Self-Organization Phenomena in Embryonic Stem Cell-Derived Embryoid Bodies: Axis Formation and Breaking of Symmetry during Cardiomyogenesis

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Abstract
Aggregation of embryonic stem cells gives rise to embryoid bodies (EBs) which undergo developmental processes reminiscent of early eutherian embryonic development. Development of the three germ layers suggests that gastrulation takes place. In vivo, gastrulation is a highly ordered process but in EBs only few data support the hypothesis that self-organization of differentiating cells leads to morphology, reminiscent of the early gastrula. Here we demonstrate that a timely implantation-like process is a prerequisite for the breaking of the radial symmetry of suspended EBs. Attached to a surface, EBs develop a bilateral symmetry and presumptive mesodermal cells emerge between the center of the EBs and a horseshoe-shaped ridge of cells. The development of an epithelial sheet of cells on one side of the EBs allows us to define an ‘anterior’ and a ‘posterior’ end of the EBs. In the mesodermal area, first cardiomyocytes (CMCs) develop mainly next to this epithelial sheet of cells. Development of twice as many CMCs at the ‘left’ side of the EBs breaks the bilateral symmetry and suggests that cardiomyogenesis reflects a local or temporal asymmetry in EBs. The asymmetric appearance of CMCs but not the development of mesoderm can be disturbed by ectopic expression of the muscle-specific protein Desmin. Later, the bilateral morphology becomes blurred by an apparently chaotic differentiation of many cell types. The absence of comparable structures in aggregates of cardiovascular progenitor cells isolated from the heart demonstrates that the self-organization of cells during a gastrulation-like process is a unique feature of embryonic stem cells.

Key Words
Embryonic stem cells · Stem cell differentiation · Embryoid bodies · Morphogenesis · Cardiomyogenesis · Mouse

Abbreviations used in this paper

| Abbreviation | Description |
|--------------|-------------|
| AP           | alkaline phosphatase |
| CMCs         | cardiomyocytes |
| EBs          | embryoid bodies |
| ECM          | extracellular matrix |
| ESCs         | embryonic stem cells |
Introduction

Embryonic stem cell (ESC)-derived embryoid bodies (EBs) provide the environment for lineage commitment to the ectodermal, mesodermal and endodermal fate [Wobus et al., 1984; Desbaillets et al., 2000; Dvash et al., 2006; Yamanaka et al., 2008]. In suspension, aggregated ESCs form an outer layer of primitive endoderm [Murray and Edgar, 2001; Rula et al., 2007] resembling the hypoblast of the implanting blastocyst. The inner cells adopt an ectodermal fate, forming a columnar epithelium [Ikeda et al., 1999; Komura et al., 2008], which resembles the epiblast and later the early egg cylinder stage at day 6.5 (E6.5) of murine embryogenesis. Maintenance of EBs in suspension culture leads to the formation of many somatic cell types of ectodermal, mesodermal and endodermal origin. Mesoderm has been shown to emerge in a Wnt-signaling-dependent, localized and asymmetric manner in EBs kept in suspension [ten Berge et al., 2008]. Alternatively, EBs may be transferred to tissue culture plates coated with components of the extracellular matrix (ECM), where they attach via the primitive endoderm, which thereupon migrate radially from the dense center of the EBs. Primitive endoderm differentiates to the extraembryonic visceral and parietal endoderm which in vivo compose the two yolk sacs of the mammalian embryo. The primitive ectoderm undergoes locally an epithelial-mesenchymal transition which results in the development and ingression of mesoderm [Behr et al., 2005; Denker et al., 2007; Maranca-Hüwel and Denker, 2010]. If EBs have been formed from 600 to 800 ESCs, the dense center of the attached EBs enlarges and occasionally a horseshoe-shaped ridge of cells forms, which surrounds the central core of the EBs [Weitzer, 2006]. Cells in the area encircled by the horseshoe-shaped ridge of cells become committed to Brachyury-expressing primitive mesoderm [Bader et al., 2001]. These mesodermal precursors give rise to hematopoietic cells and spontaneously and rhythmically contracting cardiomyocytes (CMCs). First, CMCs start to contract at the same time as the heart tube forms at E7.5 to E8.0 from the splanchnic or lateral plate mesoderm located underneath the ectodermal head fold of the embryo. Nonetheless, lineage development in EBs which commences during gastrulation in vivo has until now been mainly considered to be chaotic.

Patterning of mesoderm is orchestrated by the interplay of many growth factors with morphogenetic activities. The influence of Tgfβ and Wnt signaling is well studied [ten Berge et al., 2008]. An Activin A/Nodal gradient influences the expression of the transcription factors Brachyury, Goosecoid, and Mix [Dvash et al., 2007; Nakaya et al., 2008]. However, Nodal is also reported to inhibit mesoderm development by supporting the self-renewal of ESCs [Vallier et al., 2004, 2009]. Both low and high levels of Nodal activate Brachyury, but with different developmental outcomes. Posterior mesoderm forms at low levels of Nodal or when Wnt3a is present. Nodal maintains expression of Brachyury, inactivates Mix, and consequently Goosecoid is repressed. In contrast, anterior mesoderm forms in the presence of high Nodal levels or when Wnt signaling is inhibited. Nodal induces the expression of Mix which in turn inactivates Brachyury by activating Goosecoid. Consequently, posterior mesoderm forms at the expense of anterior mesoderm and definitive endoderm [Swiers et al., 2010], and development of the neuroectodermal lineage can proceed.

Cardiomyogenesis is induced and regulated by a set of transcription factors and extracellular signals committing primitive mesodermal cells to the cardiogenic lineage. Sequential and parallel expression of Brachyury, MesP1, Gata4, Nkx2.5, and Isl1 in mesodermal progenitors leads to the commitment of these cells to the cardiogenic lineage, and to the expression of transcription factors instructing cells to differentiate to the myocardial, smooth-muscle and endothelial fate [Laugwitz et al., 2008]. Shortly after the expression of Brachyury, cells committed to the myogenic lineage begin to express Desmin and differentiate first to CMCs, and later on to smooth muscle cells, skeletal muscle myoblasts and myotubes, both in vivo [Kuisk et al., 1996] and in EBs [Weitzer et al., 1995]. Desmin promotes cardiomyogenesis in vitro [Hofner et al., 2007; Höllrigl et al., 2007], but is dispensable during embryogenesis due to redundant functions of other type III intermediate filament proteins [Li et al., 1996; Milner et al., 1996].

From the first description of ESC-derived EBs [Wobus et al., 1984], it became more and more evident that EBs mimic the triploblastic development of eutherian embryos. Despite the fact that ectodermal, endodermal and mesodermal somatic cells could be identified in differentiating EBs, morphological processes resembling embryonic gastrulation could so far only be identified in rhesus monkey EBs when attached to a surface [Behr et al., 2005; Denker et al., 2007; Maranca-Huwel and Denker, 2010] and in suspension cultures of mouse [ten Berge et al., 2008] and human EBs [Kopper et al., 2010]. Most other data hinted at a rather chaotic development of the three germ layers in EBs [Pekkanen-Mattila et al., 2010]. In contrast, pregastrulation development of the blastocyst leading to the formation of the epiblast and the endoblast,
including its derivatives visceral and parietal endoderm, is very well recapitulated in EBs [Ikeda et al., 1999; Bader et al., 2001; Murray and Edgar, 2001; Stary et al., 2005]. Thus, here we try to demonstrate that attachment of EBs to an ECM-mimicking implantation provides the foundation for a reproducible morphological development reminiscent of early mammalian gastrulation. Indeed, EB development follows a morphological program inherent and restricted to ESC-derived EBs. Under a rigorous regime, including constant numbers of ESCs and exact timing of the attachment of EBs, self-organization of cells in EBs establishes a bilateral symmetry which is temporarily broken when cardiomyogenesis commences. Finally, these developmental processes during early cardiomyogenesis can be disturbed by interfering with Nodal signaling or by the ectopic expression of Desmin.

Materials and Methods

**Maintenance and Differentiation of ESCs**

AB2.2 [Soriano et al., 1991], W4 [Lauss et al., 2005], 663 wild-type ESCs and 664 (hdac1–/–) [Lagger et al., 2002], DC6 (des–/–des–/–) [Hofner et al., 2007], 6.24.14.24 (des–/–) [Wetzer et al., 1995], DIC2 (lif–/–) [Dani et al., 1998] and DIA-RKO (lif–/–) [Aubert et al., 1999] ESCs were maintained in DMEM containing 2 mmol/l glutamine, 50 U/ml penicillin, 50 µg/ml streptomycin, 0.1 mmol/l β-mercaptoethanol, and 15% (v/v) fetal bovine serum (HyClone –/–). Recombinant Nodal was from R&D Systems (1315ND) and DMSO (S4317) was dissolved in DMSO at a concentration of 3 mmol/l. In the next step, unspecific antibody binding was deactivated by incubation of sections with 1 % H2O2 for 15 min in the dark. Finally, remaining horseradish peroxidase activity was blocked with 0.05% Tween 20, endogenous peroxidase was blocked with 3% H2O2 for 60 min. In the next step, unspecific antibody binding was blocked using blocking reagent from Perkin Elmer and sections were incubated with primary antibodies in a humid chamber overnight at 4°C. On the following day, sections were again washed in TNT buffer and incubated with the horseradish peroxidase-conjugated secondary antibodies corresponding to the species of the primary antibody for 30 min at room temperature. After three washes in TNT buffer, signals were amplified using TSA fluorescent dyes (1:100 in amplification diluents) for 10 min in the dark. Finally, remaining horseradish peroxidase activity was deactivated by incubation of sections with 1% H2O2 for 15 min. Sections were mounted with Prolong Gold containing DAPI (Invitrogen) and analyzed on either an LSM Meta Confocal Microscope or a Fluorescence Microscope, both from Zeiss.

**In vivo Staining of Cells in EBs and Light Microscopy**

The activity of alkaline phosphatase (AP) in cells was detected after washing EBs each once with PBS and AP buffer solution (100 mmol/l Tris–HCl pH 9.5, 100 mmol/l NaCl, 50 mmol/l MgCl2) and the addition of a freshly prepared solution of 0.34 mg/ml NBT and 0.18 mg/ml BCIP in AP buffer at room temperature for 5 min.

To detect heavily glycosylated surfaces of primitive, visceral and parietal endoderm cells [Stary et al., 2005], a 0.1% (w/v) toluidine blue solution was used to stain sections.

**Confocal Immunofluorescence Microscopy**

EBs on coverslips were washed once in PBS and fixed in 96% ethanol at −20°C for 20 min, then dried and blocked with 1% bovine serum albumin in PBS for 10 min. After washing EBs in PBS for 10 min, they were stained with antibodies against cardiac Troponin T (Thermo Scientific #MS-295, 1:200), Connexin 43 (Sigma C6219, 1:400), β-catenin (Transduction Labs C19220, 1:200), Desmin (Sigma-Aldrich D8281, 1:200), Ki67 (Novocastra Laboratories, NCL Ki67p, 1:1,000), Snail 1 (Abcam, ab7732, 1:100) and E-cadherin (BD Transduction Laboratories, #610181, 1:100) for 60 min, and consecutively with FITC-, Alexa Fluor 488-, and TRITC-conjugated secondary antibodies (Dianova, 711-095-12, 1:200; 711-095-151, 1:200; 711-075-152, 1:200, Invitrogen A11008, Sigma F4018, 1:200, T5268, 1:80) for 60 min. Secondary antibodies were tested alone to exclude background staining. Nuclei were stained with DAPI or propidium iodide, and photomicrographs were taken on a Zeiss LSM 510 confocal microscope.

Alternatively, EBs maintained in suspension culture were fixed in 4% para-formaldehyde overnight at 4°C, then washed twice in cold PBS for 30 min at 4°C. Until embedding, EBs were stored in 70% ethanol. Due to the small size of EBs, 5–10 EBs were collected and sealed in low-melting agar in order to facilitate dehydration and sectioning. Consequent dehydration of EBs in agar was performed according to standard histological methods using the Excelsior tissue processor. After paraffin embedding, EBs were sectioned into 5-µm tissue sections. For staining EB paraffin sections, the TSA Kit (Perkin Elmer) was used. In short, sections were heated to 60°C for 1 h and rehydrated. Antigens were retrieved by boiling the sections in citric acid for 20 min. After three washing steps in TNT buffer (0.1 mol/l Tris–HCl pH 7.5, 0.15 mol/l NaCl, 0.05% Tween 20), endogenous peroxidase was blocked with 3% H2O2 for 60 min. In the next step, unspecific antibody binding was blocked using blocking reagent from Perkin Elmer and sections were incubated with primary antibodies in a humid chamber overnight at 4°C. On the following day, sections were again washed in TNT buffer and incubated with the horseradish peroxidase-conjugated secondary antibodies corresponding to the species of the primary antibody for 30 min at room temperature. After three washes in TNT buffer, signals were amplified using TSA fluorescent dyes (1:100 in amplification diluents) for 10 min in the dark.
Fig. 1. Radial symmetry of solitary EBs in hanging-drop cultures. A 800 ESCs were aggregated in 20-μl drops for 4.5 days and then transferred to the suspension culture. ESCs aggregate within 1.5 days (a) and the compacted EB grows in size from day 2 to day 4.5 (b). At day 4.5 most cells are still AP-positive (c) with the exception of the outer layer of primitive endoderm (arrow). Primitive endoderm increasingly expresses Snail (red) and E-cadherin (green) (d, e). At the same time a Reichert’s membrane-like borderline (arrow) between the inner cells and the outer layer of primitive endoderm forms (f). If EBs were maintained in suspension culture the AP-positive cells became less from day 5 onwards (g) and a central cavity (*) formed, EBs developed mostly an irregular shape (h) and cells were distributed in an apparently chaotic manner as made evident by E-cadherin (green) and Snail (red) staining at day 10 (i). DNA was stained with DAPI (blue). a–i Bars: 100 μm. B Cartoon demonstrating the early development of cell types in EBs. Pink: ESCs and primitive ectodermal cells; yellow: primitive endoderm. C If EBs were transferred from the hanging drops to tissue culture plates at day 2 (a) or 3 (b) they did not further differentiate but remained as ESC-like colonies (arrowheads indicate area where primitive endoderm usually starts to grow and give rise to parietal endoderm; * = ESC-like cells). When plated on day 4.5, the primitive endoderm readily attached to the gelatine-coated tissue culture plate and differentiation proceeded with a radial symmetry (bracket in c). If plated on day 5.5, many EBs developed an irregular shape (d); however, differentiation proceeded normally in most cases 1–2 days later (e). If EBs were transferred to the tissue culture plate on day 6.5 or later, their development was either totally blocked (f, arrowheads as in a) or only extraembryonic endoderm formed (brackets in g and h), while the inner cells remained entrapped within the Reichert’s membrane-like structure (arrows in g and h). Phase contrast images taken 1 day (a–d, f, h) and 2 days (e, g) after the attachment of EBs. Bars: 100 μm.
The culture medium was washed once with a freshly prepared solution of 0.2% dithiothreitol (DTT) in PBS. After 10 min at room temperature plates were washed once with PBS. Hemoglobin in hematopoietic cells was detected by mixing the culture medium 1:1 with a freshly prepared solution of 0.2% DTT in PBS to remove hemoglobin. After 10 min at room temperature plates were washed once with PBS. When attached, EBs were stained with a propidium iodide solution in PBS to identify dead cells. After 10 min at room temperature plates were washed once with PBS. Pictures were taken on an Olympus CK2 microscope under bright-field, phase-contrast, or dark-field illumination and prepared for publication by Adobe Photoshop CS2. Where indicated, pictures were composed from several microphotographic images or false coloring was overlaid to indicate areas with rhythmically contracting CMCs.

**Semiquantitative RT-PCR**

Total RNA was isolated from AB2.2 ESC-derived EBs on days 0 (ESCs), 4, 6, 9 and 12 with the Qiagen RNeasy kit. cDNA was synthesized with RevertAid M-MuLV Reverse Transcriptase (Fermentas, #EP0441). Semiquantitative RT-PCR was performed with Taq polymerase (Fermentas, #EP0402) using the following primer pairs: AFP forward: 5'-GCTCAACCAAAAGCGTCAAC-3', AFP reverse: 5'-CCTGTGAACTCTCTGATCAG-3', Brachury forward: 5'-ATCAAGGAGGCTTTAGCGG-3', Brachury reverse: 5'-GAACCTGGATACATCGGTGAG-3', Desmin forward: 5'-TGATGAGGCAGATGAGGGAG-3', Desmin reverse: 5'-CGGCCATCACGCCACAGTTT-3', GAPDH forward: 5'-GCGGATCCAGCGCAGAGG-3', GAPDH reverse: 5'-CCGTCTCCACCACTGGAGA-3', Goosecoid forward: 5'-GGAGGATGCGTCTGCTGT-3'. The number of cycles for each pair of primers was carefully optimized so that none of the obtained signals was saturated and equalized to GAPDH expression.

**Statistical Analysis**

Data are presented as the arithmetic mean ± standard deviation. Statistical significance was evaluated using the one-sample Student t test and values of p < 0.05 were considered to indicate statistical significance.

**Results**

**Self-Organization in Early EBs Alludes to the Bilaterian Body Plan**

Here we describe that EBs undergo axis formation when they are first maintained individually in hanging-drop cultures and then transferred to tissue culture plates at a density of 1 EB/cm² at day 4.5 of their development. In hanging drops, nearly all EBs develop in a radial symmetrical manner for the first 4.5 days (fig. 1A, B). Since EBs never develop uniformly in 100% of a population, quantitative data for the images shown in figures 1–6 are provided in table 1. During this time ESCs aggregate, compact (fig. 1A, a, b), give rise to an AP-expressing inner epiblast-like cell mass (fig. 1A, c) and an outer layer of primitive endodermal cells [Bader et al., 2001], increasingly expressing Snail and E-cadherin (fig. 1A, d, e). These two compartments were separated by a Reichert’s membrane-like ECM [Henry and Campbell, 1998; Murray and Edgar, 2001] (fig. 1A, f).

If further maintained in suspension mass culture, many EBs form a cavity and the number of AP-expressing cells declines (fig. 1A, g). After 3–5 days, EBs develop an asymmetrical shape (fig. 1A, h) with a rather chaotic distribution of cells at day 10 (fig. 1A, i). The time of transfer of EBs from hanging-drop cultures to plates coated with gelatine is crucial for the proper development of EBs (fig. 1C). If EBs are transferred too early, as already described in detail in Bader et al. [2001], no critical mass of cells can develop and thus no primitive endoderm forms (fig. 1C, a). Most of these too-small EBs die and

| Table 1. Quantitative morphological data supporting the significance of images shown in figures 1–6 |
|----------------------------------|---------------------------------|-----------------|-----------------|
| **Figure** | **Frequency of occurrence (mean value) % of EBs** | **Experiments n** | **Error SD** |
| 1A, a, b | 99.6 | 800 | 10 | 0.8 |
| 1A, c | 100 | 170 | 2 | 0 |
| 1A, h | 82.3 | 170 | 4 | 3.3 |
| 1C, d | 82 | 300 | 10 | 12 |
| 2A–G | 99.6 | 800 | 10 | 0.8 |
| 2H–J, 6D | 74 | 79 | 2 | 20.2 |
| 2K, L1 | 47.8 | 54 | 9 | 31.1 |
| 2M–S, 6D | 50.4 | 944 | 7 | 7.5 |
| 2X, Y | 100 | 800 | 10 | 0 |
| 3A, E | 100 | 37 | 1 | – |
| 3B, C1 | 32 | 25 | 1 | – |
| 3F | 100 | 800 | 10 | 0.0 |
| 4A–C2 | 89 | 624 | 8 | 11.0 |
| 4G2 | 81 | 148 | 8 | 19 |
| 5A–C | 100 | 14 | 4 | 0 |
| 6B1 | 99 | 446 | 10 | 1.1 |
| 6C | 89 | 706 | 4 | 2.1 |
| 6E1 | 100 | 880 | 10 | 0 |
**Fig. 2.** Attachment of EBs to a surface leads to the development of a bilateral symmetrical cell aggregate. 

A–E Attachment of EBs to gelatine-coated tissue culture plates between days 4.5 and 6. C Arrow shows first attaching primitive endodermal cells. 

D, E Brackets show proliferating primitive endoderm. F, G Development of EBs between days 5 and 6. F Arrow shows migrating primitive endodermal cells. G, H Brackets show parietal endodermal cells undergoing an epithelial-mesenchymal transition. 

H In the meantime, the core of the EBs starts to elongate. J Cartoon: cross section of a typical EB at days 5 and 6. Inner cells of primitive ectoderm (pink) are surrounded by primitive endoderm (blue) which differentiates upon contact with the gelatine-coated tissue culture plate (green) into parietal endoderm (yellow) and visceral endoderm on top of the primitive ectoderm, respectively. 

J–L Breaking of the point symmetry by elongation of the central dense core of the EBs between day 6.5 and 7.5. Dashed lines indicate the forming axis. Solid lines emanating from the center (dots) of EBs indicate the asymmetric elongation of the EB core. K, L Arrows indicate areas where the thin cell layer (mesoderm?) forms which later separates the horsehoe-shaped area from the center of the EBs. 

M–Q Development of the horseshoe-shaped ridge of cells surrounding the core of the EBs between days 7 and 8.5. R, S Epithelial sheet of cells developing at the open side of the horseshoe-shaped ridge at day 6.5. Dashed line demarcates epithelial sheet of cells; dot shows center of EB. Inset β-Catenin staining of epithelial cells (green); bar: 10 μm. T, U Hematopoietic cells develop inside of the horseshoe-shaped area of EBs. Inset Single erythrocyte-like cells; bar: 10 μm; benzidine stain. 

V Cartoon of EB at day 8: 1 transverse section through the center of the EB. W Pseudo-3-D image (tilted illumination under a stereomicroscope) of a day-8 EB where pseudocoloring indicates the area of developing presumptive mesoderm (brown), the epithelial sheet of cells at one end of the EB (yellow), and the dense horseshoe-shaped area of presumptive ectoderm (black) which is surrounded by the extraembryonic endoderm (green-blue). The central black area may be a remnant of the primitive endoderm covered by visceral endoderm. 

X, Y Blurring of the horseshoe-shaped ridge by cell growth and migration starting at day 8.5–9.0. Dashed lines indicate 2-fold axes. A–E, J–L, O, U, V Bright-field illumination. F–H, M, N, R, S Phase-contrast illumination. P, Q, X, Y Dark-field illumination. Bars: 100 μm except for S: 50 μm.
those which survive and attach to plates remain as small colonies with an ESC-like morphology (fig. 1C, b). When plated at day 4.5, almost all EBs have a uniform, radial symmetrical morphology (fig. 1C, c) with a flat outer layer of parietal endoderm. If plated after day 4.5, a large number of EBs develop an irregular shape (fig. 1C, d), but 1–2 days later proceed in their development similarly to those plated on day 4.5 (fig. 1C, e). If plated after day 6.5, the majority of EBs become encapsulated and blocked by their own overgrowing primitive endoderm (fig. 1C, f), which may give rise to parietal endoderm (brackets in fig. 1C, g, h), but at the same time physically prevents the inner cells escaping from the endodermal shell by forming a thick Reichert’s membrane-like structure (arrows in fig. 1C, g, h).

In contrast, when transferred from hanging drops to gelatine-coated tissue culture plates at day 4.5, axis formation becomes evident in EBs by the development of morphological distinct cell populations (fig. 2). Spherical EBs attach via their outermost cell layer of primitive endoderm to the culture plates (fig. 2A–C). Primitive endoderm starts to proliferate radially around the dense core of the EBs composed of undifferentiated cells (fig. 2D, E) between days 4.5 and 5.5. Then primitive endoderm cells migrate distally (fig. 2F) and simultaneously develop into parietal endoderm (fig. 2G). Radial symmetry is maintained during this short period of time. On top of the core of EBs, the remaining primitive endoderm develops into visceral endoderm [Bader et al., 2001; Stary et al., 2005] (fig. 2I). Between days 6 and 7.5 an axis forms which becomes evident, firstly, by an elliptical shape of the dense center of the EBs (fig. 2H, J, 6B), secondly, by the formation of a cavity on the elongated side of the dense center filled with a thinner layer of cells (fig. 2K, L), and finally, by the formation of a dense horseshoe-shaped ridge which migrates distally away from the also-dense core of the attached EBs (fig. 2M–Q, 6B). At the same time, radial symmetry is also broken by the appearance of an epithelial, β-catenin-positive sheet of cells which reside at the opening of the horseshoe-shaped ridge. This allows definition of, in addition to the 2-fold axis, an ‘anterior’ and a ‘posterior’ half of the EBs (fig. 2R, S). At the inner side of the horseshoe-shaped ridge hematopoietic cells develop (fig. 2T, U), which suggests that primitive mesoderm evolves in this part of the EBs, as depicted in the cartoon (fig. 2V) and the false-colored picture of an EB (fig. 2W). This finding is in line with a previous observation of Brachyury-expressing cells in this part of EBs [Bader et al., 2001].

This morphological development only takes place when EBs are not disturbed by secreted factors coming from other EBs in the close vicinity, and when the initial number of aggregated ESCs ranges between 500 and 800 cells. The latter also correlates well with the observation that cardiomyogenesis was optimal when these ESC numbers were used to generate EBs [Bader et al., 2001]. Development of these morphological features was observed in 3 independently generated and genetically distinct ESC lines, AB2.2, W4 and 663. After day 8, the axis became blurred by cell proliferation and/or migration within the center and horseshoe-shaped ridge of the EBs (fig. 2X, Y), or by excessive proliferation of visceral endoderm which formed large epithelial bubbles on top of the EBs (fig. 3F).
During the same time interval as primitive ectoderm gives rise to mesoderm and the horseshoe-shaped ridge defines a bilateral axis and an anterior and posterior end of EBs, the epithelial primitive endoderm on top of the center of EBs (fig. 3A, dark purple) develops into both epithelial visceral endoderm (dark purple) doming above the invisible ectodermal core, and parietal endoderm (bright rose) circumjacent to the central part of EBs (fig. 3B). Between days 6 and 7 endodermal cells are always more dense at one side of the EBs than in other areas (fig. 3B, C), which may well reflect the formation of a bilateral axis in the underlying primitive ectoderm. These densely packed cells on one side of the EBs might well correlate with the epithelial cell sheet next to the opening of...
the horseshoe-shaped ridge (fig. 2R). Very rarely visible, the endoderm remains attached to the core of the EBs, as suggested in the cartoon in figure 2V, and the underlying horseshoe-shaped morphology becomes visible (fig. 3D).

After day 8 of the EB development, again the axis becomes blurred by chaotic growth of all three germ layers (fig. 3E) and finally endoderm develops into large, randomly located bubbles composed of epithelial cells which cover and obscure the inner parts of all EBs (fig. 3F).

**Cardiomyogenesis Temporarily Breaks the Bilateral Symmetry in EBs**

In mammals, the heart is the first organ which forms during embryogenesis. Similarly, CMCs are the first somatic cells whose function becomes apparent in EBs. Rhythmically contracting CMCs emerge exclusively in the mesodermal part between the dense core and the dense horseshoe-shaped ridge of EBs (fig. 4A). To determine whether CMCs form in random distribution...
Throughout the mesodermal area or in some ordered fashion in relation to the 2-fold axis of the EBs, we determined the location of the first beating cluster of CMCs in EBs between day 6.7 and day 7.5 (fig. 4B). The percentage of EBs with beating CMCs when the maximum was reached on day 11 after aggregation was 89\% (table 1). CMCs started to beat in 83\% of these EBs (n = 373) in the lower half, where the horseshoe-shaped ridge opened and the epithelial, β-catenin-positive sheet of cells formed. From these EBs, 69\% had a single asymmetrically located cluster of beating cells on one side of the 2-fold axis near the epithelial cell sheet, and 32\% had 2 symmetrically located clusters of beating CMCs on both sides of the 2-fold axis. Of those with only a single cluster on one side of the axis, 52 ± 18\% started to contract in the lower left area (observed from the ‘dorsal’ side of the EBs facing the medium), and 25 ± 8\% in the lower right area (n = 261). In the upper area of the EBs, 14 ± 4\% started to contract on the left side and 9 ± 6\% on the right side. The color scheme on top of the representative EB (fig. 4C) depicts the location where the majority of first CMCs emerged (purple area) and the area where CMCs may have emerged later on during development (green area). To identify early CMCs in the lower left quadrant of EBs, we used Desmin antibodies because Desmin is one of the first muscle-specific proteins ex-
pressed in CMCs long before regular myofibrils form [Kuisk et al., 1996], in conjunction with Connexin 43 antibodies which detect both gap junctions in epithelial cells and costameres in CMCs (fig. 4D). These CMCs still have the capacity to proliferate (fig. 3E) and develop into myofibrils containing CMCs (fig. 4F).

These data together suggest that the morphology of EBs influences the local appearance of the first rhythmically contracting clusters of CMCs. The clear bias towards the 'lower left quadrant' of the EBs (fig. 4B, C) breaks the axial symmetry of the EBs and may allude to the asymmetric location of organs in bilateria later in development. At day 8.2 ± 0.5, many EBs have large and interconnected clusters of rhythmically contracting CMCs which surround the center of EBs in a ring-like structure which always remains localized clearly within the horseshoe-shaped ridge (fig. 4G).

Mesoderm Formation and Cardiomyogenesis Requires ALK-4-Mediated Signaling in EBs

Nodal is a key regulator of mesoderm development which targets mesoderm-regulating genes such as Lefty A/B and Brachyury via the ALK4 receptor kinase and Smad proteins [Dvash et al., 2007]. To test whether mesoderm formation indeed takes place in the area between the dense core and the horseshoe-shaped ridge, and whether the horseshoe-shaped ridge in EBs depends on Nodal signaling, we added the ALK4 inhibitor SB431542 to the medium starting from day 3 of EB development, right before Brachyury expression became evident in the EBs (fig. 6A). In the presence of the 3.8 μmol/l SB431542 the number of dying cells was apparently increased in a ring-like area around the center of the EBs at day 7 (fig. 5A). In the absence of SB431542 but in the presence of DMSO, only very few cells, scattered all over the EBs, were dying. Formation of a horseshoe-shaped ridge was never observed. At day 11, dead cells accumulated in a ring-like area around the center of the EBs (fig. 5B), in which most cells had become permeable for propidium iodide (fig. 5C). Increased death of cells in the mesodermal compartment of EBs resulted in a significant delay of cardiomyogenesis by 2 days (fig. 5D) and decreased cardiomyogenesis (fig. 5E) in the presence of SB431542. Most importantly, the negative effect of SB431542 on cardiomyogenic mesoderm was significantly attenuated when desmin [Hofner et al., 2007], known to promote the differentiation and survival of myocardial mesoderm, was overexpressed in EBs. Notably, the addition of 58 nmol/l Nodal from day 3 onwards did partially inhibit cardiomyogenesis, most likely because Nodal itself also prevents differentiation of ESCs [Vallier et al., 2004, 2009]. Again, Desmin expression attenuated the negative effect of Nodal in EBs, which suggests that SB431542 indeed specifically blocks ALK4 and affects the development of mesoderm. These data together support the morphological observations, are in line with previous reports [Behr et al., 2005; Denker et al., 2007; Maranca-Huwel and Denker, 2010] and suggest that mesoderm indeed forms in EBs by the ingestion of cells around the center of EBs between days 6 and 7.

Ectopic Expression of Desmin in EBs Randomizes the Localization of the First CMCs and Disturbs the Morphology of EBs

We previously demonstrated that ectopic and constitutive expression of Desmin protein promotes cardiomyogenesis in EBs by the upregulation of mesodermal transcription factor Brachyury and Nkx2.5 [Hofner et al., 2007] and that a desmin null mutation and the ectopic expression of dominant negative mutant desmin alleles hamper cardiomyogenesis [Höllrigl et al., 2007]. RT-PCR analysis of the expression of mesodermal transcription factors Brachyury and Goosecoid in developing EBs demonstrates that desmin mRNA is expressed in advance of Brachyury mRNA (fig. 6A). From these data we reasoned that the ectopic expression of Desmin protein may increase the commitment of primitive mesoderm towards the cardiomyogenic lineage, and consequently influence the development of the mesoderm at very early stages and the appearance of the outlying horseshoe-shaped ridge in EBs. Until day 5–6, when the asymmetry of the centers of EBs became visible, EBs with an ectopic expression of des+/+desect EBs had a uniform central mass of cells with a random distribution of contractile CMCs (fig. 6C) and no bilateral axis became visible.

The development of the asymmetric dense core of EBs between days 5 and 6.5 was indistinguishable between wild-type and des+/+desect EBs (fig. 6D, solid lines) but the frequency of horseshoe-shaped ridges was significantly reduced in des+/+desect EBs and the evolution of this structure was also delayed (fig. 6D, dashed lines). Notably, the epithelial sheet of cells at one end of the wild-type EBs was never observed in des+/+desect EBs and the first CMCs were always evenly distributed throughout the mesodermal area. However, cardiomyogenesis commenced earlier than in wild-type EBs and the number of EBs with CMCs...
clusters was significantly increased (fig. 6D, dotted lines). This effect was Desmin specific and not caused by the genetic manipulation of ESCs because homozygous knock-out of the histone deacetylase 1, the leukemia inhibitory factor and the leukemia inhibitory factor receptor loci did not interfere with the morphological development observed in wild-type EBs, but did influence cardiomyogenesis later on [data not shown; Bader et al., 2001; Lagger et al., 2002; Lauss et al., 2005; Lagger et al., 2010]. Therefore, disturbing the development of mesoderm by overexpression of Desmin, pushing the cells to become CMCs, or selectively killing some mesodermal cells with SB431542 negatively affects morphogenesis in EBs.

To test whether self-organization and morphological development are a general phenomenon in aggregated stem cells or restricted to pluripotent ESCs, we aggregated cardiovascular progenitor cells in the same way as ESCs. Cardiovascular progenitor cells were isolated from newborn mouse heart, (unpubl. results). After aggregation for 4.7 days, cardiovascular progenitor cell aggregates were transferred to gelatine-coated tissue culture plates, where they attached and cells started to divide and migrate distally similar to ESC-derived EBs (fig. 6E). These cells give rise to CMCs, smooth muscle and endothelial cells [Srivastava and Ivey, 2006]. Notably, none of these aggregates ever developed a horseshoe-shaped ridge or an epithelial sheet of cells, which break the point symmetry in EBs. First CMCs developed in even distribution throughout the entire center of all aggregates similar to those EBs where Desmin was constitutively expressed in two dimensional system similar to ESC-derived EBs (fig. 6F). These cells give rise to CMCs, smooth muscle and endothelial cells [Srivastava and Ivey, 2006]. Notably, none of these aggregates ever developed a horseshoe-shaped ridge or an epithelial sheet of cells, which break the point symmetry in EBs. First CMCs developed in even distribution throughout the entire center of all aggregates similar to those EBs where Desmin was constitutively expressed in all cells. From these results we may infer that self-organization of germ lineages is restricted to ESC-derived EBs.

**Discussion**

ESC were studied mainly to understand the molecular mechanisms involved in self-renewal and the differentiation to somatic cells. Aggregation of ESCs in EBs turned out to be a strong signal inducing the differentiation of ESCs, which leads to the development of numerous somatic cells descending from all three germ layers. Cell biological analysis of differentiated cells, as well as the expression of lineage-specific genes during the differentiation processes, suggests that EBs undergo a developmental process resembling gastrulation in triploblastic embryos. Beyond this molecular data, only a few authors have thus far contributed to answering the question whether these differentiation processes are based on, or correlate with, morphological phenomena in EBs such as the self-organization of specific cells or pattern formation. Apart from very early approaches to describe the self-organization of cells in embryocarcinoma aggregates by Stevens [1959, 1960] and Pierce and Dixon [1959a, b], only Denker and colleagues [Behr et al., 2005; Denker et al., 2007; Maranca-Hüwel and Denker, 2010] and Nusse and colleagues [ten Berge et al., 2008] have convincingly demonstrated that gastrulation and axis formation may occur in EBs.

Here we demonstrate that a timely implantation-like process is a prerequisite for the breaking of the radial symmetry of EBs. Once attached to a surface, EBs develop a bilateral symmetry and mesodermal cells develop between the dense center of the EBs and a distal horseshoe-shaped ridge. The development of an epithelial sheet of cells at one side of the EBs but symmetrical to the axis allows definition of an ‘anterior’ and a ‘posterior’ end of the EBs. In the mesodermal area, first CMCs develop mainly next to the epithelial sheet of cells at one side of the bilateral EBs. Development of twice as many CMCs on the ‘left’ side of the EBs suggests that cardiomyogenesis causes or reflects a (temporal) asymmetry in EBs, which breaks the bilateral symmetry.

In vitro, in EBs it has so far generally been accepted that differentiation of germ layers is chaotic, that no organized migration of cells takes place and that the oxygen and nutrition supply by vessels is rudimentary or missing, and thus inefficient in supporting the self-organization of cells in EBs. Since chaos may be described as the most complex conceivable system, our constant effort is to reduce this complexity and demonstrate the simplicity of certain developmental processes in EBs. Therefore, we and many others developed a protocol which allowed the reproducible generation of uniform EBs. Reduced complexity resulted in the visibility of self-organization of cells in EBs. Most important was that aggregation of EBs started from a defined number of viable ESCs in the absence of any residual feeder cell [Bader et al., 2000a, 2001]. Secondly, the time of transfer of EBs to an ECM-coated tissue culture plate turned out to be essential for the consecutive ordered development. Another important condition was that EBs should not be influenced by close neighbors. If this occurred, we did not obtain a horseshoe-shaped ridge but rather irregular, or at best circular, structures around the center of EBs. Finally, the transfer of EBs at day 4.5 may have mimicked the implantation process in the murine uterus and induced primitive endoderm to differentiate to visceral and parietal endoderm, respectively. Parietal and visceral endoderm are well-described sources of growth factors supporting the...
differentiation of various cells which emerge during gas-
trulation. Notably, in many EBs, endodermal cells are
more densely located at one side of the EB, which might
coincide with or even reflect the development of the epi-
thelial sheet and axis formation.

Mesoderm formation followed the development of
parietal endoderm presumably in cavities where cells
seemed to ingress in elongating EBs. This could be the
beginning of gastrulation by an epithelial-mesenchymal
transition as described by Denker and colleagues [Behr et
al., 2005; Denker et al., 2007; Maranca-Hüwel and Den-
ker, 2010]. This process also marks the beginning of the
axis formation. The cavities expand around the dense
center of EBs and constitute the horseshoe-shaped ridge
of EBs. The mesodermal cells then proliferate, perhaps
pushing the horseshoe-shaped ridge more distally. Devel-
opment of hematopoietic cells and CMCs within this
horseshoe-shaped ridge strongly supports the previous
finding that mesoderm forms in a central pit-like area of
EBs. Induction of neurogenesis by retinoic acid (unpubl.
results) exclusively gave rise to neurons at the periphery
of the horseshoe-shaped ridge from day 8 onwards, which
suggests that the remaining ectoderm is pushed outwards
by the developing mesoderm. In conclusion, involution,
migration and specification of mesoderm during EB de-
velopment seems to recapitulate at least some early as-
pects of gastrulation.

While investigating the exact timing of cardiomyo-
genesis [Hofner et al., 2007; Höllrigl et al., 2007] it became
evident that the spontaneous and rhythmic contraction
of CMCs was unevenly distributed in the mesodermal
ring-like area encircled by the horseshoe-shaped ridge.
The majority of the first contracting CMCs developed on
the ‘lower left’ side of EBs. Thus, commencing cardio-
myogenesis seems to break the bilateral symmetry of EBs.
One possible explanation for this unexpected finding can
be the development of a morphogen gradient by ciliar
movement under the dome of visceral endoderm which
forms in many EBs on top of the gastrulating primitive
ectoderm. Visceral endoderm in particular may shield
the core of EBs from the medium or other environmental
influences and provide a space in which morphogenetic
gradients may form. ESCs possess primary cilia [Kiprilov
et al., 2008] which, if maintained during the development
of primitive ectoderm and gastrulation [Takaoka et al.,
2007], may well generate a morphogen gradient by syn-
chronous and unidirectional rotation. A unidirectional
flow and a morphogen gradient under the domeing vis-
ceral endoderm could contribute to the breaking of the
bilateral symmetry.

EBs also express increasing amounts of Nodal be-
tween days 3 and 5 [unpubl. results; Vallier et al., 2004] which
might be unevenly distributed in or above EBs in
a similar fashion as the flow of extraembryonic fluids cre-
ates morphogen gradients in embryos [Nonaka et al.,
1998]. We were not able to demonstrate that Nodal indeed
had this effect on breaking symmetry because the addi-
tion of recombinant Nodal always caused an increase in
parietal endoderm formation obscuring the morphology
of EBs and partially inhibited cardiomyogenesis. How-
ever, inhibition of endogenous, mesoderm-inducing
Nodal signaling, by blocking the kinase activity of ALK4
with SB431542, resulted in a selective death of mesoder-
mal cells in the ring-like area surrounding the dense core
of EBs. Alternatively, asymmetric development of the
first CMCs may be caused by paracrine factors secreted
by the epithelial sheet or some other unidentified neigh-
boring cell types. Asymmetry is observed only during the
development of the very first CMCs. Later on CMCs
emerge in a radial symmetric pattern scattered all over the
mesodermal area encircled by the horseshoe-shaped ridge.

Previously, we and others could demonstrate that the
muscle-cell-specific intermediate filament protein Des-
min influences cardiomyogenesis at very early stages
[Hofner et al., 2007; Höllrigl et al., 2007] and here we pro-
vide evidence that Desmin is expressed earlier than the
maximum expression of Brachyury and Goosecoid in
EBs. From these results we reason that the ectopic expres-
sion of Desmin might interfere with the asymmetric de-
velopment of the first CMCs in the ring-like mesoderm.
The ectopic expression of Desmin did not interfere with
axis formation in EBs, but indeed hampered the forma-
tion of the horseshoe-shaped ridge in most EBs. Concom-
antly, first CMCs evenly dispersed throughout the me-
sodermal area started to contract and cardiomyogenesis
was consequently strongly increased [Hofner et al., 2007].
Thus uniform and constitutive expression of Desmin in
all cells prevents the temporal formation of an asymmet-
ric pattern but not the development of the mesoderm it-
self.

The total absence of axis formation in aggregates made
of cardiovascular progenitor cells and the uniform distri-
bution of CMCs throughout the entire aggregate, without
any bias in regard to their localization, suggest that gas-
trulation-like processes observed in all cells prevent the
temporal formation of an asymmetric pattern but not the
development of the mesoderm itself.
that EB development clearly deviates from murine embryogenesis once organs start to form in the murine embryo between days 7 and 8. In EBs we only saw the development of certain cell types but neither formation of complex tissues nor organs.

Once these conditions are translated to human ESCs, EBs may well become a valuable model system not only for the in vitro analysis of genetic defects and gene function, which has contributed to reducing the number of animal experiments in the past, but also for early mor-phogenetic processes recapitulating pregastrulation and early gastrulation of the human embryo in vitro; this could not otherwise be studied due to ethical reasons.

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