatively steady, MDCK cells exhibited strong area fluctuation 237 during cell sheet growth (Figure 6a and b). Particle imaging velocimetry (PIV) analysis 238 showed that during cell sheet expansion, MDCK cells displayed rapid collective 239 cellular motion with a higher overall cell velocity (root-mean-square, rms velocity, v_{rms}), 240 indicating a fluid-like state (Garcia et al., 2015). In contrast, the collective behaviour of 241 HeLa cells showed slower cell motion, indicating a relatively solid-like state (Figure 242 6c, d and Supplementary Movie 3). The fluidity of these two cell types was further 243 confirmed by the cell shape index p_0 , the median ratio of the perimeter to the square 244 root area of the cells. According to the vertex model, if the cell shape index of the 245 system increases to $p^*_0 \approx 3.81$, a transition from a jammed, solid-like state to an 246 unjammed, fluid-like state occurs (Bi et al., 2015). Over the course of cell sheet growth, 247 the average cell shape index of MDCK cells was constantly above 3.81, whereas HeLa 248 cells approached the jamming threshold $p*_0$ (Figure 6e and f). Moreover, the cell shape 249 index was lower in the central region than in the peripheral region of the HeLa cell 250 sheet (Supplementary Figure S8). At the edges of MDCK cell sheet, especially during 251 the formation of protrusions, significant collective cell migration (Figure 5a and 252 Supplementary Movie 4) and abundant cell intercalations such as T1 transitions and 253 rosettes formation were observed (Figure 7a and b). Cell intercalation is reported as a 254 mechanism for driving tissue extension in embryonic development (Guillot and Lecuit, 255 2013) and could be controlled by external constraints acting on the tissue (Aigouy et 256 al., 2010). In our experiment, most of cell intercalation was observed in cells with relatively smaller areas (less than 300 μ m²), indicating a high level of compressive 257 258 strain (Figure 7c). Moreover, the cell junction remodelling during intercalation process 259 was closely correlated with the 2D deformation of cell sheet, indicated by the small 260 angles between new junctions and tissue protrusion directions (Figure 7d) and higher

261 tissue elongation rate at the direction of new junctions (Figure 7e). To investigate the 262 effect of the tissue fluidity on the 2D/3D morphogenetic transition, cellular migration ability was inhibited using the small GTPase Rac1 inhibitor NSC23766 (Raftopoulou 263 264 and Hall, 2004). The results showed that Rac1 inhibitor treatment significantly reduced 265 tissue fluidity (Figure 7f and g and Supplementary Movie 5) and induced higher level 266 of compressive strain in MDCK cell sheet (Figure 7h and i). Importantly, treatment of 267 NSC23766 significantly promoted the emergence of tissue stratification (Figure 7j). In 268 comparison, inhibition of Rho-associated, coiled-coil containing protein kinase (ROCK) 269 by Y27632 slightly enhanced tissue fluidity of MDCK cells (Figure 7f and g and 270 Supplementary Movie 5) and had little effect on morphogenesis (Figure 7h-j). These 271 results suggest that the morphogenesis driven by interfacial mechanical interactions 272 between tissue layers is dependent on tissue fluidity that higher level of tissue fluidity 273 may prevent the storage of the compressive strain energy through in-plane cell motion. 274

275 Progression of compression gradient contributes to epidermal cell stratification 276 during chicken skin development

277 Asymmetric cell division is a common mechanism involved in tissue stratification 278 and cell fate differentiation during embryogenesis and cancer progression (Lechler and 279 Fuchs, 2005; Neumuller and Knoblich, 2009). There is evidence that during the 280 development of chicken skins, the deformation of basement membrane between 281 epidermis and dermis by the aggregation of dermal cells plays crucial role for the 282 initiation of follicle primordium formation (Ho et al., 2019; Shyer et al., 2017). 283 According to our results, tissue stratification could be initiated by the mechanical 284 interaction at interface between adjacent layers through compression-induced cell 285 division reorientation. Thus, we proceeded to verify whether this mechanism is applied 286 to the stratification of epidermis during chicken feather follicle morphogenesis. First, 287 we compared the cell division orientation of the epidermis in single-layered stage and 288 multi-layered stage during skin development. We found that when epidermis was 289 stratified, most of the mitotic spindles of epidermal cells were reoriented from 290 symmetric to asymmetric and the spindle-axis angle was closely correlated with cell 291 deformation degree (Figure 8a-c). Moreover, reinforcing the progression of 292 compression gradient by reducing tissue fluidity using Rac1 inhibitor significantly 293 promoted the progression of epidermis stratification and follicle formation, indicated 294 by the epidermal cell layer number and pattern geometry of follicle primordium (Figure 295 8d-i). Reducing tissue fluidity also increased the nuclear localization of β-catenin 296 protein, indicating the promotion of follicle cell fate determination (Figure 8h and i). In 297 addition, disrupting the interfacial mechanical interaction between epidermal and 298 dermal layers by inhibiting epidermal cell/base membrane adhesion using integrin 299 inhibitor largely attenuated the morphogenesis of primordium (Supplementary Figure 300 S9). Thus, our findings suggest a model for the initiation of epidermis stratification 301 during feather follicle formation that dermal cell aggregation induces mismatch 302 deformation between epidermal layer and basement membrane which generates 303 progressive compression gradient in epidermal layer and the latter triggers cell division 304 reorientation leading to epidermis stratification.

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306 Discussion

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308 In this work, we propose a self-organization mechanism for the initiation of 309 morphogenesis during tissue growth in which a progressive compression gradient 310 caused by macroscopic interfacial mechanics drives the initiation of morphogenesis at 311 certain developing stages in multi-layered tissues. Our experiments and theoretical 312 analysis show that mismatch deformation between adjacent layers during tissue growth 313 induces progression of compression gradient by interfacial interactions which triggers 314 cell delamination and cell division reorientation in solid-like state tissues and ultimately 315 induces 3D deformation and tissue stratification. Meanwhile, in fluid-like state tissues, 316 the interaction between adjacent layers induced dynamic progression gradient leading 317 to 2D tissue deformation instead of 3D deformation (Figure 8j). We also proposed the 318 crucial role of progressive compression gradient in epidermis generated by the 319 interaction between epidermal and dermal layers in the initiation of epidermal cell 320 stratification during chicken skin development. Interestingly, in zebrafish embryos, the 321 friction force between mesoderm and neurectoderm layers during cell moving has been 322 reported to be a key determinant in the positioning of neural anlage (Smutny et al., 323 2017). Thus, interfacial mechanical interaction between adjacent layers may be a 324 conserved force origin during embryonic development. Resembling to the instability 325 process in film/substrate material system, we regard the cell sheet as a film, which is 326 subjected to the interfacial shear stress during cell sheet expansion. Thus, the growing 327 cell sheet will be compressed and buckled. We found that cell-cell adhesion, cellsubstrate adhesion, and cell fluidity all affected the degree of cell sheet compression 328 329 and tissue stratification. Interestingly, recent studies regard the process of cell 330 aggregation and layering as the wetting problem and also reveal that inhibiting cell 331 contraction, cell-to-cell adhesion, and cell-to-substrate adhesion have significant

influences on the dewetting process (Perez-Gonzalez et al., 2019; Ravasio et al., 2015).

333 Morphogenetic processes always occur at certain developmental stages during tissue 334 growth. In Drosophila wing disc development, different cell proliferation rates induce 335 mechanical strain to shape tissues and control tissue size by regulating proliferation and 336 division orientation (Legoff et al., 2013; Mao et al., 2013). In our experimental model, 2D and 3D morphogenetic process emerges at the critical tissue size, which is 337 338 determined by the critical value of the compressive strain generated by interfacial 339 interaction during tissue growth. This macroscopic mechanical dissipation provides a 340 possible regulatory mechanism for the spatiotemporal control of morphogenesis and 341 size control of growing tissues. It has been recognized that crowding is a common 342 phenomenon in physiological and pathological processes and regulates tissue 343 homeostasis maintenance (Eisenhoffer et al., 2012; Marinari et al., 2012). A recent 344 study reports that during the development of zebrafish heart, proliferation-induced 345 crowding leads to tension heterogeneity that drives cell stratification (Priva et al., 2020). 346 According to our results, critical compressive strain has profound effects on cell 347 behaviours, including cell delamination, cell division reorientation and in-plane cell 348 rearrangement, which all contribute to tissue shaping, homeostasis maintenance and 349 size control. Moreover, in our model experiment, the compression gradient is 350 autonomously generated by tissue growth and therefore can well simulate the in vivo 351 crowding conditions with the controlled crowding levels.

As an important characteristic of tissues, fluidity (jamming/unjamming state) is determined by collective cell motion and dynamically altered during tissue growth (Sadati et al., 2013). Recent studies suggest that tissue fluidity plays crucial roles in homeostasis maintenance and tumour invasion (Garcia et al., 2015; Miroshnikova et al., 2018; Park et al., 2016). Saadaoui *et al.* reported that myosin contractility-induced

357 global tissue flow contributed to morphogenesis during avian gastrulation (Saadaoui et 358 al., 2020). In our work, we found that tissue fluidity could affect the local accumulation 359 of compressive strain under interfacial interaction between adjacent layers, which 360 finally regulates tissue deformation patterns (2D or 3D). These findings may provide 361 insights into the regulatory mechanism underlying morphogenesis by tissue fluidity.

There is no doubt that mechanical forces play essential roles in the regulation of 362 363 morphogenesis. The role of cell-scale forces generated by cytoskeleton contraction and 364 transmitted by cell adhesions has been intensively studied in regulating tissue 365 morphogenesis (Heisenberg and Bellaiche, 2013). However, the upstream regulation mechanism of these molecular machines, especially at the tissue scale, is rarely 366 367 investigated. Our results reveal that progression of compression gradient controlled by 368 the tissue-scale interfacial shear stress has propounding effects on local cell 369 delamination, reorientation of division plane and cell intercalation to initiate 370 morphogenesis. Moreover, the interfacial mechanics could be stemmed from mismatch 371 deformation during cell migration, aggregation, proliferation, etc. Thus, this scale-372 spanning mechanical loop from the cell scale to the tissue scale and then returns to the 373 cell scale may be a fundamental self-organized mechanism during morphogenetic 374 process in growing multi-layered tissues. Further study is needed to elucidate the 375 molecular mechanotransduction mechanism underlying the regulation of these cell 376 behaviours by the progression of compression gradient under interfacial mechanical 377 interactions.

Taken together, our findings unveil a self-organized mechanism that drives the initiation of morphogenesis in growing multi-layered tissues. These results also open up a new avenue that tissue-scale forces regulate cell behaviours, which could in turn facilitate tissue self-organization.

383 Materials and Methods

384 Cell culture and Immunofluorescence

| 385 | HeLa, HepG2, Madin-Darby canine kidney (MDCK), MDA-MB-231, U2OS, B16F10 |
|-----|-----------------------------------------------------------------------------------------------|
| 386 | cells were cultured in DMEM medium (containing with 4.5 g/L glucose, L-glutamine, |
| 387 | and sodium pyruvate) supplemented with 10% FBS (Life technologies, CA, USA), and |
| 388 | 100 IU/mg penicillin-streptomycin (Life technologies, CA, USA) and 1 % (v/v) NEAA |
| 389 | (Life technologies, CA, USA). For the experimental treatment, we use clones of cells |
| 390 | grown for the same amount of time. The pharmacological agents were added including |
| 391 | Blebbistatin (sigma, 25 μ M), RGD (Abcam, 50 μ g/ml), E-cadherin neutralizing |
| 392 | antibody (Biolegend, 10 $\mu g/ml$), NSC23766 (Abmole, 20 μM), Y27632 (Abmole, 20 |
| 393 | μ M) if applicable. Images were taken on the Nikon microscope. Cells were grown at |
| 394 | 37 °C in an incubator with 5% CO ₂ . Cells grown on glass bottom dishes were fixed |
| 395 | with 4% paraformaldehyde for 10 min at room temperature. Incubate the cells with 5% |
| 396 | BSA, in PBST (PBS+0.1% Tween 20) for 2 hours. Cells were incubated with primary |
| 397 | antibodies at the optimal concentrations (according to the manufacturer's instructions) |
| 398 | at 4 °C overnight. After washing, cells were incubated for 2 hours with secondary |
| 399 | antibodies: 488/568/633 IgG (H+L) and/or Alexa Flour 568/647 phalloidin (Invitrogen) |
| 400 | for 1 hour. Cell nucleus were stained with DAPI (4',6-diamidino-2-phenylindole, |
| 401 | Invitrogen) for 10 min at room temperature. Confocal images were taken on the Leica |
| 402 | microscope equipped with a 10x, 40x or 63x objective. Experiments were replicated at |
| 403 | least three times. |
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407 Live imaging

408 Live cell imaging was performed in Leica microscope or HoloMonitor M4, enclosed in 409 an incubator to maintain the samples at 37 °C and 5% of CO₂ throughout the 410 experiments. Images were acquired every 10 min with Leica software. Spindle-rocking 411 experiments were acquired every 3 min with Leica microscope. HoloMonitor M4 is a 412 Ouantitative phase imaging-based cell analyzer utilizing the principle of digital 413 holographic microscopy. Live cell imaging was performed in HoloMonitor M4, enclosed in an incubator to maintain the samples at 37 °C and 5% of CO₂ throughout 414 415 the experiments. Images were acquired every 10 min with HStudio 2.7.

416

417 **PIV (Particle Image Velocimetry) measurement**

418 PIV analysis was conducted using a custom algorithm based on the MatPIV software 419 package for MATLAB. We used a series of live cell images of HeLa and MDCK to 420 calculate the velocity of the cells in the cell sheet. The mean velocity was subtracted 421 from calculated velocity fields to avoid any drift-related bias and to get the velocity 422 fields of the cells (the net movement of the cell cluster for its drift is less than 10% and 423 can be ignored). The heat maps of magnitude are also exported subtracting the mean 424 velocity. From the exported text files, we measured the overall cell speed, or root-mean-425 square (rms) velocity vrms and calculated the average vrms of each type of cell. The 426 correlation algorithm was coded using MATLAB in our lab. Experiments were 427 replicated at least three times.

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431 **Image processing, segmentation and quantification**

432 Cell data analysis of HoloMonitor M4: Cell area, cell thickness, cell sheet area and cell
433 sheet thickness were analyzed by the software HStudio 2.7. HStudio 2.7 can
434 automatically segment and extract physical parameters of the cell. For more details,
435 please refer to the official manual.

436 The area of cell clones and individual cells: Cell boundary labeling and areas were437 determined manually in Fiji.

Cell thickness, width and division angle: The collected confocal tomography images were imported into Bitplane imaris for 3D reconstruction. After reconstruction, XZ and YZ profiles were randomly selected to measure thickness, width and cell division angle manually. Data was measured on three samples, using three regions of the image. Thickness, width and division angle in chicken embryo skin: The collected confocal tomography images were imported into Bitplane imaris. Cell thickness, length, and division Angle were measured manually on software.

445 Pattern geometry in Chicken embryo skin: Diameter and spacing was measured446 manually using Fuji.

447 Shape index: To determine the cell boundary, we used a semi-automatic segmentation 448 pipeline. We manually enhanced the blurry boundaries of the cells. After processing 449 the image by binary Ostu, we produced the finally image by Median filtering. We 450 extracted the parameters by image segmentation to obtain the area and perimeter.

451 Cell trajectory: The cell tracking image is produced by software Bitplane imaris.

452 Cell junction analysis: Cell boundary labeling and areas were determined manually in

453 Fiji(Firmino et al., 2016).

454

455 **Strain field analysis**

he area of each cell was calculated by Fuji software, and the compression strain of each
cell was calculated according to Eq (3) in Supplementary Materials. Finally, origin
software is used to draw the compression strain diagram. The strain field is obtained by
analyzing variation of cell area using Eq. (4) in Supplementary Materials.

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461 **Finite element simulations**

Finite element simulations were performed using commercial software ABAQUS (2016). Details of the finite element method were present in the Supplementary Materials. In the finite element model, the growing cell monolayer sheet was placed on a stiff substrate with the interfacial fraction factor being controlled. The cell sheet was under isotropic expansion to simulate cell proliferation and growth. The cell sheet was modeled as the linear elastic material. No other boundary conditions were applied on the cell sheet.

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470 Skin culture and immunofluorescence

Fertilized eggs were bought from local farms. The eggs were cultured in a moist environment at 37 °C and staged according to Hamburger and Hamilton. Dorsal skin pieces were dissected from E6 embryos and spread flat on the Polycarbonate membrane (pore size: $0.4 \mu m$, Corning, Cat.No.3413). Culture is DMEM with 2% chick serum and 10% FBS along with pharmacological agents NSC23766 (Abmole, 20 μ M), RGD (Abcam, 100 μ g/ml) if applicable. Skin pieces were cultured for 48 hours at 37 °C before being fixed in 4% paraformaldehyde in PBS. Images were taken on the Nikon 478 stereoscopic microscope. Pattern geometry was calculated using Fiji to measure the 479 spacing and diameter. Skin culture details were performed as previously described. For 480 immunofluorescence staining, Embryos or cultured dissected skin pieces were fixed in 481 4% paraformaldehyde in PBS and embedded in OCT. The tissue blocks were placed in 482 liquid nitrogen for 1 minute and frozen section. The section thickness is about 14 μ m. 483 We incubate the sections with 5% BSA, in PBST (PBS + 0.1% Tween 20) for 2 hours. 484 The sections were incubated with primary antibodies at the optimal concentrations 485 (according to the manufacturer's instructions) at 4 °C overnight. After being washed 486 with PBS, the sections were incubated for 2 hours with secondary antibodies: 487 488/568/633 IgG (H+L) for 1 hour. Nucleus were stained with DAPI (4', 6-diamidino-488 2-phenylindole, Invitrogen) for 10 min at room temperature. Confocal images were taken on the Leica microscope equipped with a 10 x, 40 x or 63 x objective. 489 490 Experiments were replicated at least three times.

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492 Quantification and Statistical analysis

The number of biological replicates in each experimental result was indicated in the figure legends. Data were presented as means \pm SEM. Significance was determined using Student's t test to compare the differences between two experimental groups.

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- 621

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| 637 | performed and interpreted experiments. X.B.Z. and S.K.L. helped with cell experiments |
| 638 | and statistical analysis. J.L.C helped with PIV analysis. Y.Z. and Y.P.C. did mechanical |
| 639 | analysis. J.D., Y.P.C., and Y.B.F. conceived and supervised this project and prepared the |

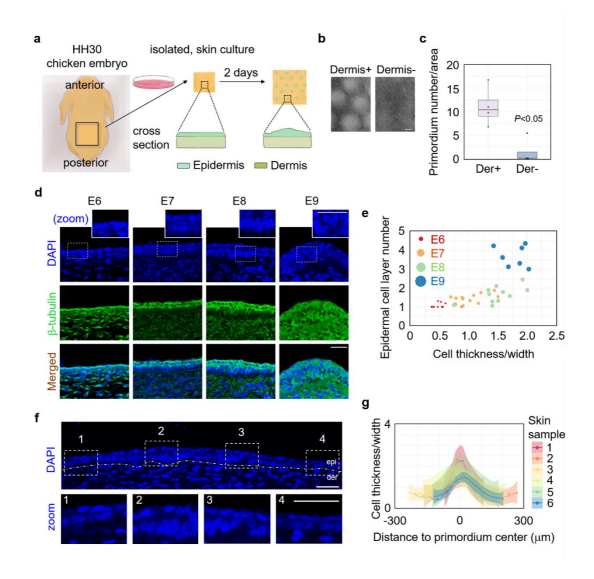
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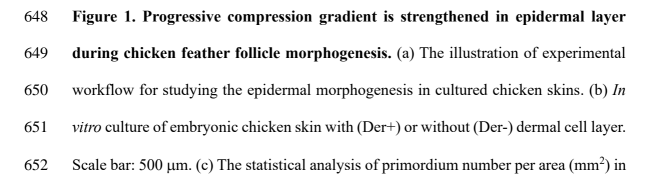
642 **Competing interests:** There are no competing interests.

644 Figures and Tables

645 **Figure 1**

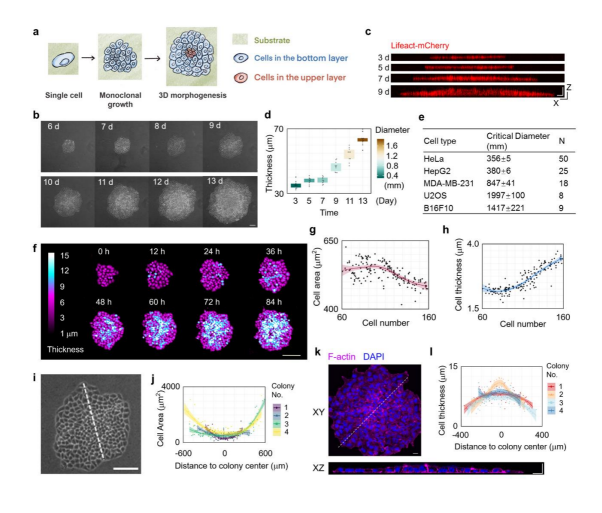


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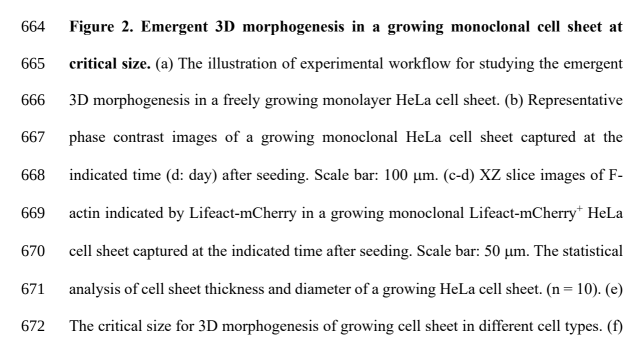


| 653 | (b). (d) Images of skin tissues from different stages of chicken embryos showing the |
|-----|-----------------------------------------------------------------------------------------------|
| 654 | deformation of epidermal cell shape. Scale bar: 25 μ m. (e) The statistical analysis of |
| 655 | cell deformation (thickness/width) and cell layer number of epidermis in skin tissues |
| 656 | from different stages of chicken embryos. Each dot represents the average value of an |
| 657 | embryo. (f) Images of epidermal cell shape deformation at different location around the |
| 658 | center of primordium. Scale bar: 25 μ m. (g) The statistical analysis of cell deformation |
| 659 | (thickness/width) with different distance to the center of primordium. |
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Figure 2

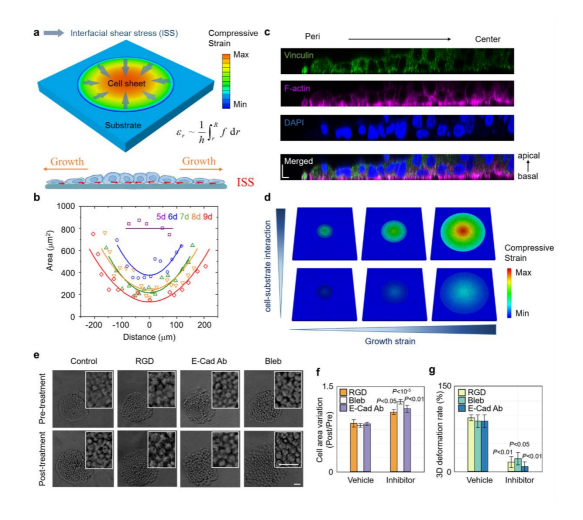


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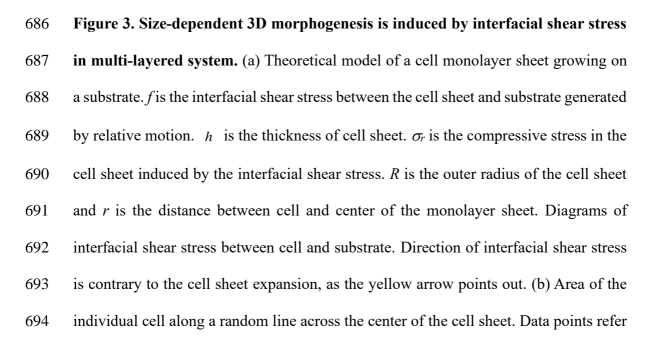


673 The representative live images of a growing HeLa cell sheet using HoloMonitor M4 674 time-lapse cytometer. Scale bar: 150 µm. (g) The statistical analysis of the area of individual cell during HeLa cell sheet growth. (h) The statistical analysis of the 675 676 thickness of individual cell during HeLa cell sheet growth. (i) The magnified view of 677 HeLa cell sheet at 8 d in (b). Scale bar: 100 µm. (j) The statistical analysis of the individual cell area along the lines in (i). (k) Representative XY and XZ slice images 678 of F-actin and nucleus stained by Phalloidin and DAPI respectively in HeLa cell sheet. 679 680 Scale bar: 20 µm. (1) The statistical analysis of the individual cell thickness along the 681 lines in (i).

683 Figure 3

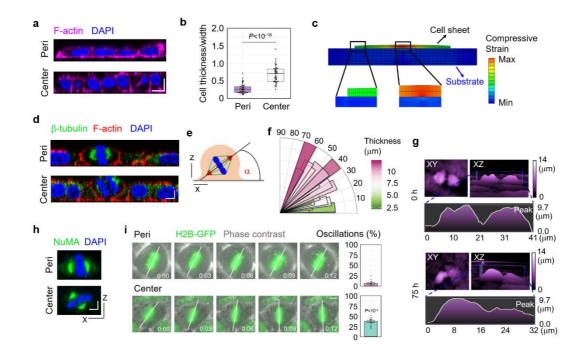


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695 to the experimental results and corresponding lines are theoretical predictions obtained 696 by fitting the experimental data using Eq. 6 in Supplementary Materials. (c) Representative XZ slice image of Vinculin, F-actin and nucleus stained by Vinculin 697 698 antibody, Phalloidin and DAPI respectively in half of a HeLa cell sheet. Scale bar: 10 699 μm. (d) Finite element simulation of the stress field in the growing cell monolayer with 700 different cell-substrate interactions ($f_{top} / f_{bottom} = 5$). Growth strains from left to right 701 are 10%, 60% and 120%, respectively. Color bar shows the distribution of the 702 compressive stress in the cell monolayer. (e) Representative phase contrast images of 703 HeLa cell sheet in the presence of inhibitors or vehicle. Scale bar: 50 µm. (f) The 704 statistical analysis of the individual cell area of HeLa cell sheet in the presence of 705 inhibitors or vehicle. (n = 10). (g) The morphogenesis percentage of HeLa cell sheet in 706 the presence of inhibitors or vehicle. (n = 3).

708 Figure 4



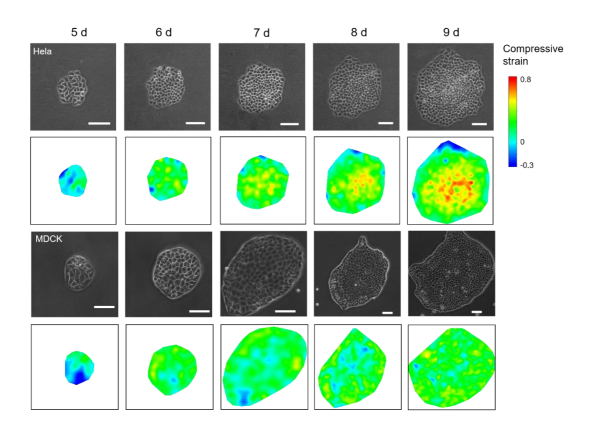
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711 Figure 4. Critical Compression triggers cell division reorientation to induce tissue 712 stratification. (a) Representative XZ slice image of F-actin and nucleus stained by 713 Phalloidin and DAPI respectively in the central region and peripheral region of HeLa 714 cell sheet. Scale bar: 10 µm. (b) The statistical analysis of cell deformation 715 (thickness/width) in the center region and peripheral region of Hela cell sheet. (n = 50). 716 (c) Cell shape variation induced by interfacial shear stress. (d) Representative XZ slice 717 image of β-tubulin, F-actin and nucleus stained by β-tubulin antibody, Phalloidin and 718 DAPI respectively in the central region and peripheral region of HeLa cell sheet. Scale 719 bar: 5 µm. (e) The schematic experimental setting of the mitotic spindle orientation. (f) 720 Distribution of the spindle-axis angles of cells with different thickness in HeLa cell sheet. (g) The representative images of dividing cells before (0 h) and after (75 h) 721 722 critical compression during HeLa cell sheet growth using HoloMonitor M4 time-lapse 723 cytometer. (h) Representative XZ slice image of NuMA and nucleus stained by NuMA

- antibody and DAPI respectively in the central region and peripheral region of HeLa
- 725 cell sheet. Scale bar: 5 μm. (i) Analysis of spindle oscillation in in the central region
- and peripheral region of GFP-H2B⁺ U2OS cell sheet. The extent of oscillation was
- calculated and plotted in bar graphs on the right. Scale bar: 5 μ m. (n = 14).

729 **Figure 5**

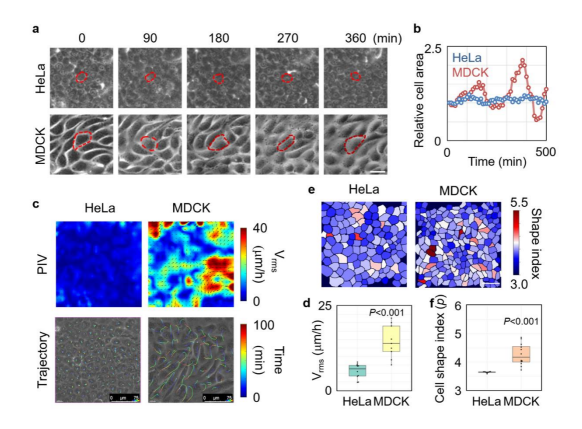
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Figure 5. Strain field evolution of HeLa and MDCK cell sheets. The calculated strain
field evolution during the growth of Hela and MDCK cell sheets at the indicated time
(d: day) after seeding. Compressive strain field was calculated using Eq. (4) in the
Supplementary materials by measuring the areas of each cell. The normal area of a fullgrown cell without sustaining compression was set as the average area of the cell sheet
in 1 d since the compressive gradient is small in the initial stage. Scale bar: 100 µm.

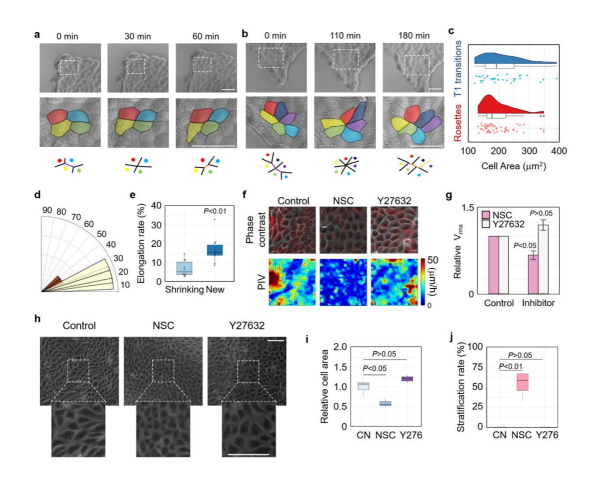
739 **Figure 6**

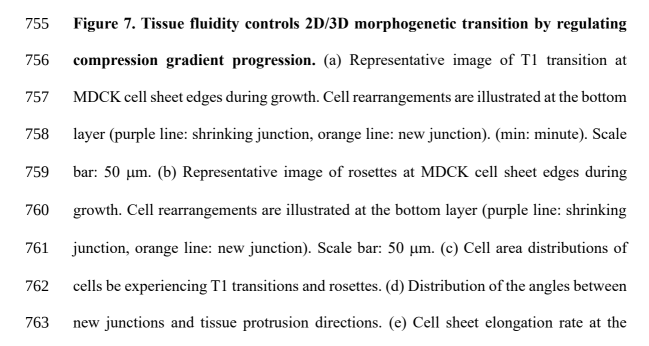


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742 Figure 6. HeLa and MDCK cell sheets show different fluidity during growth. (a) The fluctuation of individual cell shape during the growth of HeLa and MDCK cell 743 744 sheets. Scale bar: 25 µm. (b) The area alteration with time of the cell indicated by dotted 745 line in (a). (c) Cell velocity field analyzed by PIV and cell trajectories in HeLa and 746 MDCK cell sheets. Scale bar: 75 µm. (d) The statistical analysis of cell speed (rms velocity) measured by PIV in HeLa and MDCK cell sheets. (n = 10). (e) The cell shape 747 748 index distribution of HeLa and MDCK cell sheets. Scale bar: 50 µm. (f) The statistical 749 analysis of cell shape index of HeLa and MDCK cell sheets. (n = 10). 750

Figure 7

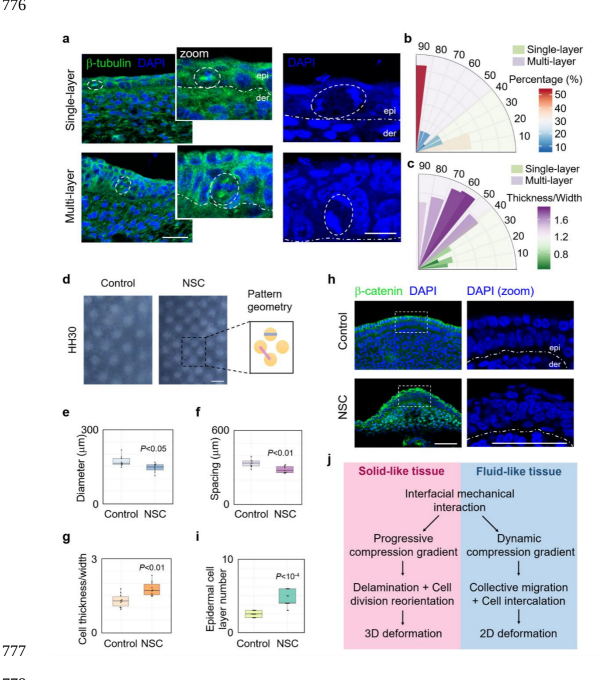




| 764 | direction of shrinking junctions or new junctions. ($n = 10$). (f) The velocity field |
|-----|--------------------------------------------------------------------------------------------------|
| 765 | superimposed on the corresponding phase contrast (upper panel) and velocity map |
| 766 | (lower panel) images measured by PIV of MDCK cell sheet in the presence of inhibitors |
| 767 | or vehicle. Scale bar: 20 μ m. (g) The statistical analysis of cell speed (rms velocity) |
| 768 | measured by PIV of HeLa and MDCK cell sheets in the presence of inhibitors or vehicle. |
| 769 | (n = 4). (h) Representative phase contrast images of MDCK cell sheet in the presence |
| 770 | of inhibitors or vehicle. Scale bar: 100 μ m. (i) The statistical analysis of the individual |
| 771 | cell area of MDCK cell sheet in the presence of inhibitors or vehicle. $(n = 3)$. (j) The |
| 772 | stratification percentage of MDCK cell sheet in the presence of inhibitors or vehicle. (n |
| 773 | = 4). |
| | |

775 Figure 8

776



778

779 Figure 8. Progression of compression gradient contributes to epidermal cell 780 stratification during chicken skin development. (a) Images of embryonic chicken 781 skin showing the orientation of mitoses relative to the basement membrane (white dotted line) in the magnified images (zoom), separating epidermis (epi) from dermis 782 783 (der). White dotted circles indicate the mitotic cells in metaphase (left) and anaphase

| 784 | (right). DAPI marks the DNA and β -tubulin antibody immunofluorescence staining |
|-----|------------------------------------------------------------------------------------------------|
| 785 | marks the spindle. Scale bar: 10 μ m. (b) Distribution of the spindle-axis angles of cells |
| 786 | in single-layered or multi-layered epithelia of embryonic chicken skin. (c) Distribution |
| 787 | of the spindle-axis angles of cells with different cell deformation (thickness/width) in |
| 788 | single-layered or multi-layered epithelia of embryonic chicken skin. (d) Reconstitution |
| 789 | culture of embryonic chicken skin with or without NSC23766 (NSC). Scale bar: 500 |
| 790 | μ m. Quantification of (e) spacing and (f) diameter as pattern geometry parameters as |
| 791 | illustrated in the right panel of (d). $(n = 10)$. (g) Quantification of cell deformation of |
| 792 | epidermis in in the presence of NSC or vehicle. ($n = 10$). (h) Representative cross |
| 793 | section image of β -catenin and nucleus stained by β -catenin antibody and DAPI |
| 794 | respectively in reconstitution cultured embryonic chicken skin with or without |
| 795 | NSC23766 (NSC). Scale bar: 50 μ m. (i) Quantification of layer number of epidermis |
| 796 | in (h). $(n = 14)$. (j) Model of 2D and 3D morphogenesis controlled by the interfacial |
| 797 | mechanical interactions in multi-layered systems with different tissue fluidity. |
| | |

798

800 Supplementary Materials

801 **Theory model**

802

The theoretical model is shown in Fig. 3a. The cell monolayer is modeled as a thin 803 film, which is attached on the surface of a rigid substrate. Direction of the tangential 804 805 adhesion force between the cell and substrate is contrary to the relative motion. Thus, 806 compression would be induced when the cell monolayer is growing on the substrate. 807 The tangential adhesion force is assumed as f = f(r), where r is the distance 808 between the cell and the center of the monolayer. f is concerned with the tangential adhesion between the cell and substrate, which is related to the cell type, stiffness of 809 810 substrate and the rate of cell division or growth. By analyzing the stress state of the 811 monolayer and solving the equilibrium equation, one can obtain the distribution of the 812 equi-biaxial compressive stress in the monolayer

813
$$\sigma_r = \frac{1}{h} \int_r^R f \, \mathrm{d}r \tag{1}$$

814 Here *R* is the outer radius of the cell monolayer, and h is the thickness of cell sheet. 815 The compressive strain can be obtained by the constitutive relation as

$$\mathcal{E}_r = \frac{1 - \nu}{Eh} \sigma_r \tag{2}$$

817 where E, ν are the modulus and Poisson's ratio, respectively. The central region of 818 the monolayer would sustain higher level of compression than the cells in other regions. 819 Thus, cell extrusion is most likely to occur in the central region, which is consistent 820 with the experimental observations. The maximum compressive strain in the cells can 821 be obtained as

822
$$\varepsilon_{\max} = \frac{1 - \nu}{Eh} \int_0^R f \, \mathrm{d}r \tag{3}$$

which depends on the cell-substrate interactions and area of the cell monolayer. Basedon Eq. (2), one can also obtain the cell area

825

$$A/A_0 = (1 - \varepsilon)^2 \tag{4}$$

where A_0 is the normal area of a full-grown cell without sustaining compression. Eq. (4) demonstrates that distribution of cell area can reflect the strain state in the cell monolayer. With cell proliferation, radius of the cell monolayer increases, and area of cells in the central region is reduced, indicating that the cell monolayer has a high level of compression in the central region.

Given the distribution of the interfacial adhesion force, one can obtain the distribution of the compressive strain and cell area of the cell monolayer. In this work, the tangential adhesion force is assumed to be uniform over the cell monolayer. Thus, the maximum compressive strain in the cell and the distribution of the cell area can be given by

836
$$\varepsilon_{\max} = \frac{1 - \nu}{Eh} fR \tag{5}$$

837
$$A/A_0 = \left[1 - \frac{1 - \nu}{Eh} f(r - R)\right]^2$$
(6)

Based on Eq. (4), distribution of the compressive strains in the HeLa cells in experiments can be obtained by analyzing the distribution of cell area, which is shown in Fig. 2b. The sequence in Fig. 2b shows the morphologies of growing HeLa cells with the distribution of the compressive strains given in Fig. 5. The size of the cell monolayer sheet grows bigger along with the cell proliferation, generating higher level of compressive stress in the monolayer. Areas of individual cells in the central region are reduced due to the increased compressive strains. The experimental observations and

845 calculations are consistent with the theoretical predictions.

846 To validate the theoretical model, finite element simulations are performed to explore the relation between the cell-substrate interactions and stress field in growing cells. 847 848 Results of finite element simulations are shown in Fig. 3d. In the finite element model, 849 the growing cell monolayer sheet was placed on a stiff substrate with the interfacial 850 fraction factor being controlled. The cell sheet was under isotropic expansion to 851 simulate the growth. More than 12,000 linear hexahedral elements were adopted to in 852 the simulations. The cell sheet was modeled as the linear elastic material. With the 853 increase of the growth strain, compressive stresses in the cell monolayer are generated 854 due to the cell-substrate interactions, and the central region has a higher level of 855 compressive stress. The stress level can be reduced by regulating the cell-substrate 856 interactions (Fig. 3d). Finite element simulations are consistent with experimental 857 observation and theoretical analysis.

858 The elastic strain energy stored in the cell would also increase with the cell 859 proliferation. When the elastic strain energy in the cell is small, interfacial normal 860 adhesion would impose restriction on cell extrusion, making the cell monolayer grow 861 in plane. Thus, higher compressive stress would be generated further. When the compressive stress reaches a critical value, elastic strain energy stored in the cell may 862 863 be greater than the energy needed for the occurrence of the interfacial delamination. In 864 this critical condition, cell extrusion may happen, and the elastic strain energy can be 865 released. The critical condition for the cell extrusion can be written as

866
$$\gamma A = \frac{E}{1 - \nu} \varepsilon_{\max}^2 V \tag{7}$$

The left hand and right hand of the equation refer to the energy for the interfacial delamination and elastic strain energy in the cell, respectively. γ is the energy per area for the interfacial delamination, which is related to the interfacial normal adhesion

between the cell and substrate. ε_{max} is the maximum strain in the cell, and V is the cell volume. Eq. (6) demonstrates that there exists a critical area or size of the cell monolayer at the critical condition of cell extrusion. The critical size of the cell monolayer sheet depends on the mechanical properties of cell and cell-substrate interactions. The cell extrusion can be controlled by regulating the modulus of the cell and cell-substrate interactions.

876 When the normal adhesion between the cell monolayer and the substrate is enhanced, 877 cell extrusion would be more difficult to happen, and the cell would bear larger compression before extrusion. When the tangential adhesion is larger, higher level of 878 879 compression would be induced, and the cell extrusion is more likely to happen. The 880 adhesion between the cells may have effect on the cell sheet stress. Maximum 881 compressive stress in the cell monolayer may be reduced when lowering down the adhesion between the cells. Thus, cell extrusion would be more difficult to happen. 882 883 Cytoskeleton is the main components that determine the cell mechanics. The 884 cytoskeleton can bear compressive stress and store elastic strain energy. If the 885 cytoskeleton in the cell is suppressed, cell extrusion is hard to happen. These predictions 886 were confirmed by experiments (Fig. 3e-g).

Shape of the cell can be changed due to the compression induced by the interfacial shear stress. Thus, the orientation of the cell division may be altered. During the evolution of the cell monolayer sheet, cell shape in the central region was changed from flat to columnar due to the high level of compression stress. While in the periphery of the monolayer, cell shape maintained flat. Stretch ratio of the cell in the direction of thickness can be written as

$$\lambda = 1 + \frac{2\nu}{1 - \nu} \varepsilon_r \tag{8}$$

Base on the stretch ratio in Eq. (8), cell thickness/width characterizing cell shape canbe obtained as

896
$$\frac{h}{b} = \frac{\lambda}{1 - \varepsilon_r} \frac{h_0}{b_0}$$
(9)

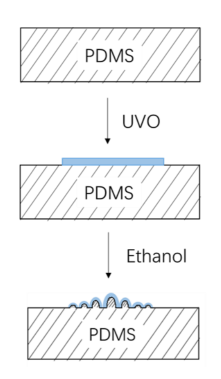
Here *h* and *b* refer to the cell thickness and width. h_0 and b_0 refer to the initial thickness and width of the cell without sustaining compression. In the periphery of the monolayer, cell thickness/width is small since the compressive strain in the cell is small. While in the central region with high level of compressive strain, cell thickness/width can be very large. This phenomenon of cell shape variation induced by the interfacial mechanical interaction is also confirmed by the finite element simulations, as shown in Fig. 4c.

905 Supplementary Figures

906

907 Supplementary Figure S1

908



909

- 914 in blue) on the surface. Then dropping an ethanol/glycerol mixture solution containing
- 915 60% 100% ethanol by volume to induce surface wrinkling.

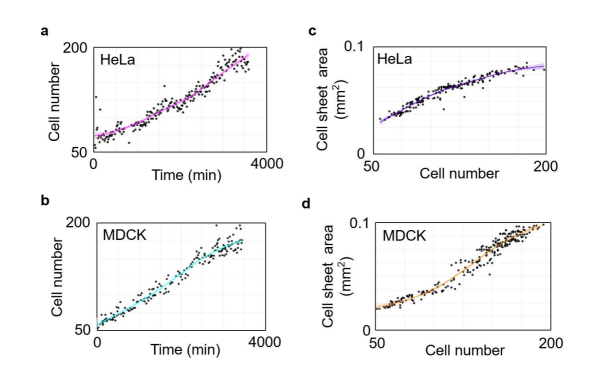
⁹¹¹ Figure S1. Surface wrinkling induced by differential expansion in a film-substrate

⁹¹² system (Zhao et al., 2015b). Polydimethylsiloxane (PDMS) was exposed to UV/Ozone

^{913 (}UVO) for 10-55 minutes to form a stiff solvent-responsive oxide layer (highlighted

916 Supplementary Figure S2

917

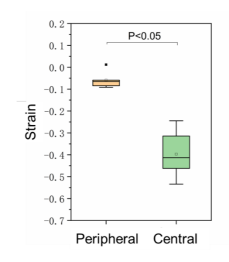


918

Figure S2. Proliferation and expansion of cell sheets during growth. The statistical
analysis of the cell number during HeLa cell sheet (a) and MDCK cell sheet (b) growth.
The statistical analysis of the cell sheet area during HeLa cell sheet (c) and MDCK cell
sheet (d) growth.

924 Supplementary Figure S3





926

927

928Figure S3. Single cell tracing of compressive strain. The plane strain of single cells929in central and peripheral regions of cell sheets during 24 hours analyzed by holographic930imaging cytometer (n = 6).

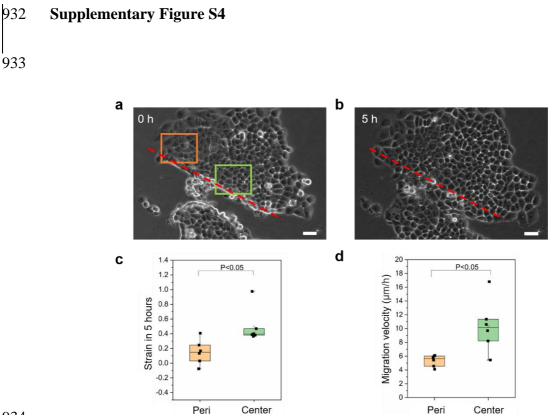
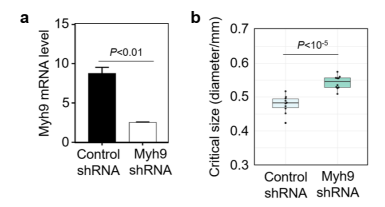


Figure S4. Cell sheet scratch assay. The representative image of cell sheet upon (a) and 5 hours after (b) scratch passing through the center (green box) and edge (orange box) of the cell sheet. The expansion (c) and migration speed (d) were compared between cells in the central region and peripheral region of the cell sheet. Scale bar: 50 μ m, (n = 6).

940 Supplementary Figure S5

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942

943 Figure S5. Inhibition of Myosin effects the emergence of 3D morphogenesis. (a)

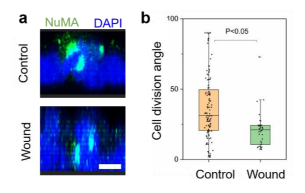
The mRNA level of myosin heavy chain 9 gene (myh9) in HeLa cells transfected with

945 myh9 shRNA and control shRNA. (n = 3). (b) The critical sizes for morphogenesis in

946 myh9 shRNA cells and control cells. (n = 10).

948 Supplementary Figure S6

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951

952 **Figure S6. Cell division angle after scratch.** (a) Representative XZ slice image of

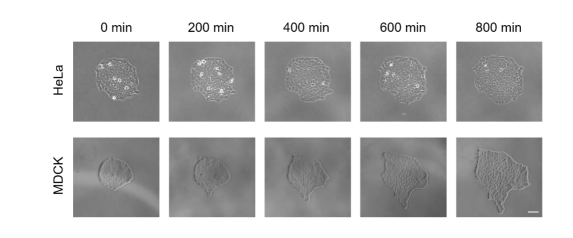
953 NuMA and nucleus stained by NuMA antibody and DAPI respectively in the control

and wound of HeLa cell sheet. (b) The cell division angle in the center of cell sheet

955 without (Control) or with (Wound) scratch. Scale bar: $10 \ \mu m$.

957 Supplementary Figure S7

958



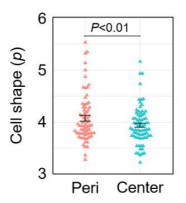
959 960

961 Figure S7. Different deformation pattern between HeLa and MDCK cell sheets

962 **during growth.** Representative phase contrast images of a growing monoclonal HeLa

and MDCK cell sheet captured at the indicated time (min: minute). Scale bar: 100 µm.

964 Supplementary Figure S8



965

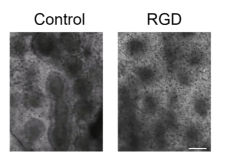
966

967 Figure S8. Cell shape index in the peripheral and central regions of HeLa cell sheet.

968 The statistical analysis of cell shape index of cells in the peripheral region and central

- 969 region of HeLa cell sheet. (n = 75).
- 970

971 Supplementary Figure S9



972 973

- 974 Figure S9. Disrupting the interfacial mechanical interaction between epidermal
- 975 and dermal layers attenuated the morphogenesis of primordium. *Ex vivo* culture of
- 976 embryonic chicken skin with or without RGD. Scale bar: 500 μm.

978 Supplementary Table

979

| ECM type | Critical diameter(µm) | Ν |
|--------------------------|-----------------------|----|
| No pre-coating ECM | 463.8±18.9 | 10 |
| Fibronectin | 493.8±23.9 | 10 |
| Gelatin | 489.1±19.8 | 10 |
| Collagen | 520.3±21.8 | 10 |
| Collogen (40 kPa PA gel) | 413.8±9.6 | 5 |
| Collogen (1 kPa PA gel) | 342.6±26.0 | 4 |

980 981

982 **Table S1.** The critical size for 3D morphogenesis of growing cell sheet on different

983 types of ECM and stiffness (polyacrylamide gel).

| 985 | |
|------------|-----------------------------------------------------------------------------------|
| 986 987 | Supplementary Movies |
| 988 989 | Supplementary Movie 1. Symmetric cell division before critical compression during |
| 990 | HeLa cell sheet growth visualized by HoloMonitor M4 time-lapse cytometer. |
| 991 | |
| 992 | |
| 993 | Supplementary Movie 2. Asymmetric cell division after critical compression during |
| 994 | HeLa cell sheet growth visualized by HoloMonitor M4 time-lapse cytometer. |
| 995 | |
| 996 | |
| 997 | Supplementary Movie 3. PIV analysis during HeLa and MDCK cell sheets growth. |
| 998 | |
| 999 | |
| 1000 | Supplementary Movie 4. Time-lapse imaging of MDCK cell sheet during growth. |
| 1001 | |
| 1002 | |
| 1003 | Supplementary Movie 5. PIV analysis during MDCK cell sheet growth in the presence |
| 1004 | of inhibitors or vehicle (control). |
| 1005 | |
| 1006 | |