A Comparative Analysis of Extra-Embryonic Endoderm Cell Lines

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Abstract

Prior to gastrulation in the mouse, all endodermal cells arise from the primitive endoderm of the blastocyst stage embryo. Primitive endoderm and its derivatives are generally referred to as extra-embryonic endoderm (ExEn) because the majority of these cells contribute to extra-embryonic lineages encompassing the visceral endoderm (VE) and the parietal endoderm (PE). During gastrulation, the definitive endoderm (DE) forms by ingestion of cells from the epiblast. The DE comprises most of the cells of the gut and its accessory organs. Despite their different origins and fates, there is a surprising amount of overlap in marker expression between the ExEn and DE, making it difficult to distinguish between these cell types by marker analysis. This is significant for two main reasons. First, because endodermal organs, such as the liver and pancreas, play important physiological roles in adult animals, much experimental effort has been directed in recent years toward the establishment of protocols for the efficient derivation of endodermal cell types in vitro. Conversely, factors secreted by the VE play pivotal roles that cannot be attributed to the DE in early axis formation, heart formation and the patterning of the anterior nervous system. Thus, efforts in both of these areas have been hampered by a lack of markers that clearly distinguish between ExEn and DE. To further understand the ExEn we have undertaken a comparative analysis of three ExEn-like cell lines (END2, PYS2 and XEN). PYS2 cells are derived from embryonal carcinomas (EC) of 129 strain mice and have been characterized as parietal endoderm-like [1], END2 cells are derived from P19 ECs and described as visceral endoderm-like, while XEN cells are derived from blastocyst stage embryos and are described as primitive endoderm-like. Our analysis suggests that none of these cell lines represent a bona fide single in vivo lineage. Both PYS2 and XEN cells represent mixed populations expressing markers for several ExEn lineages. Conversely END2 cells, which were previously characterized as VE-like, fail to express many markers that are widely expressed in the VE, but instead express markers for only a subset of the VE, the anterior visceral endoderm. In addition END2 cells also express markers for the PE. We extended these observations with microarray analysis which was used to probe and refine previously published data sets of genes proposed to distinguish between DE and VE. Finally, genome-wide pathway analysis revealed that SMAD-independent TGFbeta signaling through a TAK1/p38/JNK or TAK1/NLK pathway may represent one mode of intracellular signaling shared by all three of these lines, and suggests that factors downstream of these pathways may mediate some functions of the ExEn. These studies represent the first step in the development of XEN cells as a powerful molecular genetic tool to study the endodermal signals that mediate the important developmental functions of the extra-embryonic endoderm. Our data refine our current knowledge of markers that distinguish various subtypes of endoderm. In addition, pathway analysis suggests that the ExEn may mediate some of its functions through a non-classical MAP Kinase signaling pathway downstream of TAK1.

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Introduction

Studies in amphibians, avians and mice demonstrate that endodermal cells play both inductive roles and make important cellular contributions to organ formation. Endodermally derived organs such as the liver and pancreas serve important secretory functions that are required for homeostasis in the adult organism and because of this, much effort has been exerted in recent years toward the development of protocols for the directed differentiation of specific endodermal subtypes. Toward these efforts, the identification of secreted endodermal factors that mediate their inductive functions would also be highly desirable. However, these efforts have been hampered by a lack of markers that efficiently distinguish one type of endoderm from another. One possible reason for this is that endoderm constitutes only a small percentage of cells in the developing embryo, and consequently, slow progress has been made in the identification of regional specific markers within the endoderm. Furthermore, it has been noted that there is tremendous overlap in marker expression between the visceral extra-embryonic endoderm and the gut endoderm of the embryo.
Recent efforts to characterize markers that distinguish these lineages have relied on endoderm derived from ES cell sources followed by FACS purification with the aid of antibodies that recognize different types of endoderm [2,3]. While these approaches have identified multiple lineage restricted endodermal markers not all of the “hits” have been validated by further experimentation.

In the mouse it has long been assumed that there are two distinct phases of endoderm formation such that extra-embryonic endoderm forms prior to gastrulation and is derived from the primitive endoderm (Fig. 1G, red), while the definitive endoderm rises from the epiblast during gastrulation. Prior to gastrulation, the original primitive endoderm expands with the growing embryo and becomes subdivided into PE (Fig. 1G, yellow) and VE (Fig. 1G, green) based on their position relative to the egg cylinder. The VE itself is further divided into sub-regions, including the anterior visceral endoderm (AVE) (Fig. 1G, blue). In this study, we characterize and compare three cell lines that are either derived from the primitive endoderm or have been reported to resemble these primitive endoderm-derived lineages. Three ExEn cells lines were examined in detail by immunocytochemistry, qRT-PCR and microarray analysis, using well-characterized markers for ExEn and definitive endoderm, and a more elaborate panel of putative ExEn markers, previously identified as distinguishing between different subtypes of endoderm. These studies confirm that each of these ExEn cell lines exhibits high molecular correlation to visceral and parietal endoderm and little or no similarity to definitive (epiblast-derived) endoderm. This comparative gene analysis also refines a growing list of markers that have been proposed to distinguish between VE and DE. By providing a clearer picture of endodermal subtypes, these studies should assist the development of experimental protocols that require a distinction between embryonic and extra-embryonic lineages.

Finally, consistent with our previous embryological studies, pathway analysis from microarray data reveals that molecules downstream of TGFbeta-family members are highly represented.

Figure 1. Morphological characterization of END2, PYS2 and XEN cells. DIC (1 A–C) and Scanning EM (1 D–F) images of END2 (1 A, D), PYS2 (1 B, E) and XEN (1 C, F) cells reveal morphological details of the cell lines used in these studies. G. Cartoon depicting early endodermal lineages in the mouse embryo prior to gastrulation. The primitive endoderm (red) forms in the pre-implantation blastocyst stage embryo and subsequently expands and differentiates into parietal endoderm (yellow), visceral endoderm (green) and anterior visceral endoderm (blue). Visceral endoderm is also sub-divided into embryonic and extra-embryonic regions based both on location relative to the embryonic/extra-embryonic junction of the epiblast (grey), fate and marker expression.

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in these cell lines and suggests that both SMAD-dependent and SMAD-independent TGFβ signaling could mediate the inductive function of these cell lines.

Results

Extra-embryonic endoderm stem cells (XEN cells) and PYS2 cells but not END2, express markers characteristic of the primitive endoderm

END2 and PYS2 cells have been described previously, based on cell morphology and marker expression [1,4,5], to be similar to visceral endoderm (VE) and parietal endoderm (PE), respectively. Because these cells were originally derived from EC cell lines, they may not represent true endodermal lineages but rather, endoderm-like populations. Recently described protocols allow for the isolation of ExEn stem cells (XEN cells) directly from blastocyst stage mouse embryos [6], and as such, are more likely to represent endogenous endodermal cell types. For this study, we derived a XEN cell line from wild type mouse blastocysts of the ICR strain. Although each of the three cell lines are relatively flat and exhibit a cobblestone appearance when confluent, bright field microscopy reveals that each of the three cell lines is morphologically distinct from the other two (Fig. 1A–C). Scanning electron microscopy of cells plated at low density reveals that all three cell types are rounded in appearance and densely covered with microvilli, with END2 and PYS2 cells forming large lamellipodia (Fig. 1D–F). These data demonstrate that, like END2 and PYS2 cells, XEN cells exhibit an endodermal morphology.

Each of these cell lines was assessed by immunocytochemistry, for a panel of markers characteristic of the primitive endoderm including SOX7, GATA4 and GATA6 [7,8,9,10]. Both XEN and PYS2 cells are recognized by antibodies against GATA4, GATA6, and SOX7 (Fig. 2 B, C, E, F, H, I). However, only a subset of XEN cells express SOX7 (Fig. 2I). By contrast, END2 cells express only low levels of GATA6 (Fig. 2D) and do not express GATA4 or SOX7 (Fig. 2 A, G). In these studies, PYS2 cells showed the most uniform expression of these primitive endoderm markers. By contrast, END2 and XEN cells may represent mixed or fluctuating populations of primitive endoderm and other lineages since their expression of these markers was more heterogeneous. In particular, the failure of END2 cells to express GATA4 and SOX7, suggests that there are few if any primitive endoderm cells within this line. While they do exhibit heterogeneous expression of GATA6 it should be noted that this gene also marks other endodermal subtypes including the VE. The heterogeneity of END2 cells is also demonstrated by the expression of BMP2, which is proposed to be a major signaling molecule from the endoderm [11,12,13,14,15,16,17]. BMP2 is uniformly expressed in PYS2 and XEN cells (Fig. 2L, M) but only expressed in a small subset of END2 cells (Fig. 2J). Overall these data suggest that END2 cells represent a heterogeneous endodermal population with little resemblance to the primitive endoderm.

Detailed marker analysis demonstrates that END2 cells are molecularly divergent from XEN and PYS2 cells

To further characterize these cell lines, we used qRT-PCR (Fig. 3A), to examine a panel of markers representing several ExEn lineages (Fig. 1G and Fig. 3, insert) including PrE, PE, VE and anterior visceral endoderm (AVE). Sox7 [7], Pdgfra [18], Gata4 [10], and Gata6 [19] are expressed in the primitive endoderm of the mouse blastocyst and are thought to be among the earliest ExEn markers. XEN and PYS2 cells express all of these markers,
but END2 cells express only Gata6. Note that all of these markers are also expressed in derivatives of the primitive endoderm including the PE and VE and as a consequence, there are no known markers that are uniquely expressed in the primitive endoderm. All three cell lines express markers for the PE including t-type Plasminogen activator (tPA) [20], Cytokeratin 19 (Krt19) [21], Laminin B1 (Lamb1) [22] and SPARC [23], although END2 cells express these at relatively lower levels as compared to the other cell lines. 

We next determined the transcriptional status of markers that are widely expressed throughout the VE including FoxA2 [24], Transferrin (Tr) [25] u-type Plasminogen activator (uPA) [26] and Hepatocyte nuclear factor 4 (HNF4a) [24]. Tr and uPA are expressed by all three of these cell lines (although at lower levels in END2). HNF4a is expressed by PYS2 and XEN cells, but not by END2. FoxA2 is only expressed by PYS2 cells. None of these cell lines express Vilin (data not shown).

We next assessed markers whose expression is restricted (either spatially or temporally) within the VE. The extra-embryonic VE that lies proximally over the extra-embryonic ectoderm of the mouse embryo (Fig. 1G) expresses Sox7, Sox17 and also upregulates alpha fetoprotein (Afp) after gastrulation has been initiated. Prior to gastrulation, Afp and Sox17 also mark the distally positioned VE that overlies the epiblast (the embryonic VE) (Fig. 1G and 3, insert) [27]. Of these, Sox7 and Sox17 are present in both PYS2 and XEN but not END2 cells. Afp is only expressed in PYS2 cells. Thus, PYS2 and XEN cells express markers for both the extra-embryonic VE and the embryonic VE, whereas END2 cells only expressed panVE markers such as Ttr.

Finally, we assessed a panel of markers that are spatially restricted in the VE. Dkk-1 [28], Cerl [29,30,31], and Hex [32] are all known to be expressed in the AVE of the mouse embryo. As has previously been shown for XEN cells [6], all three ExEn cell lines express Hex. PYS2 and XEN cells also express Dkk1. None of the

Figure 3. Heart inducing cell lines express markers characteristic of several primitive endoderm lineages. A. Summary of Real-Time PCR on END2, PYS2 and XEN cells. Insert, cartoon showing embryonic lineages assessed, primitive endoderm (reds), parietal endoderm (oranges/yellows), visceral endoderm (greens) and AVE (blues). The panel of markers assessed include markers for primitive endoderm (Sox7, Pdgfra, Gata4, Gata6), parietal endoderm (tPA, Krt19, Lamb1 and SPARC), visceral endoderm, (FoxA2, Tr, uPA and HNF4a), anterior visceral endoderm (Dkk1, Cerl, Hex), the regionally restricted VE marker Bmp2 and the definitive/pan endoderm marker Sox17. B. Linear regression analysis comparing real-time PCR data to averaged fluorescence detection in the Illumina Microarray. 80% of markers that we compared showed strong correlation between the qRT-PCR data and microarray detection. The data discussed in this publication have been deposited in NCBI’s Gene Expression Omnibus [78] and are accessible through GEO Series accession number GSE19564 (http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE1956). doi:10.1371/journal.pone.0012016.g003
cell lines express Cerl (data not shown). Consistent with our immunocytochemical analysis, BMP2, which is also spatially restricted within the VE, is expressed by all three of the ExEn cell lines.

Both the patchy expression of these markers when assessed by immunocytochemistry and the relatively lower expression of mRNAs for these genes when assessed by qRT-PCR, are consistent with the idea that END2 cells are a heterogeneous population in which a small subset of cells express markers for the VE (or more likely, a subtype of VE), whereas PYS2 and XEN cells are more homogeneous and express a broad array of markers for the primitive endoderm and its derivatives.

To further analyze these cell lines, we performed a comparative microarray analysis. To confirm the consistency between the array data and data collected from qRT-PCR and immunocytochemistry analyses, we examined the same panel of markers initially assessed by qRT-PCR. Averaged fluorescent detection for each marker was plotted versus qRT-PCR data normalized to Gapdh and R\(^2\) values were determined from the line of best-fit. Importantly, a high degree of correlation is found between the two data sets, and over 80% of the genes tested had R\(^2\) values close to one (Fig. 3B). We found only three notable exceptions. First, \(\beta 2\) showed the same basic trend between qRT-PCR and microarray, but with low numerical correlation. This could reflect non-specific amplification by qRT-PCR or a problem with the array probe. In addition, a single probe for \(Gata4\) is highly recognized in the array by END2 cells. \(Gata4\), however, is absent in END2 cells by both qRT-PCR (Fig. 3A) and immunocytochemistry (Fig. 1A). Finally, \(Bmp2\) is not detected in the array but is highly expressed by PYS2 and XEN cell lines, as assessed by qRT-PCR and immunocytochemistry (Fig. 2 J–M and Fig. 3A). This suggests that these particular probes may either recognize non-specific transcripts or splice variants of the target genes and highlights the need for independent verification of candidates identified by probing microarrays. Overall, there is significant agreement between microarray data and our other assays.

Having confirmed a high degree of correlation between qRT-PCR data and the array, a large-scale analysis of markers for endodermal cell types was undertaken using microarray data (Fig. 4). Studies using immunohistochemistry and in situ hybridization studies have identified a relatively small number of genes that are differentially expressed in the endoderm. Indeed, many of these markers are expressed in more than one endodermal subtype. For example, the primitive endoderm markers used in this study (\(Gata4\), \(Gata6\), \(Sox7\) and \(Pdgfr\)) are also expressed in other endodermal subtypes. Thus few truly diagnostic markers that distinguish endodermal subtypes have been identified. Two recent studies aimed at identifying markers that distinguish between VE and DE were based largely on comparative analysis of endodermal cell populations sorted by expression of endodermal markers. First, Sherwood et al. used differential expression of the epitopes for EpCAM, Dha and Sox4-\(\beta\) [2]. Later, Yasunga et al. used differential expression of Goosecoid and Sox17 [3]. Here we sought to analyze these data sets by comparative analysis of the three ExEn cell lines. First, we analyzed our array data based on well-characterized markers for endodermal subsets (Fig. 4). As expected all three of these cell lines express most of the VE specific markers. Notably, all three express high levels of \(Foxd3\), \(Em2p\), \(Sdc4\) and \(Gata6\), suggesting that these markers, in particular, are highly diagnostic for the VE fate. By contrast, a second subset of VE markers including \(Tb\), \(Dab2\), \(Pph1\) and \(Cldn4\) are highly expressed by PYS2 and XEN cells but not by END2 cells, suggesting that they mark a specific subset of cells within the VE. In addition, all three cell lines express the AVE markers \(Hex\) and \(Dkk1\) but not \(Cerl\).

None of the cell lines express markers reported to be diagnostic for DE including \(Gata4\), \(Gp1\) and \(Tmie2\) [3]. This would seem to confirm that these cell lines do not possess characteristics of the DE and conversely support the notion that these markers are diagnostic for DE but not ExEn cell types.

We further analyzed these cells for PE markers and found, as expected based on our previous analysis, that PYS2 and XEN but not END2 cells show high expression of PE markers. Finally, in confirmation of our previous findings, XEN cells and PYS2 cells but not END2 cells express high levels of markers for the primitive endoderm. In addition, each cell had a specific subset of uniquely expressed markers that are either higher or lower as compared to the other two cell lines. Since these cell lines have in other assays been shown to have inductive effects, such as activating heart formation [4,5,33,34,35], these differences might be exploited to identify specific inductive signals within the individual cell lines.

The microarray data were then examined for expression of a large panel of markers identified in Sherwood et al. that are described as distinguishing between DE and VE (Fig. 5). Many, but not all, of the pan-endodermal markers are expressed by these cell lines [9/18 (50%), END2, PYS2 and 7/18 (39%), XEN cells]. These data indicate that a subset of these markers are not truly pan endodermal but add further support to the characterization of \(Sox17\), \(Spns3\) \(Rab\) 15, \(Dg2\), \(Rbp4\), \(Axs4\) and \(Emb\) as true pan endodermal markers. In addition, markers that were described as VE-enriched are highly expressed in the ExEn cell lines (65%, END2, 60% PYS2 and 62.5% XEN). By contrast, ExEn cells express only a small percentage of markers that distinguish DE from VE (9.6% END2, PYS2, 19% XEN cells). It should be noted, however that these markers are not exclusive to the DE and VE, and thus their expression in ExEn cell lines may indicate the presence of other lineages such as parietal endoderm.

Altogether these findings suggest that all of the cell lines examined in these studies are similar to extra-embryonic endodermal lineages and probably represent mixed populations of PE derivatives. In addition, microarray analysis reveals distinct differences between these cell lines. To further understand these differences, cluster analysis (Fig. 6A) was performed. This supported the existence of significant molecular differences between the three ExEn cell lines. The greatest overall differences were found when END2 cells were compared to the other two cell lines (PYS2 and XEN). This finding is consistent with our previous analysis. We also compared the number of genes that are differentially expressed in pair-wise comparisons of the three cell lines, confirming our findings from the cluster analysis (Fig. 6B).

Therefore, while demonstrating that the three cell lines exhibit characteristics of the VE, these studies also highlight the fact that they do, nonetheless, have significant molecular differences between them. Since a subpopulation of the VE, the AVE has been shown to have heart-inducing ability, it is possible that these molecular differences might also reflect differences in the ability of these cell lines to activate and/or enhance cardiac differentiation in ES cells. Indeed both END2 and PYS2 cells have already been shown to possess heart inducing ability [4,5,33,34,35].

Microarray analysis
To examine the microarray data in more detail, we performed pathway and tissue expression analysis on a total of 6094 annotated gene IDs that were detected as present in at least one of the three cell lines (based on a p-value of detection <0.01) using the Database for Annotation, Visualization and Integrated Discovery (DAVID) [36,37]. Tissue expression analysis revealed that the top non-cancer tissue hit for this list of gene IDs was for
liver (with a \( p \)-value of 5.3E-115), which is expected given the high degree of overlap between markers for the liver and the VE.

To determine the signaling pathways that characterize these cell lines, the DAVID bioinformatics tool was used to compare the 6094 gene IDs present in the arrays to the BIOCARTA pathways database. This analysis revealed that the top pathways (not directly related to cell cycle) in the ExEn lines were the MAP Kinase (\( p \)-value, 2.7E-3) and TGFbeta (\( p \)-value, 3.1E-3) signaling pathways. Gene expression in these pathways was described by comparing our gene list to the TGFbeta and MAP Kinase pathways described in the Kyoto Encyclopedia for Genes and Genomes (Fig. 7 and Fig. 8). A detailed analysis of the TGFbeta pathway suggests that all three of these cell lines are capable of responding to all known subgroups of TGFbeta family members (Fig. 7). By comparison, an analysis of known MAP Kinase signaling pathways suggests that only XEN cells have a fully intact classical MAP Kinase signaling pathway since it was the only cell line in which \( \text{Grb2} \) is present by microarray (Fig. 8A). By contrast microarray data suggests that

Figure 4. Heat map analysis of well-characterized markers for different endodermal cell types. Illumina microarray data for genes that are expressed in various endoderm subtypes (depicted as heat maps) includes regionally restricted markers representing the VE, DE, PE and PrE. Each of the three cell lines expresses markers for VE, PE and PrE. None of the cell lines express markers that are diagnostic for DE. Fluorescence data was indicated as 0.00 if the \( p \)-value of detection was greater than 0.01.

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Figure 5. Heat map representation of detailed marker analysis. Microarray data of markers distinguishing VE from DE according to Sherwood et al. [2]. Only a subset of the previously examined pan endodermal markers is expressed in the array. This refines the list of true pan endodermal markers to include Sox17, Spink3, Rab15, Dsg2, Ripk4, AnxA4 and Emb. "VE enriched" genes indicate factors that were found to be expressed in VE plus other lineages but not DE. Our analysis suggests that a subset of these factors may not be present in all VE subtypes and thus may represent regionalized VE markers. "DE enriched" represents genes found, in Sherwood et al. [2], to be expressed in DE and other subtypes but not in VE. These data strongly suggest that the ExEn cell lines are not similar to DE. Fluorescence data was marked as 0.00 if the p-value of detection was greater than 0.01.

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### Pan Endodermal

| Gene   | END2 | PYS2 | XEN |
|--------|------|------|-----|
| Bnpi1  | 0.00 | 0.00 | 0.00 |
| Cacna1b| 0.00 | 0.00 | 0.00 |
| Cldn8  | 0.00 | 0.00 | 0.00 |
| Clic6  | 0.00 | 0.00 | 0.00 |
| Npnt  | 0.00 | 0.00 | 0.00 |
| Rbm35a | 0.00 | 0.00 | 0.00 |
| Sh3gl2 | 0.00 | 0.00 | 0.00 |
| Sft14  | 0.00 | 0.00 | 0.00 |
| Tmem30b| 0.00 | 0.00 | 0.00 |
| Tnpr5s2| 0.00 | 0.00 | 0.00 |
| Sox17  | 0.96 | 11.74 | 11.13 |
| Spink3 | 7.11 | 12.44 | 12.08 |
| Rab15  | 7.15 | 10.77 | 10.45 |
| Foxa1  | 7.43 | 6.95  | 0.00 |
| Dsg2   | 7.82 | 7.29  | 7.33 |
| Ripk4  | 8.27 | 9.90  | 10.28 |
| AnxA4  | 9.05 | 9.41  | 7.85 |
| Emb    | 9.31 | 12.27 | 10.58 |

### DE enriched

| Gene   | END2 | PYS2 | XEN |
|--------|------|------|-----|
| Amt2   | 0.00 | 0.00 | 0.00 |
| Brachyury | 0.00 | 0.00 | 0.00 |
| Cdx4   | 0.00 | 0.00 | 0.00 |
| Cdt2   | 0.00 | 0.00 | 0.00 |
| Dlx2   | 0.00 | 0.00 | 0.00 |
| Dlx3   | 0.00 | 0.00 | 0.00 |
| Dlx5   | 0.00 | 0.00 | 0.00 |
| Dmrt1  | 0.00 | 0.00 | 0.00 |
| Evx1   | 0.00 | 0.00 | 0.00 |
| Sox3   | 0.00 | 0.00 | 0.00 |
| Foxg1  | 0.00 | 0.00 | 0.00 |
| Gli3   | 0.00 | 0.00 | 0.00 |
| Hey2   | 0.00 | 0.00 | 0.00 |
| Hoxa3  | 0.00 | 0.00 | 0.00 |
| Hoxa9  | 0.00 | 0.00 | 0.00 |
| Hoxb1  | 0.00 | 0.00 | 0.00 |
| Hoxb2  | 0.00 | 0.00 | 0.00 |
| Hoxb3  | 0.00 | 0.00 | 0.00 |
| Hoxc4  | 0.00 | 0.00 | 0.00 |
| Hoxd1  | 0.00 | 0.00 | 0.00 |
| Hoxd8  | 0.00 | 0.00 | 0.00 |
| Idb4   | 0.00 | 0.00 | 0.00 |
| Ix2    | 0.00 | 0.00 | 0.00 |
| Ix3    | 0.00 | 0.00 | 0.00 |
| Ix5    | 0.00 | 0.00 | 0.00 |
| Is1    | 0.00 | 0.00 | 0.00 |
| Mesp1  | 0.00 | 0.00 | 0.00 |
| Nkx1-2 | 0.00 | 0.00 | 0.00 |
| Pax1   | 0.00 | 0.00 | 0.00 |
| Pax3   | 0.00 | 0.00 | 0.00 |
| Pax6   | 0.00 | 0.00 | 0.00 |
| Pax8   | 0.00 | 0.00 | 0.00 |
| Pax9   | 0.00 | 0.00 | 0.00 |
| Pknox2 | 0.00 | 0.00 | 0.00 |
| Rhi3   | 0.00 | 0.00 | 0.00 |
| Sif1   | 0.00 | 0.00 | 0.00 |
| sidx3  | 0.00 | 0.00 | 0.00 |
| Sox11  | 0.00 | 0.00 | 0.00 |
| Sox9   | 0.00 | 0.00 | 0.00 |
| Sp6    | 0.00 | 0.00 | 0.00 |
| Sstb2  | 0.00 | 0.00 | 0.00 |
| Tcfap2a| 0.00 | 0.00 | 0.00 |
| Tcfap2b| 0.00 | 0.00 | 0.00 |
| Tpf63  | 0.00 | 0.00 | 0.00 |
| Zfhx2  | 0.00 | 0.00 | 0.00 |
| Pbx1   | 0.00 | 0.00 | 0.00 |
| Hoxa1  | 0.00 | 0.00 | 0.00 |
| Hoxa3  | 0.00 | 0.00 | 0.00 |
| Mrg1   | 0.00 | 0.00 | 0.00 |
| 8.88   | 7.11  | 7.92  |

| Normalized detection in array |
|--------------------------------|
| absent                         |
| 6.0-6.99                      |
| 7.0-7.99                      |
| 8.0-8.99                      |
| 9.0-9.99                      |
| 10.0-10.99                    |
| 11.0-11.99                    |
| 12.0-13.5                     |
END2 and PYS2 cells likely signal through the TAK1/p38/JNK and the TAK1/NLK pathways. Although we found a better than 80% correlation between microarray and qRT-PCR data, we decided to confirm the presence or absence of Grb2 expression in these cell lines by qRT-PCR (Fig. 8B). By PCR, we found that each of the three cell lines expresses mRNA for Grb2 at similar levels. Taken together these analyses suggest that each of the three endodermal cell lines can signal through both classical and non-classical MAP Kinase signaling pathways.

In addition, we have previously shown [40] that signaling of TGFβa family members in the endoderm is required for heart development. Together these pathway analyses (Fig. 7, 8) reveal that the effects of TGFβa signaling could be mediated by either traditional SMAD-dependent signaling or by a SMAD independent pathway involving either TAK1/p38/JNK or TAK1/NLK.

Finally, since END2 and PYS2 cells have been shown to secrete factors that induce differentiation in ES cells it is likely that XEN cells will have a similar inductive ability. Before this can be established, however, it is important to show that XEN cells are stable when maintained in culture. To address this question, we collected XEN cells at 70% confluence to ensure that cell density is equivalent for all analyses over seven passages. We then assessed the expression of a number of ExEn markers at each passage. In parallel, XEN cells were passaged in a serum-free medium to determine if gene expression in these cells is sensitive to culture conditions (Fig. 9). We found that despite spikes in some markers at certain passages, that cells grown in standard serum-containing medium are quite stable over seven passages and that there were no obvious trends in marker expression during the duration of this experiment. This suggests that XEN cells are stable in standard medium for at least short-term culture. When XEN cells are grown in serum-free medium we again noticed spikes in expression of some markers at some passages but again found no obvious trends in marker expression. We did however note that overall expression of Gata6 and Tbr was different between the two culture conditions whereas other markers were expressed at statistically the same levels at most passages.

Discussion

The endoderm makes up only a small percentage of cells of the early embryo. Cells within the endoderm layers of the early embryo exist in simple cuboidal or squamous epithelia, making them difficult to isolate mechanically from embryos. Because of these challenges to the embryological study of endodermal cells in the mouse embryo, many recent studies have relied on the use of endodermal-like cell lines that can mimic the functions of the early endoderm. For example END2 cells have been shown to enhance myocardial differentiation of both human and mouse ES cells [4,33,34] and to mimic the effect of the VE in activating cardiac formation from the undifferentiated mesoderm from the mouse embryo [35]. While it is clear from these studies that END2 cells mimic the effects of the AVE, the fact that they were originally derived from ECs raises some doubt as to whether they perfectly recapitulate the endogenous signals secreted by the AVE.

An added complication is the significant overlap between genes that mark the ExEn and those that mark the DE. This makes the analysis of in vitro differentiation of endodermal cell types difficult and thus poses a major hurdle for attempts to derive endodermal cell types for ES or other sources that might be used for therapeutic purposes.

Our studies address both of these concerns. First, we have undertaken an in depth analysis of the newly characterized XEN stem cells [6] which like the EC-derived END2 and PYS2 cells, express markers for the AVE. XEN cells can thus be used to study the inductive effects that have been attributed to the AVE, including heart formation, primitive streak initiation and forebrain induction. In addition, since XEN cells are derived from mouse blastocysts, it will be possible to derive XEN cells from mice with deletions of genes thought to be involved in AVE function thereby

END2 and PYS2 cells likely signal through the TAK1/p38/JNK and the TAK1/NLK pathways. Although we found a better than 80% correlation between microarray and qRT-PCR data, we decided to confirm the presence or absence of Grb2 expression in these cell lines by qRT-PCR (Fig. 8B). By PCR, we found that each of the three cell lines expresses mRNA for Grb2 at similar levels. Taken together these analyses suggest that each of the three endodermal cell lines can signal through both classical and non-classical MAP Kinase signaling pathways.

In addition, we have previously shown [40] that signaling of TGFβa family members in the endoderm is required for heart development. Together these pathway analyses (Fig. 7, 8) reveal that the effects of TGFβa signaling could be mediated by either traditional SMAD-dependent signaling or by a SMAD independent pathway involving either TAK1/p38/JNK or TAK1/NLK.

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Figure 6. Cluster analysis of microarray data. A. Cluster Dendrogram representing the amount of variance in markers commonly expressed through microarray analysis by each of the ExEn cell lines (in triplicates). B. Venn Diagrams representing the number of genes in the array that are either upregulated (red) or downregulated (blue) in pair-wise comparisons between the different cell lines. Probes are called as “present” if the p-value for detection was less than 0.01.
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providing an assay to directly test their function in the primitive endoderm. Finally, by comparing the three related but dissimilar cell lines, we are able to refine the list of markers that distinguish between different subtypes of endoderm including those that distinguish DE from VE.

**Inductive and morphogenetic functions of the extra-embryonic endoderm**

Endoderm both embryonic and extra-embryonic has been shown in various model systems to play an inductive role in the differentiation of mesodermal and ectodermal tissues that adjoin it. The VE in particular has been proposed to play important roles in heart formation, primitive streak formation and the development of the forebrain.

Studies in amphibian embryos dating back to the 1960s demonstrate that signals from the pregastrula endoderm support myocardial differentiation [38,39,40,41,42,43,44,45]. This finding is supported by studies that the AVE of the mouse [46] and the hypoblast of avian embryos showing that rotation of the hypoblast (which is equivalent to the AVE [55]) results in a repositioning of the primitive streak [55,56,57]. A more in depth molecular analysis of this process has revealed greater complexity than was previously anticipated. First, removal of the hypoblast does not eliminate streak formation but rather results in the formation of multiple streaks. This suggests that the hypoblast does not activate streak elongation but rather acts to limit streak formation to a single location [58]. Simultaneously, FGF signaling co-cultured with mouse ES cells [50]. Importantly, these studies indicate that cardiac specification requires a VE signal only transiently, from early to mid-gastrulation [47,51,52] and that signals from the DE are required only later, for proliferation of cardiomyocytes and the initiation of beating [53]. Thus it will be of particular importance to separate the molecular signals of the DE from the VE in order to uncover the specific signals that mediate endoderm’s ability to activate and later support myocardial differentiation.

The VE has also been implicated as providing signals required for streak elongation. This hypothesis has come largely from the observation of an anterior migration of the AVE just prior to the onset of gastrulation [54] and studies in avian embryos showing that rotation of the hypoblast (which is equivalent to the AVE [55]) results in a repositioning of the primitive streak [55,56,57]. A more in depth molecular analysis of this process has revealed greater complexity than was previously anticipated. First, removal of the hypoblast does not eliminate streak formation but rather results in the formation of multiple streaks. This suggests that the hypoblast does not activate streak elongation but rather acts to limit streak formation to a single location [58]. Simultaneously, FGF signaling

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**Figure 7. TGFbeta signaling pathways are active in heart inducing cell lines.** Diagram of TGFbeta signaling pathways from the Kyoto Encyclopedia of Genes and Genomes [75,76,77] showing pathway components that are considered to be present based on a p-value of detection <0.01. Factors indicated in blue are present in all of the ExEn cell lines. Factors present in green are present in a subset of the cell lines (the particular cell lines are indicated by letters adjacent to the box indicating the factor). doi:10.1371/journal.pone.0012016.g007
from the hypoblast activates localized expression of genes associated with the establishment of planar cell polarity and defines the site of active streak elongation [59].

Finally, two specific observations lead to the hypothesis that AVE might serve as an inducing population for the vertebrate forebrain. First, ablation of the AVE of the mouse leads to a loss of Hesx expression in the anterior neural folds of the mouse (although other markers assessed were normal) [60]. Second, chimeric analysis of mouse embryos possessing homozygous deletion of Otx2 [61,62] and Lim1 [63] or that are double mutant for Lim1 and HNF3B [64], suggest that AVE expression of these genes is necessary for normal axis and forebrain development. Together these data suggest that signaling from the AVE is necessary for normal forebrain development. However, the AVE does not directly activate forebrain differentiation. Grafting of the rabbit AVE [65] or chick hypoblast to naïve epiblast that is capable of forming neural tissue results in the transient ectopic expression of neural markers but these markers are not maintained and host tissues do not form neural plate structures [55]. Similarly, explant co-cultures of AVE and mouse epiblast do not induce the expression of anterior neural markers but instead suppress posterior neural differentiation [66]. So what is the mechanism by which VE helps to pattern the forebrain if it does not act as a forebrain inducer? 1) Transient activation of early neural markers suggests that the VE may prime the ectoderm for neural development, 2) The AVE appears to repress posterior development and 3) it appears that the VE directs morphogenetic movements in the ectoderm and mesoderm that are required for normal axis formation [reviewed in: [55,66]. This hypothesis is consistent with the hypoblast rotation experiments described in the previous section.

Together, these studies highlight the importance of the VE generally and the AVE specifically in the early patterning of the embryo and subsequent organ formation.

Does the extra-embryonic endoderm make cellular contributions to endodermal organ formation?

Until recently, the dogma of endoderm formation in the mouse has asserted that the visceral endoderm that surrounds the embryonic epiblast prior to gastrulation is actively displaced by the forming definitive endoderm and contributes only to extra-
embryonic structures [67]. This conclusion was based largely on the observation of gene expression patterns showing markers such as alpha-fetoprotein being lost in the endodermal cells overlying the epiblast during gastrulation and low-resolution fate mapping studies. With the advent of multiple lineage labels and time-lapse live imaging, we have clearly demonstrated that all or most the visceral endoderm that overlies the pre-gastrula epiblast is integrated into the definitive endoderm rather than being displaced by it. Concomitantly, these cells lose expression of visceral endoderm markers and gain expression of markers for the definitive endoderm. This lineage study also demonstrates that VE-derived cells contribute to gut formation [27]. This shows that there are distinct regions of embryonic (EmVE) and extra-embryonic (ExVE) visceral endoderm with unique fates in the embryo and suggests that these EmVE cells may contribute to endodermal organ formation. While none of the ExEn cell lines that we assessed expressed markers for the definitive endoderm, it remains to be determined if these cells can be coaxed by the addition of growth factors to differentiate along definitive endoderm lineages and adopt fates associated with the gut or its associated organs. If this turns out to be the case, then in addition to being a useful tool for the study of the inductive properties of the

Figure 9. Marker expression in XEN cells is stable over several passages when cells are grown under standard serum-containing medium. qRT-PCR data comparing marker expression in XEN cells grown under standard serum-containing conditions to XEN cells grown in serum-free medium. Despite some spikes in marker expression at some passages, over the entire course of this experiment there were no obvious trends (either upward or downward changes in expression) in any of the markers assessed. In addition, while some markers were differentially expressed between the two culture conditions, there was also no obvious change in marker expression in the serum-free medium when assessed over the entire course of the experiment. This suggests that XEN cells are stable when grown in culture over several passages.

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ExEn. XEN cells might also serve a stem population from which to derive differentiated endodermal cell types.

TAK1/p38/JNK/NIK pathways in these cell types

Above, we presented genetic and embryological data that the VE acts, in part, to direct cell movements in the embryo. Consistent with this, we found that each of the three ExEn cell lines possesses intact pathways for non-classical MAP Kinase signaling acting through JNK [68] and NLK [69]. Since both of these factors are associated with planar cell polarity, it supports the notion that these pathways may play a pivotal role in the endoderm's ability to direct morphogenetic movements. Indeed, JNK Kinase has been shown to be necessary for heart induction downstream of the non-canonical Wnt, Wnt11 [70]. In addition, TAK1, which is immediately upstream of NLK is essential for cardiac differentiation in P19 [71], and TAK1 [72] and TAB1[73] mutants show defects in cardiac morphogenesis. It remains to be determined whether these defects result from a specific endodermal requirement for these signals or arise from a more general requirement in embryonic tissues. Nonetheless these findings lend support to a model in which a non-classical MAP Kinase pathway mediated by TAK1/p38/JNK or TAK1/NIK mediate some functions of the endoderm.

Conclusions

Endoderm, both embryonic and extra-embryonic plays important morphogenetic functions in the mouse and other vertebrate embryos. We are only just beginning to refine our understanding of the different types of endoderm and the molecular mechanisms by which they mediate their functions in development. Here, we undertake a thorough analysis of ExEn cell lines to help refine the list of markers that define various endodermal cell types. Specifically we provide further characterization of XEN cells [6] which may serve as a useful tool in the study of ExEn differentiation and function.

Materials and Methods

Cell Culture

XEN cells were derived from ICR strain blastocyst stage embryos according to standard procedures [6]. END2 cells were derived from P19 embryonal carcinoma cell lines [74] and PYS2 cells were derived from 129 strain mice tumor cells [1]. XEN and END2 cells were maintained in high glucose Dulbecco’s Modified Eagles Medium (DMEM) devoid of L-glutamine and sodium pyruvate (Mediatech). DMEM was enriched with 10% ES serum containing and serum-free conditions. PYS2 cells were both maintained in high glucose Dulbecco’s Modified Eagles Medium (DMEM) devoid of L-glutamine and sodium pyruvate (Mediatech). XEN cells were passaged on to gelatin-coated plastic coverslips. A secondary antibody was added and cells were incubated overnight at 4°C. Finally, cells were washed with PBS and cover slipped with Vectashield mounting medium containing DAPI.

Immunohistochemistry

Cells were washed with PBS and fixed in 4% PFA 30 minutes, then blocked with 3% FBS-0.3% Triton in PBS. Primary antibodies were then added, and incubated overnight at 4°C. Cells were washed with PBS and blocked for 30 minutes at room temperature. Secondary antibodies were added and cells were incubated overnight at 4°C. Finally, cells were washed with PBS and cover slipped with Vectashield mounting medium containing DAPI.

Scanning Electron Microscopy

Cells were passaged on to gelatin-coated plastic coverslips. A day later, they were rinsed once with PBS and fixed at room temperature in 2.5% Glutaraldehyde/2% PFA in 0.075M Sodium cacodylate buffer.
Cacodylate buffer pH 7.5 for one hour. They were then dehydrated in a graded ethanol series. Cells were then critical-point dried in a Denton JCP-1 Critical Point Drying Apparatus and subsequently coated with gold/palladium in a Denton Vacuum Desk 1V sputter coating system. Imaging was carried out with a Zeiss Field Emission Supra 25 Scanning Electron microscope.

Microarray Analysis

Total RNA was isolated with QIagen RNeasy Mini Kit and used to probe Illumina expression arrays (MouseWG-6_V2_0_R0_11278593 in triplicate for each of three heart-inducing cell lines using Illumina BeadStudio version 3.4.0. The raw Illumina data (9 arrays) was analyzed using Bioconductor packages. The data was first normalized using LumiExpresso (function. The differentially expressed genes in each pairwise comparison were obtained using Limma (function. For gene ontology studies, Illumina probes were mapped to gene symbol names using getAnnotate.illumina ("MouseWG-6_V2_0_R0_11278593_AhHz") downloaded from Bioconductor website: http://www.bioconductor.org/download.

Pathway and expression analysis was carried out using DAVID Bioinformatics Resources 2008 sponsored by the National Institute of Allergy and Infectious Diseases (NIAID), NIH, at http://david.abcc.ncifcrf.gov/[36,37] and the Kyoto Encyclopedia of Genes and Genomes http://www.genome.jp/kegg/ [75,76,77]. This data is MIAME compliant and has been deposited in NCBI’s Gene Expression Omnibus [78]. All data is accessible through GEO Series accession number GSE19564 (http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE1956).

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Author Contributions

Conceived and designed the experiments: AKH ACF. Performed the experiments: KB SL JA MXD. Analyzed the data: KB RK AKH ACF. Contributed reagents/materials/analysis tools: KK. Wrote the paper: KB ACF ACF.

References

1. Lehman JM, Speers WC, Swartzendruber DE, Pierce GB (1974) Neoplastic differentiation: characteristics of cell lines derived from a murine teratocarcinoma. J Cell Physiol 84: 13–27.
2. Sherwood RL, Jitianu C, Cleaver O, Shaywitz DA, Lamenzo JO, et al. (2002) Prospective isolation and global gene expression analysis of definitive and visceral endoderm. Dev Biol 304: 541–555.
3. Yasunaga M, Tada S, Torikai-Nishikawa S, Nakano Y, Okada M, et al. (2005) Induction and monitoring of definitive and visceral endoderm differentiation of mouse E8.5 cells. Nat Biotechnol 23: 1542–1550.
4. Mummery CL, van Achterberg TA, van den Eijnden-van Raaij AJ, et al. (1991) Visceral-endoendoderm-like cell lines induce differentiation of murine P19 embryonic carcinoma cells. Differentiation 46: 51–60.
5. Stary M, Pasteiner W, Summer A, Hrdina A, Eger A, et al. (2005) Parietal endoderm secreted SPARC promotes early cardiomyogenesis in vitro. Exp Cell Res 310: 331–343.
6. Kunath T, Arnaud D, Uy GD, Okamoto I, Chureau C, et al. (2005) Imprinted X-inactivation in extra-embryonic endoderm cell lines from mouse blastocysts. Development 132: 1649–1661.
7. Kanaz-Azuma M, Kanai Y, Gao JM, Tajima Y, Taya C, et al. (2002) Depletion of definitive gut endoderm in Sox17-null mutant mice. Development 129: 2367–2379.
8. Koutouzakis M, Langeveld A, Patient R, Beddington R, Grosveld F (1999) The transthyretin gene is expressed in the extraembryonic endoderm of the mouse embryo. Development 126: 723–732.
9. Chazaud C, Yamanaka Y, Pawson T, Rossant J (2006) Early lineage segregation between epiblast and primitive endoderm in mouse blastocysts through the Gfb2/SPARC pathway. Dev Cell 10: 615–624.
10. Pasha B, Pilszok A, Frankenberg S, Artus J, Hadjantonakis AK (2008) Distinct sequential cell behaviours direct primitive endoderm formation in the mouse blastocyst. Development 135: 3081–3091.
11. Shi Y, Katsos S, Cai C, Eaves S (2000) BMP signaling is required for heart formation in vertebrates. Dev Biol 224: 226–237.
12. Barron M, Gao M, Lough J (2000) Requirement for BMP and FGF signaling during cardiogenic induction in non-precordial mesoderm is specific, transient, and cooperative. Dev Dyn 210: 383–393.
13. van Wijk R, Moorman AF, van den Hoff MJ (2007) Role of bone morphogenetic proteins in cardiac differentiation. Cardiovasc Res 74: 224–235.
14. Schlange T, Andree B, Arnold HH, Brand T (2000) BMP2 is required for early heart development during a distinct time period. Mech Dev 91: 239–270.
15. Beldir A, Zingman LV, Hodgson DM, Rauzier JM, Kane GC, et al. (2002) Stem cell differentiation requires a paracrine pathway in the heart. Fasebj J 16: 1558–1566.
16. Coucouvanis E, Martin GR (1999) BMP signaling plays a role in visceral endoderm differentiation and cavitation in the early mouse embryo. Development 126: 535–546.
17. Uchimura T, Komatsu Y, Tanaka M, McCann KL, Mishina Y (2009) Bmp2 and Bmp7 genetically interact to support multiple aspects of mouse development including functional heart development. Genesis 47: 374–384.
18. Mercola D (1992) Platelet-derived growth factor, transformation and antiendocrine models wanted. In: Erickson RP, Leant J, eds. Gene Regulation by Antisense RNA and DNA. New York: Raven Press. pp 329–353.
19. Chazaud C, Rossant J (2006) Disruption of early proximodistal patterning and AVE formation in Apc mutants. Development 133: 3357–3367.
20. Cheng L, Grabel LB (1997) The involvement of tissue-type plasminogen activator in parietal endoderm outgrowth. Exp Cell Res 230: 187–196.
21. Marotti IJ, Taylor A, Williams JG, Sage H, Hogan BL (1998) Evidence from molecular cloning that SPARC, a major product of mouse embryonic parietal endoderm, is related to an endothelial cell ‘culture shock’ glycoprotein of Mr 43,000. Endo J 5: 1463–1472.
22. Mason IJ, Taylor A, Williams JG, Sage H, Hogan BL (1998) Expression of transcription factor HNF-4 in the extraembryonic endoderm, gut, and nephrogenic tissue of the developing mouse embryo. Dev Biol 218: 383–393.
23. Mason IJ, Taylor A, Williams JG, Sage H, Hogan BL (1998) Expression of transcription factor HNF-4 in the extraembryonic endoderm, gut, and nephrogenic tissue of the developing mouse embryo. Dev Biol 218: 383–393.
24. Duncan SA, Manova K, Chen WS, Hoodless P, Weinstein DC, et al. (1994) Expression of transcription factor HNF-4 in the extraembryonic endoderm, gut, and nephrogenic tissue of the developing mouse embryo: HNF-4 is a marker for primary endoderm in the implanting blastocyst. Proc Natl Acad Sci U S A 91: 7508–7502.
25. Costa RH, Van Dyke TA, Yan C, Kuo F, Darnell JR, Jr. (1998) Similarities in transthyretin gene expression and differences in transcription factors: liver and yolk sac compared to choroid plexus. Proc Natl Acad Sci U S A 87: 6598–6593.
26. Marotti KR, Belin D, Strickland S (1982) The production of distinct forms of plasminogen activator by mouse embryonic cells. Dev Biol 90: 154–159.
27. Shavalk W, Deng JM, Behringer RR (1998) Expression of the mouse cerberus-related gene, Covt, suggests a role in anterior neural induction and somitogenesis. PNAS 85: 6198–6203.
28. Inman KE, Downs KM (2006) Localization of Brachyury (T) in embryonic and extraembryonic tissues during mouse gastrulation. Gene Expr Patterns 7: 783–793.
29. Shawlot W, Deng JM, Behringer RR (1998) Expression of the mouse cerberus-related gene, Covt, suggests a role in anterior neural induction and somitogenesis. PNAS 85: 6198–6203.
30. Inman KE, Downs KM (2006) Localization of Brachyury (T) in embryonic and extraembryonic tissues during mouse gastrulation. Gene Expr Patterns 7: 783–793.
31. Bilen C, Stanley E, Fabri L, Kotrecha S, Rhim M, et al. (1998) Murine cerberus homologue mCer-1: a candidate anterior patterning molecule. Dev Biol 194: 153–151.
32. Thomas PQ, Brown A, Bedington RS (1998) Hex: a homeobox gene revealing peri-implantation asymmetry in the mouse embryo and an early transient marker of endodermal cell precursors. Development 125: 85–94.
33. Mummery C, Ward-van Oostraad D, Doevendans P, Spijker R, van den Bos J, et al. (2003) Differentiation of human embryonic stem cells to cardiomyocytes: role of corticosteroids in perinatal heart disease and adult heart failure. Circulation 107: 2733–2740.
34. Passier R, Oostraad DW, Snapper J, Kluots J, Hassink RJ, et al. (2005) Increased cardiomyocyte differentiation from human embryonic stem cells in embryoid bodies with drug exposure. Stem Cells 23: 772–780.
35. Nijmeijer RM, Leeuwis JW, DeLisio A, Mummery CL, Chuva de Sousa Lopes SM (2009) Visceral endoderm induces specification of cardiomyocytes in mice. Stem Cell Res 3: 170–178.
Sater, A.K., Jacobson, A.G. (1989) The specification of heart mesoderm occurs during gastrulation in salamanders. Journal of Experimental Zoology 167: 79–103.

Rudy-Reil, D., Lough, J. (2004) Avian precardiac endoderm/mesoderm induces visceral embryonic endoderm and primitive streak for terminal differentiation. Development 131: 343–353.

Matsui, H., Sakabe, M., Sakata, H., Yanagawa, N., Ikeda, K., et al. (2008) Induction of cardiac differentiation occurs in the visceral embryonic endoderm and primitive streak by antagonizing nodal signaling. Dev Cell 3: 735–744.

Gannon, M., Bader, D. (1995) Initiation of cardiac differentiation occurs in the mouse inner cell mass by blastocyct injection. J Embryol Exp Morphol 129: 413–417.

Antin, P.B., Taylor, R.G., Yatskievych, T. (1994) Precardiac mesoderm is specified during gastrulation in quail. Developmental Dynamics 200: 144–154.

Matsui, H., Sakabe, M., Sakata, H., Yanagawa, N., Ikeda, K., et al. (2008) Induction of cardiac differentiation occurs in the visceral embryonic endoderm and primitive streak for terminal differentiation. Development 131: 343–353.

Arai, A., Yamamoto, K., Toyama, J. (1997) Murine cardiac progenitor cells require visceal embryonic endoderm and primitive streak for cardiac myocyte differentiation in murine embryonic stem cells. Circ Res 94: 2561–2570.

Voiculescu, O., Bercovich, F., Wolpert, L., Keller, R.E., Stern, C.D. (2007) The amniote primitive streak is defined by epithelial cell intercalation before gastrulation. Nature 449: 1049–1052.

Thomas, P., Beddington, R. (1996) Anterior primitive endoderm may be responsible for patterning the anterior neural plate and the mouse embryo. Current Biology 6: 1487–1496.

Rhim, M., Dierich, A., Shavlovit, W., Behringer, R.R., Le Meur, M., et al. (1998) Sequential roles for Otx2 in visceral endoderm and neuroectoderm for forebrain and midbrain induction and specification. Development 125: 845–856.

Perea-Gomez, A., Shavlovit, W., Sasaki, H., Behringer, R.R., Ang, S. (1999) HNF3beta and Lim1 interact in the visceral endoderm to regulate primitive streak formation and anterior-posterior polarity in the mouse embryo. Development 126: 4925–4932.

Knoch, H., Viebahn, C., Kesel, M. (1999) Heart induction in the chick by primitive endoderm of mammalian, but not avian origin. Development 126: 615–625.

Kumura, C., Yoshinaga, K., Tan, E., Suzuki, M., Aiawa, S., et al. (2000) Visceral endoderm mediates forebrain development by suppressing polarization signals. Dev Biol 235: 304–321.

Gardner, R.L., Rossant, J. (1979) Investigation of the fate of 4-5 day post-coitum mouse inner cell mass cells by blastocyct injection. J Embryol Exp Morphol 52: 141–152.

Gansel, B., Rowning, B.A., Vogeli, K.M., Rothbacher, U., Fraser, S.E., et al. (2000) Dishevelled controls cell polarity during Xenopus gastrulation. Nature 405: 81–85.

Choi, K.W., Benzer, S. (1994) Rotation of photoreceptor clusters in the developing Drosophila eye requires the nemo gene. Cell 78: 125–136.

Pandur, P., Lasche, M., Eisenberg, L.M., Kuhl, M. (2002) Wnt-11 activation of a non-canonical Wnt signalling pathway is required for cardiogenesis. Nature 418: 636–641.

Monzen, K., Shiojima, I., Hiroi, Y., Kudoh, S., Oka, T., et al. (1999) Bone morphogenetic proteins induce cardiomyocyte differentiation through the mitogen-activated protein kinase kinase kinase TAK1 and cardiac transcription factors Csx/Nkx2.5 and Gata-4. Molecular and Cellular Biology 19.

Jadicz, J.L., O'Connor, M.B., Corenvasian, E. (2006) The TGF beta activated kinase TAK1 regulates vascular development in vivo. Development 133: 1529–1541.

Komer, M., Shihoya, H., Takeda, N., Ninomiya-Tsuji, J., Yasui, T., et al. (2002) Targeted disruption of the Tab1 gene causes embryonic lethality and defects in cardiovascular and lung morphogenesis. Mech Dev 119: 239–249.

Mummery, C.L., Frieman, A., van der Saag, P.T., van den Brink, C.E., de Laat, S.W. (1985) Clonal variants of differentiated P19 embryonal carcinoma cells exhibit epidermal growth factor receptor kinase activity. Dev Biol 109: 402–410.

Kamehara, M., Goto, S. (2000) KEGG: kyoto encyclopedia of genes and genomes. Nucleic Acids Res 28: 27–30.

Kamehara, M., Goto, S., Furuichi, M., Tanabe, H., Hirakawa, M. (2000) KEGG for analysis of large gene lists using DAVID bioinformatics resources. Nat Protoc 4: P3.