Gastrulation in amphibian embryos, regarded as a succession of biomechanical feedback events

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ABSTRACT  Gastrulation in amphibian embryos is a composition of several differently located morphogenetic movements which are perfectly coordinated with each other both in space and time. We hypothesize that this coordination is mediated by biomechanical interactions between different parts of a gastrulating embryo based upon the tendency of each part to hyper-restore the value of its mechanical stress (see Belousov and Grabovsky, 2006). The entire process of gastrulation in amphibian embryos is considered as a chain of these mutually coupled reactions, which are largely dependent upon the geometry of a given embryo part. We divide gastrulation into several partly overlapped steps, give a theoretical interpretation for each of them, formulate the experiments for testing our interpretation and describe the experimental results which confirm our point of view. Among the predicted experimental results are: inhibition of radial cell intercalation by relaxation of tensile stresses at the blastula stage; inversion of convergent intercalation movements by relaxation of circumferential stresses at the early gastrula stage; stress-promoted reorientation of axial rudiments, and others. We also show that gastrulation is going on under a more or less constant average value of tensile stresses which may play a role as rate-limiting factors. A macromorphological biomechanical approach developed in this paper is regarded as complementary to exploring the molecular machinery of gastrulation.

KEY WORDS: gastrulation, morphogenesis, mechanical stresses, Xenopus laevis

It is not birth, marriage or death, but gastrulation which is truly the most important time in your life.
L. Wolpert (1986)

Introduction

Gastrulation in general and, particularly in amphibian embryos (taking the Anurans and, in particular, Xenopus laevis as examples) is a space/temporal composition of several morphogenetic events. The main ones are: (1) radial cell intercalation (RCI) in the blastocoel roof; (2) formation of bottle-shaped cells in the marginal zone, starting from the dorsal side of an embryo; (3) convergent cell intercalation (CCI) in the suprablastoporal zone (SBZ) directed towards dorso-medial embryo midline (Keller, 1987; Keller and Daniilchik, 1988; Wilson et al., 1989); (4) involution around the blastoporal lip (Holtfreter, 1944; Keller, 1981, 1984; Keller and Hardin, 1987). Although it has long been known (Spemann, 1936; Holtfreter, 1944; Keller and Jansa, 1992) that all of these processes can proceed, in experimental conditions, independently from each other, their rates, locations and directionality are, after such an isolation, largely distorted. Meanwhile, in an entire embryo they are perfectly coordinated and enhance each other. Moreover, gastrulation-like movements can be easily reproduced in small pieces of embryonic tissues (Holtfreter, 1944) which shows that they are not unambiguously related to any definite regional properties of embryo material. That makes it possible that the gastrulation movements are the components of a structurally stable chain of self-organizing events linked with each other by some kinds of feedback. In this study we use the theoretical constructions described in the accompanying paper (Belousov and Grabovsky, this volume) and suggest that the feedback may be based upon the tendency of any part of an embryo which is mechanically affected (stretched, relaxed or compressed) by other part(s) to hyper-restore its initial stress value, reciprocally affecting other part(s). We divide the entire

Abbreviations used in this paper: CCI, convergent cell intercalation; CE, contraction-extension (feedback); EE, extension-extension (feedback); HR, hyper-restoration; MZ, marginal zone; RCI, radial cell intercalation; RD, residual deformation; SBZ, suprablastoporal zone.

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Fig. 1. Biomechanical interpretation of gastrulation events in amphibian embryos. (A-D, E1, E2) Biomechanical "maps" of successive stages and most important regions of embryos. Active forces are displayed by red arrows, while passive stresses are indicated by blue arrows. (A) Stretching of the blastocoel roof by turgor pressure in the blastocoel. (B) RCI in the blastocoel roof producing pressure forces onto vegetal regions. (C) A detailed scheme of the region framed in (B). (D) Two alternating phases of biomechanical interactions between the marginal zone (MZ) and the suprablastoporal zone (SBZ). (E1, E2) Successive stages of the involution in sagittal section. Frames (ab, c, d, e) display suggested hyper-restoration responses that correspond to the upper row frames indicated by the same capital letters (ab corresponds to A and B together). Dotted arrows connect the active branches of preceding hyper-restoration responses with the shifts of stress values which they produce in the neighboring parts of embryos triggering the latter’s active responses. Pre, preinvoluted; pst, postinvoluted layer.

gastrulation process, starting from the blastula stage, into four successive (but partly overlapping) steps regarding each of them as a single HR feedback loop creating the initial conditions for the next one. Two first steps are just preparatory, while latter two comprise the gastrulation proper. For each step, we firstly suggest a theoretical construction, then formulate some definite experimentally verified predictions and, at last, submit the supporting experimental evidence. Theoretical constructions are summarized in Fig. 1. For more detailed explanation of the terminology and schemes associated with HR hypothesis we refer the readers to the accompanying paper. Stage numbers are given by Nieuwkoop and Faber (1956).

**Step 1: Radial cell intercalation in the blastocoel roof**

**Theoretical interpretation**

RCI is a hyper-restoration reaction to the stretching of the blastocoel roof by an increased turgor pressure within a blastocoel (Fig. 1 A,B: ab). As such, it obeys the contraction-extension (CE) feedback (see accompanying paper, Fig. 4).

**Predictions**

1. A typical shape of amphiblastula with a thin roof and thick bottom is structurally stable and can thus be developed from a spherically symmetric state without any special initial conditions. Suggest that the blastocoel is ideally centered and surrounded by walls of equal thickness. From our point of view, such a shape, if being pressurized from inside, will be unstable. If indeed one part of the blastocoel wall (A) becomes a bit thinner than another one (B) the pressure-generated tensile stress applied to A will be greater than that applied to B and the extension-promoted RCI will function predominantly in A, leading to its further thinning. At the same time, according to CE feedback, RCI within the A wall will stimulate the tangential contraction and cell immigration in B, making it more thick.

2. Relaxation of the blastocoel roof should inhibit RCI.

**Experimental results**

1. If one extirpates a small tissue piece from a blastocoel roof of an early—mid-gastrula (stages 10-11) embryo (Fig. 2A), within several dozens minutes it will roll into a solid ball covered by an outer ectodermal layer. Soon a cavity forms inside, whose spherical shape indicates that it is under turgor pressure. The cavity gradually shifts out of the central position, one of its walls becoming very thin while the opposite one becomes thick and indented, indicating cell immigration (Fig. 2 B-D; see for more details Beloussov and Petrov, 1983). This shows that RCI-mediated eccentricity of a blastula cavity is a structurally stable property, not associated with some unique initial conditions.

2. If one relaxes circumferential tensions at the blastula stage (9-10) by inserting a wedge of endodermal tissue into a vegetal hemisphere of an embryo (Fig. 3A), the process opposite to RCI is taking place, namely the clustering of deep cells of the blastocoel roof into a condensed ensemble (Fig. 3 B,G). Additionally, several hours after the operation the animal surface of relaxed embryos becomes ruffled and acquires several indentations, later on taking the shape of large folds and protrusions (Fig. 3C). Obviously, all these events promote a circumferential contraction of the blastula surface and hence an increase of tensions on its surface. For example, the formation of just a single protrusion (like that shown in Fig. 3C) stretches the surrounding surface area for no less than about 15 % and usually more than one protrusion is created. As a rule, such embryos do not exhibit gastrulation-associated involution movements. The axial organs, if formed, are arranged quite abnormally (Ermakov and Beloussov, 1998; Beloussov and Ermakov, 2001). Meanwhile, if we restore the

![Image](Image)
tensile stresses soon after their relaxation (either by putting embryos in hypotonic solution which increases the water flow into the blastocoel or by direct embryos stretching with the use of two needles) the RCI is renewed and a subsequent development normalized (see Fig. 3 D,G, and E,H,F). The corresponding quantitative data are summarized in Fig. 3I (for more details see Belousov and Ermakov, 2001).

Step 2: Formation of bottle-shaped cells in the marginal zone

Theoretical interpretation
This is the HR reaction to this zone relaxation/compression by the actively extended blastocoel roof (Fig. 1 B,C, ab, c).

Predictions
(1) Bottle-shaped cells should actively contract themselves in the direction of their relaxation/compression by an actively extended blastocoel roof (that is, in the dorso-ventral direction).
(2) Artificial relaxation of embryonic tissue can induce the formation of bottle-shaped cells in unusual locations.

Experimental results
1. The existence of tangential pressure exerted by the blastocoel roof towards more vegetal embryo regions is confirmed by planimetric measurements performed at the late blastula stage (Keller, 1978): the square of the blastocoel roof is significantly increased within this period, while that of the MZ remains constant. The expected dorso-ventral (meridian) contraction and hence transversal (latitudinal) flattening of bottle-shaped cells has been directly observed (Hardin and Keller, 1989). The authors found such a highly anisotropic contraction not to be an intrinsic property of bottle-shaped cells, because in isolated explants they contract isotropically.

2. The excessive bundles of bottle-shaped cells are produced in the following abnormal locations as a consequence of tissue relaxation:
(a) onto the SBZ surface relaxed at stages 10-11 by P-shaped incision and separation from underlain tissue (Fig. 4A; see for details Belousov, 1988);
(b) on the bottom of a newly arisen pit formed onto the tip of a relaxed at stages 10 1/2 - 11 dorsal blastoporal lip (Fig. 4 B-D). The lip was firstly isolated and then relaxed by a longitudinal incision (Fig. 4B). Its reaction to the relaxation consisted of two successive steps. The first one (Fig. 4C) is passive. It takes no more than a few seconds and can proceed under low temperatures (4 – 60°C for Rana temporaria and 10 – 120°C for Xenopus laevis embryos). The second one (Fig. 4D) is active (occurring only under normal temperature) and takes several dozen minutes. As a result, the lip’s curvature is inverted (cf. Fig. 4 B,D). This is a clear overshoot reaction, because for restoring tensions without an overshoot it would be enough for a dissected lip to return to the curvature of the blastula surface.

(c) on the ventral embryo surface after relaxation of circumferential tensions at the early gastrula stage (10 – 10 1/2) by a dorso-medial cut (see below for more details).

Step 3: Gastrulation proper; epiboly and involution

Theoretical interpretation
We suggest that the main driving force of gastrulation is the contraction-extension (CE) feedback established between MZ and the surrounding embryonic surface, first of all between the dorsal blastoporal lip and SBZ (Fig. 1 C,D,d). This feedback is
initiated by the above mentioned meridian contraction of the bottle-shaped cells which triggers the active (convergent) extension of SBZ. This latter in its turn relaxes/compresses the MZ, promoting its contraction and so on. As a result, MZ corresponds to the $\alpha$ zone, while SBZ corresponds to the $\beta$ zone (see Fig. 4 from Beloussov and Grabovsky paper, this volume). As gastrulation proceeds, CE feedback extends to more vegetal MZ regions. The involution of cell material around the blastoporal lip is a clear indication of an overshoot SBZ response because the linear rate of the active SBZ extension exceeds that of the MZ contraction (such an excess has been directly demonstrated by time-lapse filming: Glagoleva et al., 2003). At the same time, the blastoporal lip, while becoming a part of a toroidal surface, starts to work as a funnel (see Fig. 3D from Beloussov and Grabovsky paper, this volume), converging the adjacent cells towards the dorsal embryo midline.

Predictions

1. The location of the dominating meridian of cell convergence as well as the direction of convergence should depend upon the tensile pattern within the MZ, rather than being an autonomous property of converging cells.

As argued by Beloussov and Grabovsky (this volume), the MZ is a semi-toroidal surface pressurized from inside, with its merid-
transversal axis, that is, perpendicularly to its normal direction. At the same time, such a stretching should relax $\sigma_c$ in the latero-ventral MZ regions and hence promote ventralwards cell convergence as in the previously suggested experiment (Fig. 5D).

2. Cell convergence towards meridians should take place in any internally pressurized fold (even if prepared artificially), or in a tissue “scroll”, irrespective of its specificity.

**Experimental results**

1(a). In *X. laevis* embryos destined for the operations, at 32 blastomeres stage two ventral (Fig. 6 A-C) or two dorsal (Fig. 6D) blastomeres were labeled by microinjecting fluorescein-dextran. At the early gastrula stage ($10 - 10^{1/2}$) a dorso-medial cut was made through the SBZ and the dorsal blastoporal lip. The wound gap was covered by a piece of ventral ectoderm in order to weaken wound healing reactions at the gap edges. In the embryos with ventral tissue labeled, within a few hours after operation, condensation of the labeled material along the ventral embryo midline was observed. Later on an abnormal tail formed on the ventral (rather than dorsal, as normally) lip of the blastopore (Fig. 6B). This was in a sharp contrast with the same cells movements in unoperated embryos, which are always directed dorsally (Fig. 6A). Correspondingly, due to the same operation the labeled dorsal material also moved ventralwards and formed stripes aligned along the lateral lips of the blastopore (Fig. 6D), which is strictly opposite to the normal dorsalwards convergent movements. Hence, according to our expectations, a complete inversion of cell convergence movements took place in these experiments.

1(b). Before making a similar dorso-medial cut, we decreased $\sigma_m$ by cutting off an entire MZ from the rest of embryo. In these, “sausage-like” MZ preparations no ventralwards convergence took place at all (Fig. 6D). Instead, a patch of labeled cells has extended towards the edges of the explant.

1(c). We first give an overall view of the movements of the labeled cells and then show the morphological results of transversal SBZ stretching.

As previously, two dorso-medial blastomeres were labeled at

![Fig. 6. Inversion of cell convergence movements as a result of a dorso-medial cut of early gastrula stage *X. laevis* embryos.](image)
the 32 cells stage. When the embryos reached early gastrula stage, the SBZ was stretched by use of 4 needles (Fig. 7 A,B). By 2.5 h after fixing the needles in the stretched positions a ventralward movement of the labeled cell material could be traced. It was so extensive that the tip of needle #1 was involved, in spite of its lower part being anchored in the agarose substrate (Fig. 7 B,C). Later on this movement was continued, leading in 7 h to shift of the labeled cells towards the lateral lips of the blastopore, that is, greatly overlapping the needles’ positions. This result is quite similar to that obtained by a dorso-medial cut (see Fig. 6).

If the SBZ undergoes two rounds of stretching (being stretched in the total for no less than about 120% of its initial length) and maintained stretched for about 20 h, a reorientation in the stretch direction of entire embryo body, including not only the axial organs (notochord, neural tube and somite series), but also the yolk-containing compartments. In (D) and to some extent in (F) part of the notochord is shifted towards the lateral blastoporal lip (G). A scheme of a similar operation performed at the early neurula stage. (H) Its result, total view. Contrary to (A-F), the larva fully preserves its initial axis, remaining bound to one of the needles by a thin unstretched tissue thread (later on disrupted). In (H), pointers indicate the positions of the needles.

Fig. 7 (Left). Induction of the active lateralwards movements of the labeled descendants of dorsal cells as a result of transversal suprablastoporal zone stretching by four needles. To the left: optical light and luminescent views just after inserting needles and before stretching them (A), immediately after stretch (B) at 2.5 h (C), 3.5 h (D) and 7 h (E). In optical light views, needles are denoted by black spots. To the right are shown the positions of the needles 1-4 before stretching, immediately after stretch and 2.5 h later. Note a considerable post-stretching increase of the distance 1-2 in spite of the needles being anchored to the agarose substrate. As a result, the dorsal cell material is very much shifted to the ventral, considerably overlapping the needles’ positions. This result is quite similar to that obtained by a dorso-medial cut (see Fig. 6).

Fig. 8 (Right). Morphological results of suprablastoporal zone (SBZ) transversal stretching, as compared with a similar stretch of neurula stage embryos. (A) Scheme of SBZ stretching by two needles. (B-F) 24 h results. (B,C) Total views (on frame B, the direction of stretching is indicated), whereas (D-F) are histological sections made in transversal embryo planes. Note a complete transversal reorientation of the embryo body including not only the axial rudiments (notochord and somite series), but also the yolk-containing compartments. In (D) and to some extent in (F) part of the notochord is shifted to the lateral blastoporal lip. (G) A scheme of a similar operation performed at the early neurula stage. (H) Its result, total view. Contrary to (A-F), the larva fully preserves its initial axis, remaining bound to one of the needles by a thin unstretched tissue thread (later on disrupted). In (H), pointers indicate the positions of the needles.
only result of such manipulation was the extrusion of a thin tissue thread (later on disrupted) towards one of the needles (Fig. 8H). Therefore, the capacity for a profound stretch-dependent reorientation of cell movements was a property of gastrulation stage embryos only. Later on the stage-specificity of biomechanical cell reactions will be discussed in more detail.

2. Cell convergence in the artificially prepared folds and tissue “scrolls”.

If it is true that the very geometry of a fold provides cell convergence towards its midline, this effect should take place not only on the “natural”, but also on the artificially prepared folds. To check this hypothesis, we excised from a midgastrula embryo (stage 11 - 11 1/2) a piece of ectoderm situated laterally to the dorsal midline, folded it by its outer surface outside and fixed the fold with a glass needle (Fig. 9A). At about 20 h after operation we found, in 4 cases out of 6, rather long protrusions that had grown from a midline of the folded surface (Fig. 9 B.C). The formation of protrusions indicates cell convergence going towards the midline meridian of a bent surface. Such an effect took place only with gastrula stage embryos and not later. However, at advanced stages (12 – 17) a similar reaction could be observed on dissected pieces of the lateral ectoderm (Fig. 9D) which after isolation spontaneously rolled down into scrolls, the latter’s long axes coinciding with those of the entire embryos. When observing these formations after several dozens minutes, we could easily trace an extensive longitudinal contraction and formation of a series of transversal grooves and ribs (Fig. 9 E–G). This indicates cell convergence towards several more or less arbitrarily located meridians of a scroll. These experiments show that the process of meridian cell convergence is rather non-specific.

3. Relaxation rates of tensile and compression stresses and their relations to the rates of gastrulation movements.

To maintain CE feedback (Belousov and Grabovsky, this volume), embryonic tissues should be tensed. This will be possible only if the time required for the relaxed/compressed sample for at least restoring (if not hyper-restoring) its initial tension due to its active contraction (let us denote this time as $T_{ac}$) is much smaller than the time required for a passively extended tissue area to develop the active intercalation-promoted pressure ($T_{pp}$): in the opposite case the entire system will be pressurized, rather than tensed. On the other hand, the reverse $T_{pp}$ (i.e., the rate of the active pressure-generated extension) should not be too small in comparison with the rate of the epibolic tissue movements.

Fig. 9 (Left). Evidence of cell convergence on artificial folds and ectodermal “scrolls”. (A) A scheme of fold preparation; an ectodermal tissue piece situated lateral to the dorsal mid-line of a gastrula stage embryo is cut off, bent as shown by a dashed line and fixed in this position by a needle. (B,C) Protrusions formed in 10-15 h out of the bent folds (opposite to the needles). (D) If a framed region of lateral ectoderm is extirpated from the neurula stage embryo, it rolls spontaneously into a scroll, its longitudinal axis coinciding with that of embryo (E). Soon the scroll is transformed into a meridionally shrunk body with many transversal grooves and ribs (F,G). Times after scroll formation are shown.

Fig. 10 (Right). Residual deformations (RDs) in stretched or shrunk explants of embryonic tissues after their unloading. Vertical axes: percents of the positive (fixed elongation) or negative (fixed shrinkage) RDs. Initial length is taken as zero. Horizontal axes: time, minutes. (A) Protocol graphs of a stretched explant (1) and two shrunk explants (2,3). Explant 1, stretched to 70% at 5 min time and unloaded 6 min later, gradually returned to its initial length. Meanwhile, explant 2 which was shrunk to 50% and kept in this state for only 2 min gave -35% RD; explant 3 shrunk for 5 min and gave -100% RD. (B) RD after unloading 10-20% stretch in 5, 30 and 60 min after its application. (B1) Explants of the early gastrula ventral ectoderm, (B2) SBZ tissue, (B3) explants of the lateral ectoderm from neurula stage embryos. While in 5 min after force application the samples $B_1$ and $B_2$ undergo 20-10% contraction, (that is, negative RD), after 60 min stretch, $B_1$ samples gave +20% RD, $B_2$ samples just a slight RD and $B_3$ samples no RD at all.
during gastrulation, because otherwise the gastrulation movements would be hampered by increased tensions.

We tested $T_{ac}$ and $T_{ip}$ by shrinking or stretching the sandwich explants of the ventral ectoderm and SBZ taken from 10 1/2-11 1/2 stage embryos during different time periods and to different extents. The residual deformations (RD) after unloading have been measured (see Glagoleva et al., 2003, for more details). Some of explants were marked by carbon particles for testing cell movements either in a shrunk or in a stretched state. It turned out, that the time of stabilization of a shrunk state did not exceed 5 min and in about 10 min after shrinkage the active contraction movements started. Therefore, $T_{ac}$ did not exceed 10 min. On the contrary, $T_{ip}$ was no less than 30 min. If relieved earlier, a stretched explant contracted up to its unstretched state (Fig. 10A). A substantial RD could be traced only in 60 min after imposing a 10-20% stretch (Fig. 10 B.). It is noteworthy, under the same amount of deformation and stretching time the RD of SBZ samples were very small (Fig. 10 B.), reaching 20% only after 40-80% stretch (not shown). Hence, the rate of relaxation of moderate stretching stresses in the ventral ectoderm was greater than in the SBZ. Obviously, during epiboly this should promote a dorso-medial shift of embryonic tissue from the rapidly to slowly relaxed area, that is from the ventral towards dorsal. This is just what is normally observed.

Now let us compare the rate of the linear extension of an animal embryo hemisphere during epiboly with the rate of tension relaxation in the same tissue. The first value, measured just between stages 10 1/2 and 11 1/2 was found to be from 8 to 12 % per hour (Keller, 1978). Meanwhile, the explants of the blastocoel roof, stretched for an hour up to 20% of their initial length retained, after stretch relief, just 10% of their initial length. Consequently, the rates of normal gastrulation movements roughly satisfy the condition of constant tension. One may conclude that the rate of tension relaxation is the limiting parameter of gastrulation movements.

On the other hand, the lateral ectoderm explants taken from early neurula embryos (stages 13-15) did not show RD at all after any amount and duration of stretch (Fig. 10 B.). Hence, at these stages the imposed tensions are not relaxed at all. One concludes that the ability to relax tensions by cell intercalation is a stage-specific property of gastrula stage embryos only.

Step 4: Extension of the postinvoluted cell material

Theoretical interpretation

EE feedback (see Belousov and Grabovsky, this volume) is expected to be established between preinvoluted (pre) and postinvoluted (pst) cell material. Namely, at the beginning of involution pst is assumed to be passively extended by the actively extending pre, while later on pst becomes an active component of the mutually promoting EE feedback (see Fig. 1 E1, E2, e).

Predictions and experiments

While carefully detaching the adjacent pre and pst areas from each other, one should expect that at the start of involution pst will be immediately and extensively contracted (relieving the passive tension), while at the later stages it can even overlap the over lain pre. Just this was observed: if we made such a detachment at stage 11 (mid-gastrula), an extensive and immediate pst contraction and curling took place (Fig. 11A). Meanwhile, at stages 13-14 the pst, instead of being contracted, in few seconds extended, overlapping pre (Fig. 11B).

Discussion

The main goal of this study was to show that the entire process of amphibian gastrulation can be represented as a succession of mutual biomechanical reactions between the different parts of an embryo, the earlier reactions creating the initial conditions for the next ones. In this account, for the sake of brevity, several points have been omitted or just briefly discussed. Among those is the latero-ventral circular spreading of a blastoporal slit and some detail of involution. The adequate biomechanical interpretation of these events is given by Cherdantzev et al. (this volume).

We do not know any experiments or the observations of a normal gastrulation process which, after a proper analysis, contradicts our interpretation. This includes, among others, the detailed description of morphogenetic events within the isolated MZ (Davidson et al., 2002). Moreover, we believe that our interpretation can be applied to other types of gastrulation as well, at least in the case of holoblastic embryos. In all these cases, the crucial point is the establishment of CE feedback in the blastocoel walls: one (animal) part of a spherical wall becomes actively extended while another one (vegetal) becomes actively contracted. If the blastula wall includes more than one cell layer in width, the animal extension is achieved by RCI while in one cell
layered blastulae (as in sea urchins) the same should be achieved by cell flattening. On the other hand, the vegetal contraction can proceed with participation of either cell migration, or tangential cell contraction, or both together. The above described experiments (Fig. 2 and the corresponding comments) show that a segregation of a blastula wall into extended and contracted parts can go in the absence of any pre-established polarity, although normally it coincides with the earlier positioned animal/vegetal egg axis.

At a next step, the contraction of the vegetal surface will make it flatter than the animal one and so the both zones will be separated by a circle of an increased curvature which is the rudiment of a blastoporal lip. Hence taking the shape of a semi-toroidal surface, the lip should start to contract circumferentially, thus leading to the blastopore closure. Such may be a rough sketch of the processes involved in all kinds of gastrulation.

What is the relationship of our approach to research directed towards deciphering the molecular machinery of gastrulation? In our view, both approaches are complementary (see also Keller et al., 2003), because the active responses to mechanical stresses require a refined molecular machinery, whose disturbances should largely modify the rates and other parameters of the standard morphogenetic responses. Detailed studies of these effects are in progress in our research group. On the other hand, certain elements of the genes expression patterns should be considered as an initial condition for the entire morphomechanical causal chain. This relates first of all to the establishment of dorso-ventrality, which takes place long before the beginning of gastrulation (Gerhart and Keller, 1986). It is noteworthy, that this process is not purely chemical, but includes also mechanical events, namely the rotation of the egg cortex and/or yolk flows (see Cherdantzeva and Cherdantzev, this volume). It is easy to see that these mechanical forces are directed towards the relaxation/compression of the future dorsal side of an egg, as related to its ventral part. Taking into consideration that by Farge (2003) data it is pressure that initiates the migration of β-catenin to cell nuclei of Drosophila embryonic tissues, one may speculate that similar processes involved in dorsalization of amphibian embryos (Schneider et al., 1996) may be triggered by the same mechanical impulse.

Therefore we suggest that the relationships between the morphomechanics and molecular machinery are bi-directional: a molecular machinery affects mechanics and vice versa. Such a view fits the approaches, emphasizing the role of cell movements (Gordon, 1999; Salazar-Ciudad et al., 2003) or mechanical waves (Gordon, 1999) in affecting cell differentiation, because the both kinds of events are associated with mechanical stresses. Our results presented above permit us to formulate some definite questions addressed directly to the molecular level. For example, one should ask whether CCI movements taking place, after remodeling of mechanical stresses, in quite abnormal directions and locations are associated with the activation of the same signal pathways as the “standard” convergent movements. The corresponding investigations are now in progress.

Experimental Techniques

All the operations were made on Xenopus laevis embryos obtained from hormonally stimulated adults and incubated at room temperature. Operated embryos were maintained in MMR solution (100 mM NaCl, 2 mM KCl, 2 mM CaCl$_2$, 1 mM MgCl$_2$, 5 mM HEPES, pH 7.4). The optical histology, transmission and scanning electron microscopy techniques were routine (see Belousov et al., 1990, 2000). For modifying tensile stresses, the following approaches were used:

1. Relaxation of mechanical stresses either by localized incisions, or by making a radial cut through embryonic body and inserting a piece of a homologous tissue within the wound gap (Fig. 6A; see Belousov et al., 1990 for more details).

2. Artificial stretching of embryonic tissues by two 70- to 80 μm diameter glass needles attached to the agarose substrate (prepared from 2% agarose). We used either 1-fold stretching, extending a sample up to about 150% of its initial length, or 2-fold stretching (the both sets of stretching separated by a 7-10 min time period). In the last case the overall stretching reached 200% of the initial sample’s length.

For labeling embryonic tissues fluorescein-dextran (40000 MW, “Mol. Probes”), were injected into either two dorso-medial or two ventro-medial blastomeres at 32 blastomeres stage (10 ng per blastomere). Fluorescence under UV excitation was monitored with the use of an Olympus SZX9 epifluorescent microscope with fluorescent block U-RFL-T.

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