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**Endothelium-derived essential signals involved in pancreas organogenesis**

Talavera-Adame D *et al.* Endothelium role in pancreas organogenesis

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**Abstract**

Endothelial cells (ECs) are essential for pancreas differentiation, endocrine specification, and endocrine function. They are also involved in the physiopathology of Type 1 and Type 2 diabetes. During embryogenesis, aortic ECs provide specific factors that maintain the expression of key genes for pancreas development such as pancreatic and duodenal homeobox-1. Other unknown factors are also important for pancreatic endocrine specification and formation of insulin-producing beta cells. Endocrine precursors proliferate interspersed with ductal cells and exocrine precursors and at some point of development; these endocrine precursors migrate to pancreatic mesenchyme and star forming the islets of Langerhans. By the end of the gestation and close to birth, these islets contain immature beta cells with the capacity to express vascular endothelial growth factor and therefore to recruit ECs from the surrounding microenvironment. ECs in turn produce factors that are essential to maintain insulin secretion in pancreatic beta cells. Once assembled, a cross talk between endocrine cells and ECs maintain the integrity of islets toward an adequate function during the whole life of the adult individual. This review will focus in the EC role in the differentiation and maturation of pancreatic beta cells during embryogenesis as well as the current knowledge about the involvement of endothelium to derive pancreatic beta cells *in vitro* from mouse or human pluripotent stem cells.

**Key words:** Endothelium; Endothelium-derived signals; Differentiation; Pancreas development; Organogenesis

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**Core tip:** Many studies have demonstrated that endothelial cells (ECs) have an important role in organogenesis. For instance, during embryogenesis, aortic ECs provide specific factors that maintain the expression of key genes for pancreas development. Other unknown factors are also important for pancreatic endocrine specification and formation of insulin-producing beta cells. In addition, by the end of the gestation and close to birth, pancreatic islets contain immature beta cells with the capacity to express factors that recruit ECs from the surrounding microenvironment and form a functional unit that will lasts for the whole life of the individual. In the present review, we will analyze the current endothelial-derived factors called angiocrine factors that are essential in organogenesis and we will focus the role of these factors in pancreas development and pancreatic beta cells.

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**INTRODUCTION**

At present, insulin producing cells have been derived from different sources[1–3]. With the emergence of embryonic stem cells (ESCs) and induced pluripotent stem cells (iPSCs), that can be plated *in vitro*, the potential production of several cell types including pancreas can be now achieved[2,4–11]. Endothelial cells (ECs) are present at early stages during embryogenesis[12–15]. These cells are the main component of most niches[16]. Therefore, ECs interact early with developing tissues during organogenesis even before they are able to form blood vessels and nourish specific regions[16–19]. It has been reported that the absence of ECs conducts to agenesis of some organs such as pancreas[20,21]. This fact points out the essential role of EC signaling during organogenesis. Apparently ECs are not only involved in pancreas differentiation, it has been shown that they are essential for endocrine differentiation as well[20]. Endocrine progenitors give rise to immature beta cells that recruit ECs after expression of vascular endothelial cell growth factor (VEGF)[18,20]. These ECs in turn provide factors to promote beta-cell maturation and stabilizes beta-cell function. In this review, we will be focusing in the role of ECs in the differentiation and maturation of beta cells *in vivo* and *in vitro* with the emerging technology of human PSCs that can be expanded *in vitro*.

**ENDOTHELIUM AS AN ENDOCRINE ORGAN**

Extensive studies with ECs have demonstrated that these cells play essential roles in immunity, inflammation, angiogenesis, and tumor metastasis and not only line the interior surface of blood vessels[22,23]. ECs are found at the interface between blood and other cell types. They not only maintain the blood fluid but also have a great plasticity that allows these cells to accomplish several essential functions to maintain homeostasis[18]. However, recent studies demonstrated that the endothelium is capable of releasing growth factors and cytokines that play an essential role in organogenesis[24] (Table 1). The term angiocrine has been proposed by Butler and collaborators and indicates the capacity of ECs to release growth factors and cytokines that may be involved in organogenesis[25]. For instance, bone marrow sinusoidal ECs (SECs) promote differentiation of hematopoietic stem cells (HSCs) through “angiocrine factors” such as HGF, Wnt2, and Notch[25,26]. Additionally, Butler and colleagues reported that ECs promote self-renewal of hematopoietic stem cells when these cells are interacting in co-culture [25]. Apparently, Notch ligands expressed by ECs are associated with this response[25]. Another studies indicate that ECs from liver, called sinusoidal ECs (LSECs), express factors such as VE-cadherin, Factor VIII and vascular endothelial growth factors 2 and 3 (VEGF2, VEGF3) and these cells release angiocrine factors that may be involved in liver regeneration[26]. In another work it has been described that angiocrine factors are able to regulate tumor growth[27]. Other studies demonstrated that mice deficient in FLK-1 (VEGF receptor), die at embryonic day E9.5 or E10.5 because immaturity of blood cells and blood vessels[28]. Absence of embryonic liver budding is also present in these mice indicating that ECs play an important role during the early phases of organogenesis. In a similar work, it has been observed that ECs are also involved during the early stages of pancreas development[20]. Another research work indicated that signals from myocardium, such as those exerted by bone morphogenetic protein-2 (BMP-2), can promote epithelial-mesenchymal transformation mediated by ECs[24,29]. In the kidney, VEGF, bFGF, and PDGF coordinate cellular differentiation, proliferation, and migration[30]. It has been suggested that ECs promote the differentiation of endoderm cells toward liver or pancreas through secretion of HGF[17,19,24,28,31,32] (Table 1). At the same time, reciprocal interactions between tissue-specific cell types and ECs ensure coordinated growth and adequate tissue function. For instance, it is known that neurogenesis takes place close to blood vessels in adult brains[33]. Additionally, brain-derived neutrophic factor (BDNF) secreted by ECs promotes neurogenesis and angiogenesis in the brain of song birds[34]. Another report indicates that pigment epithelium-derived factor (PEDF) is secreted by ECs and enhances self-renewal of neural stem cells (NSCs)[35]. Regarding the pancreas, it has been shown that pancreatic endoderm attract endothelial progenitor cells (EPCs) or angioblasts by expression of SDF-1/CXCL12[36]. Expression of PDX-1 appeared in endoderm cells in contact with angioblasts via LIM domain only 2 (LMO2) suggesting that angioblasts may induce expression of PDX-1[37]. Functional blood vessels may induce differentiation before they carry blood. However, it has been described that some blood factors such as sphingosine-1-phosphate (SIP) are important for the differentiation and maturation[38]. For instance, it has also been described that beta-cell differentiation can be regulated by oxygen tension *via* hypoxia-inducible factor 1 alpha (HIF-1α)[39]. This fact suggests that blood factors can also be involved in the complete differentiation and maturation of pancreatic endocrine cells.

**ENDOTHELIUM ROLE IN THE PANCREATIC NICHE**

The pancreas originates from ventral and dorsal buds formed in the foregut at 8.5 d post-coitum (d.p.c) of gestation in mice and Carnegie stage 12 (CS12) in humans[54–56]. The cells that composed these buds express transcription factors such as pancreatic and duodenal homeobox 1 (PDX-1) which is a key regulator of pancreas development[57–59]. However, before these cells express these genes, the cells interact with other surrounding cells such as those that compose the notochord and factors such as fibroblast growth factor-2 (FGF-2) and activin-βB suppress the expression of sonic hedgehog (*SHH*) locally and promotes expression of PDX-1 in the subjacent endoderm[60–63] (Figure 1).

Once the mesoderm layer start proliferating, other signals from aortic endothelial cells (AECs) and mesenchyme (MCs) continue interacting with these PDX-1 expressing cells that give rise to acinar cells that harbor pancreatic exocrine and endocrine progenitors[64] (Figure 2). As these progenitors continue receiving more surrounding instructive signals, definition of cell function is established and the exocrine cells form acini while the endocrine cells form islets of Langerhans[54] (Figure 3). However, these islets contain immature beta cells that will become more mature after islet vascularization that allows closer interaction between beta cells and ECs

***Embryonic endothelium and endoderm pre-patterning***

The vascular system is one of the first tissues that develop during embryogenesis. Mesodermal progenitors coalesce in the yolk sac and give rise to endothelium and blood cells[15]. Endothelial cells exert inductive effects in specific points were they are in contact with pre-patterned DE cells of the FG[20]. DE is forming during gastrulation and Nodal, a member of transforming growth factor (TGFβ) family, plays a central role in DE formation[65]. At these points of effective cell-cell interactions, the gut endoderm has to be competent to respond to EC-derived signals. Competence of these cells takes place during gastrulation when the mesoderm germ layer invades the middle area between primitive ectoderm and endoderm[66]. Apparently mesoderm-derived cells are required to maintain the phenotype of posterior endoderm that includes the site where pancreas and duodenum will be formed. Therefore, anterior-posterior (A-P) endoderm axis will be sustained by the presence of mesoderm-derived factors such as Wingless-type MMTV integration site family (Wnt), fibroblast growth factor (FGF), bone morphogenetic proteins (BMPs), and retinoic acid (RA)[63,67]. For instance, high signaling of the canonical WNT/β-catenin pathway promotes endoderm posterior pattern with foregut-derived structures such as pancreas and liver[68]. Interestingly it has been found that some of these factors are produced by ECs[13,42,53,69–71].

***Pancreatic specification induced by surrounding endothelium***

Endothelial signaling is required to induce insulin gene expression during pancreas development[20]. Cell-cell interactions between definitive endoderm and aortic endothelial cells take place at about 9-10 d.p.c. in mice and give rise to PDX-1 expressing cells[18,20]. Apparently, these interactions are also essential to promote insulin expression in pre-patterned endoderm[20]. However, signals from the developing pancreas to embryonic endothelium also promote endothelium-specific phenotype and this interactions are crucial for adequate organ function in adulthood[32]. As mentioned above, the first signals to promote expression of PDX-1 come from the notochord that produce factors such as activin-βB and FGF-2[60]. These cross-talk take place between cells from notochord and cells from the subjacent endoderm leading to inhibition of factors such as *SHH* and promotion of PDX-1 expression[60]. However, these permissive signals are apparently replaced by instructive signals from the growing mesoderm and AECs that become in close proximity with pre-patterned endoderm. Interestingly, ECs not only exert these signals directly to the pre-patterned endoderm but also promote the survival of adjacent mesodermal cells that produced essential factors such as Islet-1 (Isl1)[64]. Homozygous mice lacking expression of Flk-1 (*flk-1*-/-) had absence of aorta with no formation of dorsal mesenchyme that led to diminished expression of PDX-1 in the subjacent endoderm[64]. However, lateral and ventral mesenchyme were not affected and PDX-1 positive cells also appeared in the ventral endoderm[64]. These findings indicate that aortic EC signaling is essential to maintain the dorsal mesenchyme and therefore to direct differentiation of dorsal pancreatic endoderm. Additionally, at later embryo stage, *flk-1*-/- mice showed dorsal mesenchyme that does not express Isl-1 in absence or aortic ECs suggesting that endothelial-cell signaling promotes Isl-1 cell expression from dorsal mesenchymal cells[64]. In addition, it has been reported that mesenchymal cells also express bone morphogenetic proteins (BMPs) and that these proteins have a pivotal role in pancreas development[71,72].

***Signals from pancreatic mesenchyme toward exocrine pancreas***

It is well known that once the mesenchymal cells proliferate between aortic ECs and the foregut endoderm, the aorta is pulled out from the subjacent gut (compare Figure 2 with Figure 3)[20,55]. This fact implies that the subjacent foregut starts receiving signals from the mesenchyme and that a gradient can be formed with diluted signals from aortic ECs. However, it has been demonstrated that ECs maintain the integrity of the subjacent mesenchyme and that absence of aorta promotes apoptosis of the mesenchymal cells and avoids the formation of the dorsal pancreas[64]. As mentioned before, mesenchymal cells between aorta and foregut maintain the expression of PDX-1 in pancreatic endoderm cells through the expression of the transcription factor Isl-1[54]. In addition, signals from mesenchyme such as Fgf-10 are essential to promote proliferation of pancreatic buds that already received signals from ECs[64]. Apparently, the specification of pancreatic fate is determined by permissive signals from notochord and instructive signals by ECs that are maintained by mesenchyme after the aorta is pulled out from the gut. It has been suggested recently that specification of endocrine phenotype also takes place during the close interaction between aortic ECs and pancreatic progenitors that express PDX-1 within the pancreatic bud[54]. Previous experiments indicated that insulin is expressing in foregut explants only after co-culturing with aortic endothelial cells[20]. However, the signals from ECs that promote such specification in the foregut endoderm toward the formation of pancreatic endocrine progenitors are still under investigation. In the same way more characterization is required to identify the signals from ECs that promote survival and adequate function of mesenchyme. Therefore, functional mesenchyme is crucial for appropriate signaling to the subjacent foregut endoderm toward maintenance of the pancreatic phenotype and branching[43].

**ENDOTHELIUM ROLE IN PANCREATIC BETA-CELL MATURATION**

***Pancreatic endothelium during islet neogenesis***

Pancreatic endocrine progenitors receive signals from aortic ECs and mesenchyme that determined their fate. However, they remain interspersed with the ductal cells that form the epithelium of the growing branches. At certain time of development, these cells receive still unknown instructions to migrate from the ductal area to the mesenchyme region. Once in the mesenchyme, these pancreatic progenitors that are apart from ductal cells form the islets of Langerhans[73]. This fact raises two questions: (1) are there signals from ECs that promote islet neogenesis? And (2) once the blood vessels are formed are there factors in the blood stream that promote the final maturation of beta cells? The answer to these questions is still unknown. The cells that migrate are pancreatic endocrine progenitors that give rise islets of Langerhans composed of alpha, beta, delta, and PP cells that will produce glucagon, insulin, somatostatin, and pancreatic polypeptide respectively. These islets will be distributed differently into the pancreas and apparently will be subjected to different stimuli[74,75]. Although endocrine specification takes place before migration, maturation of endocrine cells occurs at islet level and coincides with islet vascularization[54,76]. There is a significant growth of islet cells that correlates with islet endothelial-cell proliferation in rats the first week after birth[76]. For instance, it has been found endocrine cells with higher proliferative capacity closer to blood vessels[76]. Furthermore, it is known that ECs are able to produce hepatocyte growth factor (HGF) which is a potent mitogen for beta cells and ECs[31,77]. Therefore, the endothelial signaling is essential for beta-cell maturation. For instance, immature beta cells are formed some days before birth and maturation occurs several days after birth[76]. During this period, immature beta cells express vascular endothelial growth factor (VEGF) and start recruiting ECs by the islet that will provide signals for further differentiation and maturation[32,78]. However, along with EC stimuli, another signals should also be considered such as hormones that can reach beta cells through the blood stream once the vascular network is established and may also be essential for cell maturation[79]. Apparently, endogenous insulin has a minor role for the glucose homeostasis before birth[80]. In this condition, insulin provided by the mother regulates glucose in the fetus[80]. This fact suggests that fetal beta cells are not mature enough at birth to maintain the glucose homeostasis and that further maturation can be promoted by ECs after birth.

**ENDOTHELIUM ROLE IN PANCREATIC BETA-CELL FUNCTION**

The role of ECs in beta-cell function and pathology has been previously described[47,81–82]. It has been found that ECs from islets correspond to fenestrated endothelium[83,84]. Apparently the characteristics of islet endothelium differ from pancreatic exocrine endothelium and endothelium from other regions[47,83,85]. For instance, pancreatic endocrine capillaries have higher diameter than exocrine capillaries and endothelium from endocrine capillaries have 10 times more fenestrae that endothelium from exocrine capillaries[83]. These facts suggest that cell-cell interactions and signaling between endothelium and the surrounding cells are different even in the same organ. Pancreatic beta cells have polarity with an apical and basolateral membrane and insulin vesicles are more dense in the apical region close to ECs[47,86]. This aspect is very important when considering the ability of beta cells to release insulin into the blood stream. After islet transplantation, the absence of suitable EC-signals for polarization can be crucial to avoid the appropriate insulin release into the capillaries. Islet ECs express common markers of ECs but one specific marker called the proteinase inhibitor and angiostatin factor α1-antitripsin[70,84]. This marker of specific islet ECs can be absent in surrogate vasculature with deregulation of islet function. Apparently endothelial progenitors (EPCs) or islet ECs are important for islet revascularization after transplantation[87–89]. In this sense, any islet injury leads to islet restoration through recruitment of EPCs toward islet re-vascularization with specific islet ECs and beta cell function[87]. For instance, normoglycemia is improved in streptozotocin-treated animals after co-transplantation of EPCs and islets[87].

**ENDOTHELIUM AND BETA-CELL REGENERATION *IN VIVO* AND *IN VITRO***

As a first approach to investigate the role of endothelium in beta-cell differentiation, we studied the role of *in vivo* surrogate vasculature in mouse embryoid body (EB) differentiation using quail chorioallantoic membranes (CAM)[90]. We found that some cells expressed cardiotin, myosin heavy chain, collagen IV, CD34, CD31, and neurofilament. Although some epithelial cells appeared, no cells derived from endoderm were identified[90]. Then, studies using co-cultures between human microvascular ECs (HMECs) and mouse EBs were performed[52,53,91]. In these studies, ECs from human dermis were able to induce differentiation of mouse EBs to pancreatic progenitors and insulin-producing cells[52]. Furthermore, BMP-2/-4 were involved in this differentiation process as evaluated by the effects of agonists (recombinant BMPs), and specific antagonist of BMP bioactivities (Noggin, Chordin). BMPs are members of the transforming growth factor beta ( TGF-β) superfamily[92]. In addition to the effects of BMP antagonists, we explored the levels of phosphorylation of SMAD1,5,8 in cells that expressed proinsulin[52,53]. The role of BMPs in pancreas development has also been explored previously[72,93]. We demonstrated that HMECs or mouse dermis as well as mouse aortic ECs (AECs) expressed BMPs and that BMP-2 and BMP-4 increased the phosphorylation levels of SMAD1,5,8 in pancreatic progenitors and beta-like cells derived from mouse ESCs[52,53,94]. These findings together with previous works pointed out the important role of ECs in beta-cell differentiation *in vitro*. We recently have observed that when human ESCs (cell line H9) are co-cultured with HMECs, the formation of proinsulin positive cells takes place in about twenty days in close proximity to internal ECs without the use of additional growth factors (Figure 4).

In the model using mouse EBs, we observed that ECs promote up-regulation of BMPs within EBs[53]. However, the target cells that produce these BMPs are still unknown. Some good candidates for these cells are mesenchymal cells since it has been demonstrated that ECs are essential to maintain dorsal pancreatic mesenchyme during pancreas moprhogenesis that may promote pancreas differentiation within EBs[64] (Figure 5). However, further studies should be done to demonstrate that internal ECs are able to trigger beta-cell differentiation through signaling to mesenchymal stem cells. At present, ECs can be generated *in vitro* from hiPSCs or hESCs and this studies can be important to answer this question[95,96].

In this model, an excess of human microvascular ECs (HMECs) are surrounding a human embryoid body which is composed of endoderm, ectoderm, and mesoderm cells. External ECs produce factors such as BMPs and other EC-derived factors that promote upregulation of endogenous BMPs in still unknown target cells (good candidates are mesenchymal or internal ECs). These BMPs together with other unknown factors may promote differentiation of multipotent cells (MC) toward beta-like cells (BC) and other cell lineages (OCL)[52,53].

**ENDOTHELIUM AND BETA-CELL PATHOGENESIS**

ECs play an important role for the pathogenesis of Type 1 (T1DM) and Type 2 diabetes mellitus (T2DM). In T1DM, ECs will allow infiltration of leucocytes[47]. Interestingly the problem that generates diabetic multiple complications relay in endothelial-cell function that can be the key to cure diabetes since recently the role of endothelium in beta-cell differentiation and maturation *in vitro* has been emphasized[52,53]. In addition, it has been suggested that EC pathology can lead to islet dysfunction suggesting that ECs are essential to maintain islet function in adults[97].

**CONCLUSION**

In this review we focused in the essential role of endothelium for pancreatic endocrine differentiation, functional maturation, and islet dysfunction. ECs play a key role during the differentiation of the dorsal pancreas by maintaining the expression of transcription factors necessary for pancreas development including endocrine progenitors. Before birth, immature beta cells recruit ECs close to their microenvironment and these ECs provide signals for further maturation and function of pancreatic beta cells. ECs involvement in cellular diapedesis, inflammation, and vessel fibrosis, which leads to islet dysfunction, has been demonstrated. In addition, ECs co-transplanted with islets have demonstrated to improve the engraftment of human islets. Finally, ECs are able to provide signals *in vitro* for derivation of functional beta-like cells from human pluripotent stem cells. Therefore, the study of interactions between EC and beta cells is relevant for future clinical applications in regenerative medicine.

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**Table 1** EC-derived factors related to organogenesis

|  |  |  |  |
| --- | --- | --- | --- |
| **Angiocrine factors** | **Source** | **Target** | **Ref。** |
| Notch ligands (1 and 2) | Bone marrow ECs; Brain ECs | Neural progenitors and HSCs  | [40,41] |
| VEGF | ECs | ECs and several tissues such as islets | [12,42,43] |
| bFGF | ECs | ECs and several tissues such as islets | [43] |
| PDGF | Brain capillaries | ECs | [42] |
| HGF | Lunga capillaries, SECs,Islet capillaries  | Lung epithelium,Hepatocytes,Islet beta cells | [26,31,44,45] |
| Endothelins | ECs | Lung,Neural Cells | [46] |
| EG-VEGF | ECs | Endocrine Glands | [32] |
| Brain-derived neurotrophic factor  | Brain Microvascular endothelium | Neuronal Precursors,Islet endothelium | [33] |
| Pigment epithelium-derived factor  | Brain capillaries  | Neural stem cells  | [35,47]  |
| Vessel-derived stromal-derived factor 1(SDF-1/CXCL12) | Microvascular Endothelium | Endoderm and pancreatic beta cells | [48] |
| Wnt2 | Sinusoidal ECs | Hepatocytes | [26] |
| S1P | Plasma (plateles) | Pancreatic multipotent progenitor cells | [38] |
| CTGF | Pancreas capillaries | Pancreatic endocrine cells | [49] |
| Laminin  | Islet capillaries | Islet endocrine cells | [50] |
| Collagen IV | Islet capillaries | Islet beta-cells | [51] |
| BMP-2 | ECs/MSCs | Islet beta-cells | [52,53] |
| BMP-4 | ECs/MSCs | Islet beta-cells; hepatocytes; cardiomiocytes | [52,53] |
| BMPR1A | ECs/MSCs | Islet beta-cells; hepatocytes; cardiomiocytes | [52,53] |

ECs: Endothelial cells; BMP: Bone morphogenetic protein; VEGF: Vascular endothelial cell growth factor; HGF: Hepatocyte growth factor; bFGF: Basic fibroblast growth factor; PDGF: Platelet-derived growth factor; CTGF: Connective tissue growth factor.



**Figure 1 Early cell-cell interactions that give rise to pancreatic cells derived from definitive endoderm (DE) of the foregut (FG).**Factors released by notochord (N) such as activin-βB and FGF-2 permit expression of PDX-1 and suppress expression of sonic hedgehog*.*



**Figure 2 Endothelial-derived signals maintain expression of PDX-1 and promote pancreatic endocrine differentiation.** Once the aorta is forming, aortic ECs (AECs) interact closely with FG/DE cells that maintain expression of PDX-1 and form the dorsal pancreatic bud (DPB). All the EC-derived signals are still under investigation (?). ECs: Endothelial cells.



**Figure 3 Endothelial-derived signals promote the survival of pancreatic mesenchyme which is essential for pancreas development.** Mesenchymal cells (MCs) appear between AECs and the dorsal pancreatic bud (DPB) and promote proliferation and survival of differentiated cells. Immature beta cells (BCs) that co-express PDX-1 and insulin migrate toward the mesenchyme and form cell clusters that will become islet of Langerhans that will recruit ECs that become islet ECs (iECs) and produce collagen IV and laminins which promote insulin expression. AECs crosstalk with MCs and maintain the integrity of these cells toward adequate exocrine and endocrine pancreas development. AECs: Aortic ECs; ECs: Endothelial cells.



**Figure 4 Analysis of blood vessels and insulin-producing cells in embryoid bodies obtained from human embryonic stem cell line H9.** Human embryonic stem cells were cultured in suspension for 5 d to obtain embryoid bodies (EB). After attachment on coverslips for 24 h. Some EBs were cultured alone or together with human microvascular endothelial cells (HMECs). Then, after 20 d both groups of EBs were fixed and stained with with anti-proinsulin (green) (a marker for pancreatic beta cells), anti-CD31 (red) (a marker for endothelial cells), and DAPI (blue) (that stains the nuclei). A: EB cells cultured alone that do not show proinsulin or CD31 expression. In contrast with, (B) EB cells co-cultured with HMECs at passage 14 in which we can find cells that express proinsulin in close proximity with cells that express CD31. HMECs did not stain positive for CD31 at the dilutions used indicating that the ECs are forming within EBs.



**Figure 5 Diagram that explains the possible effects of external Endothelial cells toward *in vitro* beta-cell differentiation in human embryoid bodies.** ?: Unknown factors, differentiation steps, or cell lineages; BMPs: Bone morphogenetic proteins; EC: Endothelial cells; EB: Embryoid bodies; BC: Beta cells; OCL: Other cells lineages; MC: Multipotent cell.