Nodal protein processing and fibroblast growth factor 4 synergize to maintain a trophoblast stem cell microenvironment

Marcela Guzman-Ayala, Nadav Ben-Haim, Séverine Beck, and Daniel B. Constam*

Developmental Biology Group, Swiss Institute for Experimental Cancer Research (ISREC), Chemin des Boveresses 155, CH-1066 Epalinges, Switzerland

Edited by Kathryn V. Anderson, Sloan–Kettering Institute, New York, NY, and approved September 16, 2004 (received for review July 27, 2004)

Before implantation in the uterus, mammalian embryos set aside trophoblast stem cells that are maintained in the extraembryonic ectoderm (ExE) during gastrulation to generate the fetal portion of the placenta. Their proliferation depends on diffusible signals from neighboring cells in the epiblast, including fibroblast growth factor 4 (Fgf4). Here, we show that Fgf4 expression is induced by the transforming growth factor β-related protein Nodal. Together with Fgf4, Nodal also acts directly on neighboring ExE to sustain a microenvironment that inhibits precocious differentiation of trophoblast stem cells. Because the ExE itself produces the proteases Furin and PACE4 to activate Nodal, it represents the first example, to our knowledge, of a stem cell compartment that actively maintains its own microenvironment.

In eutherian mammals, the exchange of gas and nutrients across the placenta depends on an elaborate vascular network, which forms during embryogenesis with the help of fetal derivatives known as the trophoblast. This tissue arises from the spherical trophoderm layer of the blastocyst surrounding the inner cell mass (ICM) and the blastocoel. Upon implantation into the uterus, the ICM cavitates and forms the epiblast, while adjacent trophoderm cells proliferate and form the extraembryonic ectoderm (ExE) and the ectoplacental cone (EPC). Together with a superficial layer of visceral endoderm (VE), these structures constitute the egg cylinder. Throughout the ExE exists a reservoir of self-renewing trophoblast stem cells (TSCs) (1, 2) that provide the EPC with progenitor cells for differentiated spongiotrophoblasts and nondividing polyplid giant cells (3–5). They express essential transcription factors such as the estrogen-related receptor β (Errβ), Eomesodermin, and Cdx2, together with Bmp4, but repress differentiation markers such as Mash2. Their capacity to self-renew and proliferate in the embryo depends on a microenvironment that is established by neighboring cells of the ICM and the epiblast. A critical component of this microenvironment is fibroblast growth factor 4 (Fgf4), but additional, unknown signals are also required (6). Pharmacological inhibition of Errβ blocks the proliferative effect of Fgf4 on TSCs and triggers their differentiation toward the polyploid giant cell fate, substantiating the conclusion that it is an essential stem cell marker (7). At the egg cylinder stage [embryonic day (E) 5.5] and throughout gastrulation, the ExE in addition produces Furin and PACE4; two secreted proteases of the subtilisin-like proprotein convertase (SPC) family also known as SPC1 and SPC4 (8, 9). Recent experiments in mice showed that these proteases act together on neighboring tissues, where they specify anteroposterior asymmetry and stimulate germ layer formation and gastrulation movements (9). Here, we asked whether these proteases also influence the fate of TSCs.

Histological and gene expression analysis of mutant embryos reveals that Furin and PACE4, and a transforming growth factor β-related substrate in the epiblast encoded by Nodal are required to sustain TSCs in the ExE during gastrulation. In part, the role of Nodal is to induce Fgf4 expression in the epiblast. In addition, we use embryo explant culture assays to show that Nodal also acts directly on the ExE, where it is required alongside Fgf4 to sustain the expression of TSC marker genes. Besides identifying Nodal as an essential component of the TSC microenvironment, these findings define a cascade of reciprocal inductive interactions between the ExE and epiblast that are essential for TSCs to retain an undifferentiated character.

Materials and Methods

Mouse Strains. Mice cis heterozygous for null alleles of Furin (10) and PACE4 (8) were maintained on a mixed C57BL/6 × 129 Sv/Ev/Svj genetic background at the ISREC mouse facility in individually ventilated cages. Timed matings among cis heterozygotes were used to obtain Furin−/−;PACE4−/− double mutants [referred to as double knockout (DKO) embryos]. Mice carrying the NodallacZ reporter allele (11) were maintained on a mixed genetic background of 129Sv EV × NMRI. Genotyping by PCR was performed as described (8, 10, 11). Outbred diabetes-resistant NMRI mice were from Harlan (Horst, The Netherlands).

Embryo Explant Cultures. For explant cultures, whole embryos and epiblasts from NMRI mice were dissected during the evening of the sixth day postcoitum (E5.75) between 1700 and 2000 hours and cultured for 20 h in OptiMEM containing 15% (vol/vol) knockout serum replacement factors (Invitrogen), 1% (vol/vol) glutamine, and 100 µg/ml gentamycin sulfate in Millipore filter inserts (pore size, 12 µm) on γ-irradiated STO fibroblasts expressing leukemia inhibitory factor. For factor treatment, epiblasts were freed from VE by using trypsin/pancreatin (13). Mature recombinant Nodal and SPC-resistant precursor were produced in stably transfected 293T cells and applied in equal amounts to embryo explants as described, based on comparative quantitation by Western blot analysis (9). At this concentration, the activity of mature Nodal in 293T cells transfected with the AR3-lux luciferase reporter reached 50–80% of the maximal response and was comparable with that of 20 ng/ml activin A. By comparison, the activity of SPC-resistant precursor reached a plateau and induced 6- to 8-fold less luciferase expression, similar to what has been described after transfection (9). Con-
Err both DKO (second and third columns) and marker Mash2 is expanded toward the epiblast boundary (stippled lines) in Right the enlarged EPC (stippled lines). The high magnification (right portion at heterozygotes ( 14), was expanded at

Moreover, the expression domain of Mash2, a basic helix–loop–helix (bHLH) transcription factor that promotes the differentiation in the epiblast was down-regulated compared with control litter mates

chorion are reduced or absent. ( 14, 16). As shown in Fig. 1 (B), expression of Fgf4, Errβ, and Cdx2 was also abolished in Nodal mutants. In contrast, Mash2 is ectopically expressed throughout the ExE, as observed in DKO embryos (Fig. 1). These results are consistent with a model wherein Furin and PACE4 up-regulate the expression of the Nodal lacZ reporter allele (Center), whereas an SP-C-resistant mutant precursor precursor (lane 1) has no effect. (D) Recombinant Nodal can also rescue expression of Fgf4, whereas conditioned medium of untransfected, parental 293T cells (mock) was inactive.

Precursor Processing Stimulates Nodal Autoinduction. The earliest known function of Nodal is to specify a population of anterior VE, which emerges around E5.5 at the apex of the egg cylinder. Concomitantly, Nodal amplifies its own expression in the epiblast and overlying VE by means of autoinduction (12). Furin and PACE4 stimulate both of these processes, possibly byactivating the Nodal precursor (9). Intact whole embryos can be cultured in serum-free medium at this stage for up to 20 h without disrupting Nodal signaling, marked for example by the expression of Cripto. By contrast, if the ExE is cut off to remove the source of Furin and PACE4, Cripto and several other target genes downstream of Nodal are silenced (9). The expression of a Nodal lacZ reporter also appeared to be reduced (9), but to a lesser extent than in epiblast explants stripped of both ExE and VE (Fig. 2 B and D). Therefore, to directly test whether precursor processing stimulates Nodal autoinduction, isolated

the expense of the TSC markers Errβ, Cdx2 (Fig. 1B), and Eomesoderm (9). This finding suggests that Furin/PACE4 activities are necessary to inhibit precocious differentiation of TSCs into ectoplacental cell types. It seemed possible, therefore, that Furin and PACE4 are epistatic or act in parallel to Fgf4. Further analysis of DKO embryos revealed that Fgf4 expression was severely attenuated (Fig. 1B, n = 6/8, class I) or undetectable (n = 2/8, class II), indicating that these proteases cleave a substrate involved in the regulation of Fgf4. A known substrate is Nodal, a precursor protein of the transforming growth factor β family expressed in the epiblast and overlying VE (11, 15). Nodal can be cleaved by extracellular forms of Furin and PACE4, which thus convert the precursor to a more active, mature form (9, 16). As shown in Fig. 1B, expression of Fgf4, Errβ, and Cdx2 was also abolished in Nodal mutants. In contrast, Mash2 is ectopically expressed throughout the ExE, as observed in DKO embryos (Fig. 1). These results are consistent with a model wherein Furin and PACE4 up-regulate Fgf4 through the known potentiation of Nodal signaling.

Results

TSC Markers and Fgf4 Are Induced Downstream of Furin, PACE4, and Nodal. Histological analysis of late gastrulation stage Furin−/−;PACE4−/− DKO embryos revealed that virtually all cells in the ExE adopt an ectoplacental morphology (Fig. L4). Moreover, the expression domain of Mash2, a basic helix–loop–helix (bHLH) transcription factor that promotes the differentiation of spongiotrophoblasts in the EPC (14), was expanded at

conditioned medium of untransfected parental cells (mock) was used as a negative control. Human activin A and BMP4 (R & D Systems) were applied at 20 and 50 ng/ml, respectively. FGF4 (Sigma) was used at 40 ng/ml together with 1 μg/ml heparin (Sigma).
epiblasts were incubated with recombinant Nodal. As shown in Fig. 2C and Table 1, processed Nodal markedly up-regulated NodallacZ, whereas uncleaved precursor had no effect. Consistent with data from genetic studies (9, 12, 17), this result directly demonstrates that Nodal amplifies its own expression in an autoinductive feedback loop that is positively regulated by Nodal alone has no significant effect on Mash2. Furthermore, in negative control experiments, none of the markers examined is influenced by conditioned medium of untransfected HEK293T cells (mock).

Recombinant Nodal Can Rescue Fgf4 Expression in Epiblast Explants. Because Furin and PACE4 produced in ExE are necessary for Nodal activation, removal of the ExE should also inhibit Fgf4 expression. Confirming this prediction, Fgf4 expression was lost in epiblast explants irrespective of the presence or absence of VE (Fig. 2D). Thus, in both situations, the activity of endogenous Nodal in epiblast explants is below the threshold required for induction of Fgf4. However, upon addition of exogenous recombinant Nodal, expression of Fgf4 was restored. In contrast, BMP4, which can mediate a stimulatory effect of Nodal on Cripto (9), failed to induce Fgf4 (n = 0/6, data not shown). These results demonstrate that Nodal acting directly on epiblast cells is sufficient to maintain Fgf4 expression.

Nodal Acts on ExE Alongside Fgf4 to Sustain TSCs. Fgf4 and additional signals from the epiblast are essential to prevent TSCs from undergoing differentiation (6), but the expression pattern of stem cell markers in isolated ExE explants at early stages has not been examined. As shown in Fig. 3A, ExE explants cultured without the epiblast rapidly down-regulated Errβ, Cdx2, and Eomes mRNA, whereas Mash2 transcripts ectopically accumulated throughout the explants. These results show that isolated ExE explants mimic the molecular defects observed in Furin/PACE4 DKO and Nodal mutants, and rapidly lose their stem cell character. However, the loss of TSCs in a class of Furin/PACE4 DKO embryos appeared to be slightly less severe compared with

Table 1. Regulation of gene expression in epiblast explants

<table>
<thead>
<tr>
<th>Gene</th>
<th>Staining</th>
<th>None</th>
<th>Mock</th>
<th>Ndl</th>
<th>pre-Ndl</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nodal</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>lacZ</td>
<td>+</td>
<td>12</td>
<td>1</td>
<td>6</td>
<td></td>
</tr>
<tr>
<td></td>
<td>++</td>
<td>2</td>
<td>6</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>+++</td>
<td>2</td>
<td>6</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Fgf4</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>–</td>
<td>6</td>
<td>6</td>
<td>11</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>+</td>
<td>0</td>
<td>0</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>++</td>
<td>0</td>
<td>0</td>
<td>8</td>
<td>7</td>
</tr>
</tbody>
</table>

Numbers correspond to the total of explants from two to four independent whole-mount in situ stainings that were recovered after treatment with the factors indicated (top row). Processed Nodal (Ndl), uncleaved precursor (pre-Ndl), or conditioned medium of untransfected HEK293T cells (mock) were applied as described in Materials and Methods. The relative intensities of staining (–, +) were scored visually.
Nodal mutants and ExE explants, possibly because of weak residual expression of Fgf4 (Fig. 1, class I). Therefore, we wished to determine whether Fgf4 alone might be sufficient to transiently restore normal gene expression in ExE explants if applied in increased amounts. Indeed, Fgf4 at a concentration of 40 ng/ml potently inhibited ectopic expression of Mash2 by contrast, recombinant Nodal, activin A (n = 0/4), or BMP4 (n = 0/3) had no effect (Fig. 3B, Table 2, and data not shown). However, although this finding shows that cultured ExE explants respond to Fgf4, we did not observe a corresponding up-regulation of Errβ or Cdx2 mRNAs, and Eomes was only weakly induced in 40% of the explants (Fig. 3B and Table 2). Therefore, we asked whether additional signals act in parallel to Fgf4. Interestingly, both Errβ and Eomes were significantly induced if FGF4 was added in combination with Nodal (Fig. 3B), the expression levels being comparable with those observed in whole-embryo cultures (Fig. 3A). This effect was mimicked by recombinant activin A (Errβ, n = 9/16; Eomes, n = 8/11), a related protein sharing its signaling receptors with Nodal, but not by BMP4 (Errβ, n = 0/15; Eomes, n = 0/9). The combination of Fgf4 and Nodal also restored expression of Cdx2 (Fig. 3B). We conclude that Errβ, Cdx2, and Eomes expression during normal development is maintained by Nodal itself, acting together with the epiblast-derived relay signal Fgf4.

**Discussion**

Overall, our results show that the ExE inhibits its own differentiation into ectoplacental tissue through reciprocal interactions with the epiblast that are mediated by Furin, PACE4, and Nodal (Fig. 4B). Previous analysis of Furin;PACE4 DKO embryos suggested that an important function of SPCs is to stimulate Nodal autoinduction in the epiblast (9). By using cultured embryo explants, we here provide direct evidence that Nodal can only amplify its expression after SPC cleavage of the precursor. Furthermore, our results show that Nodal is necessary to maintain the expression of Fgf4. In contrast to Nodal mutants, however, a majority of Furin;PACE4 DKO embryos still weakly expressed Fgf4 throughout the epiblast. The ectopic distribution of Fgf4 transcripts in this class of mutant embryos mirrors the residual expression of Nodal (9). Therefore, it will be interesting to determine in future studies whether Nodal can induce Fgf4 at a lower threshold compared with other target genes.

Additional experiments show that Fgf4 mediates an inhibitory effect of Nodal on the expression of the early differentiation marker Mash2 in the ExE. However, besides inducing Fgf4 as a secondary signal, Nodal itself also acts directly on the ExE alongside Fgf4 to maintain expression of Errβ, Cdx2, and Eomes. These transcription factors mark undifferentiated TSCs (1), and are required to sustain trophectoblast growth (18–20). Thus, our results argue that both Fgf4 and Nodal are essential components of a microenvironment that is required to maintain a population of self-renewing TSCs (Fig. 4).

How do Nodal and Fgf4 signaling synergize in the ExE to induce TSC markers? To address this question, we tested whether Nodal up-regulates the expression of FgfR2, the putative receptor mediating Fgf4 signaling. However, FgfR2 is normally expressed both in cultured Nodal−/−embryos and in a collapsed ExE explants (M.G.-A., unpublished work). Conversely, Fgf4 does not stimulate canonical Nodal signaling via the Smad pathway because it failed to potentiate induction of the luciferase reporter construct AR3-lux in 293T cells (ref. 21 and M.G.-A., unpublished work). Therefore, Nodal and Fgf4 signaling appear to act in parallel and may be integrated directly at the promoter level of target genes as observed for the T box transcription factor brachyury (22). Because all of the TSC markers examined are transcription factors, it is also possible that only one of them directly depends on synergistic activation by Nodal and Fgf4, and subsequently up-regulates the others.

In keeping with a direct role in trophoderm development, transfection of Nodal cDNA into cultured TSCs was recently found to attenuate their terminal differentiation into giant cells marked by the expression of the late differentiation marker PI-1. Conversely, in embryos that are homozygous for a hypomorphic Nodal allele, giant cell and spongiotrophoblast formation are accelerated at the expense of labyrinth growth (23). Thus, Nodal was suggested to antagonize the terminal differentiation of trophoblasts during midgestation because its expression is reactivated at that stage specifically in the spongiotic layer. We propose that a Nodal signaling network in addition already acts before gastrulation to mediate an intricate crosstalk between the ExE and epiblast that enables these tissues to sustain one another, possibly to grow and differentiate in a coordinated fashion. Thus, whereas the ExE normally serves as a reservoir of undifferentiated TSCs to sustain trophoblast growth beyond gastrulation, genetic inactivation of Nodal or its convertases, Furin and PACE4, triggers premature expression of an ectoplacental phenotype. The loss of ExE, in turn, prevents further growth of the epiblast and, hence, functions as a checkpoint to arrest development. Whereas this early developmental arrest precluded a clean and comprehensive analysis of terminal differentiation, we do not rule out the possibility that the ectopic EPC tissue arising in the absence of Nodal signaling in principle could give rise to one or several differentiated lineages, including polyploid giant cells. To address this question, cultured TSC lines may be useful as a model system to characterize in more detail the signaling pathways of Nodal and Fgf4 and their potential to perhaps modulate the fate of TSC derivatives also during later developmental stages that have not been analyzed in the present study.

We thank Janet Rossant (Mount Sinai Hospital, Toronto, Canada) for *in situ* probes of ExE markers and Andreas Trumpp and Anne Grapin-Botton for valuable discussions. This work was supported by the Barmet Foundation (to M.G.-A.) and Swiss National Science Foundation Grant 31-62031.00 (to N.B.-H. and D.B.C.).