Accessing the human trophoblast stem cell state from pluripotent and somatic cells

Rowan M. Karvas1 · Laurent David2,3 · Thorold W. Theunissen1

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Abstract

Trophoblasts are specialized epithelial cells that perform critical functions during blastocyst implantation and mediate maternal–fetal communication during pregnancy. However, our understanding of human trophoblast biology remains limited since access to first-trimester placental tissue is scarce, especially between the first and fourth weeks of development. Moreover, animal models inadequately recapitulate unique aspects of human placental physiology. In the mouse system, the isolation of self-renewing trophoblast stem cells has provided a valuable in vitro model system of placental development, but the derivation of analogous human trophoblast stem cells (hTSCs) has remained elusive until recently. Building on a landmark study reporting the isolation of bona fide hTSCs from blastocysts and first-trimester placental tissues in 2018, several groups have developed methods to derive hTSCs from pluripotent and somatic cell sources. Here we review the biological and molecular properties that define authentic hTSCs, the trophoblast potential of distinct pluripotent states, and methods for inducing hTSCs in somatic cells by direct reprogramming. The generation of hTSCs from pluripotent and somatic cells presents exciting opportunities to elucidate the molecular mechanisms of human placental development and the etiology of pregnancy-related diseases.

Keywords Placenta · Trophoblast · Pluripotency · Stem cells · Reprogramming · Organoids

Introduction

The placenta is a complex organ system that mediates the exchange of nutrients, gases and waste products between the mother and the developing fetus. Human placentas are hemochorial, which means that trophoblast cells come into direct contact with maternal blood. Placental development occurs in two morphologically and temporally distinct stages, the pre-villous and villous stages. The pre-villous placenta emerges upon implantation of the embryo within the maternal decidua starting at 7–9 days postfertilization (dpf). Multinucleated primitive syncytiotrophoblasts in the outer trophectoderm (TE) layer of the blastocyst rapidly divide and form a shell, which surrounds the epiblast (EPI, will give rise to the fetus) and primitive endoderm (PrE, will become the yolk salk) [4, 5]. Extraembryonic mesoderm descends from the embryonic compartment around 14–16 dpf and stretches through this shell, aligning with CTBs that further differentiate into two functionally distinct terminally differentiated trophoblast cell types: syncytiotrophoblast (STB) and extravillous trophoblast (EVT) [6]. After the emergence of the villous core, which is supplied with fetal blood vessels and placental macrophages (Hofbauer cells), this structure is considered the villous placenta and is maintained throughout the remainder of pregnancy.

STBs emerge directly from the underlying CTBs and are in direct contact with maternal blood. While many trophoblast cell types in the first trimester express placental
hormones, STBs are the main manufacturers of human chorionic gonadotropin (hCG), which communicates the presence of the fetus to the maternal system, and other signaling hormones responsible for altering maternal metabolism, including leptin, prolactin-growth hormone family, and various steroid hormones [7]. EVTs arise from the tips of villi that form a prominent column extending to the maternal endometrium. The base of these columns contains a proliferative pool of EVTs that eventually travel to the endometrium, complete partial epithelial-to-mesenchymal transition (EMT) [8], and invade multiple uterine structures [9]. Interstitial EVTs reside within the decidua and invade deeply into the myometrium where they interact with maternal immune cells, obtain nutrients from maternal glandular epithelial cells, and invade uterine veins [10]. Although glandular invasion is yet to be fully characterized, arterial invasion and remodeling are a well-defined feature of EVTs. EVTs remodel uterine spiral arteries by inducing apoptosis in the smooth muscle layer [11], expressing similar adhesive proteins as vascular endothelium [12], and repopulating the walls of arteries in order to establish a consistent supply of maternal blood to the fetus. Insufficient arterial remodeling by EVTs has been identified as a leading cause of severe forms of pre-eclampsia [13].

Owing to ethical and regulatory restrictions on studying the human placenta in vivo, and the scarcity of samples obtained through elective terminations, our understanding of human placental development remains limited. Primary placental cultures are particularly difficult to perform and immortalized trophoblast cell lines fail to recapitulate the emergence and development of trophoblast cells from an undifferentiated progenitor state [14–16]. Self-renewing trophoblast stem cells (TSCs) were first derived from E3.5 mouse blastocysts by Janet Rossant’s laboratory in 1998 [17]. Mouse TSCs can be propagated indefinitely in the presence of fibroblast growth factor 4 (FGF4), heparin, and fibroblast conditioned medium and differentiate into specialized trophoblast cell types by withdrawal of FGF4 and fibroblast conditioned medium, among other methods [18–20]. The derivation of equivalent TSCs in the human system remained challenging for many years, but in 2018 Takahiro Arima’s laboratory in Japan reported the successful derivation of self-renewing human TSCs (hTSCs) from first-trimester placental CTBs and blastocysts [21]. Importantly, these hTSC lines were capable of undergoing lineage-directed differentiation into specialized EVT and STB cell types [21].

Human pluripotent stem cells (hPSCs), including human embryonic stem cells (hESCs) and induced pluripotent stem cells (iPSCs), have been reported to acquire trophoblast-like fates in response to bone morphogenetic protein 4 (BMP4) [22–24]. However, hPSCs derived under conventional conditions exhibit biological and molecular features consistent with a post-implantation epiblast (EPI) identity [25]. Based on the alignment with single cell expression data from monkey embryos, conventional – also known as “primed”—hPSCs most closely resemble the late post-implantation EPI just prior to gastrulation, which arises more than a week after segregation of the trophoblast lineage [26]. Furthermore, there remains significant debate whether BMP4 treatment of primed hPSCs may give rise to amnion and mesoderm fates in addition to trophoblast [27–31]. We and others have shown that naïve hPSCs, which display transcriptional and epigenetic properties of the pre-implantation embryo, can directly differentiate into self-renewing and bipotent hTSCs [32–35]. In addition, somatic cells can be reprogrammed into human induced TSCs (hiTSCs) by overexpression of defined transcription factors [34, 36]. Here, we review the key properties that define human trophoblasts, the isolation of hTSCs from human blastocysts and placental tissues, and recently developed methods to derive hTSCs from pluripotent and somatic cell sources.

**Biological and molecular criteria for bona fide human trophoblast stem cells**

The recent years have seen a surge of interest in methods for deriving trophoblast cells from human stem cell sources. How do we assess the trophoblast identity of these cells? Which trophoblast model is most suitable for investigating placental development or disease processes? These questions are not simply answered and the choice of one trophoblast model versus another requires diligent considerations [37].

Lee et al. proposed four criteria for identification of primary first-trimester CTBs [38]. These criteria include: (i) the expression of GATA3, TFAP2C, and KRT7 at the protein level in mononuclear CTBs; (ii) hypomethylation of the ELF5 promoter region; (iii) expression of microRNAs (miRNAs) from the imprinted chromosome 19 miRNA cluster (C19MC), which is almost exclusively expressed in the placenta [39]; and (iv) the absence of classical HLA receptors (HLA-A, -B, -C). These criteria were established by comparing isolated primary first-trimester CTBs with villous stroma (extraembryonic mesoderm-derived), choriocarcinoma cell lines, primed hESCs, trophoblast-like cells generated from primed hESCs, and an embryonic carcinoma cell line. All four of these trophoblast features were identified in villous CTBs and choriocarcinoma cells. On the other hand, primed hESCs, their trophoblast derivatives, and embryonic carcinoma cells exhibited only some of these features, while villous stroma exhibited none of them.

Although the four criteria proposed by Lee et al. are characteristic of villous CTBs, it remains unclear whether they also apply to the pre-villous post-implantation stage, which is extraordinarily difficult to access. Additional criteria are
needed to stage-match CTBs across different timepoints of human post-implantation development. In addition, specialized trophoblast cell types have their own unique molecular properties and recent studies are just beginning to illuminate the complex interactions between EVTs and maternal immune cells, how they invade uterine glandular spaces, and the possibility that placental bed giant cells may arise by fusion of deeply invasive interstitial EVTs [40, 41]. To define *bona fide* hTSCs, we propose that the following additional features need to be considered as well: (i) the ability to undergo lineage-directed differentiation into EVT and STB lineages, which should include functional characterization of differentiated trophoblast cell types based on the hormone-producing syncytia and invasive potential; (ii) the capacity of hTSCs for long-term self-renewal beyond a certain number of replication cycles, termed the Hayflick limit [42], which defines a truly proliferative stem cell; (iii) rigorous transcriptional benchmarking to human trophoblast identities in vivo; and (iv) whenever possible, the use of primary tissue sections for morphological and molecular comparison, e.g. to assess the size of syncytia and marker expression.

The recent availability of single cell RNA-sequencing (scRNA-seq) datasets for several stages of early human development, including pre-implantation [43, 44], post-implantation [45, 46], and villous placental tissues [47–49] provides stringent criteria for evaluating trophoblast fate (see the companion review in this issue by Brian Cox). Still missing, however, is a comprehensive scRNA-seq analysis that traces human trophoblast development from the pre-implantation TE to post-implantation CTB and subsequent stages of placental development. This would help to address questions regarding the hierarchical relationship between trophoblast subpopulations leading to the emergence of the placenta.

**Derivation of human trophoblast stem cells from blastocysts and placental tissues**

The discovery of in vitro culture conditions for establishing authentic human trophoblast stem cells (hTSCs) has transformed our ability to model placental development [21]. To identify pathways required for hTSC derivation, Okae et al. performed RNA-seq analysis on CTBs, EVTs, and STBs isolated from first-trimester placental tissues. Since the Wingless/Integrated (WNT) and epidermal growth factor (EGF) signal transduction pathways were enriched in CTBs, they first tried to culture CTBs in the presence of the GSK3 inhibitor CHIR99021, which activates WNT signaling, and recombinant EGF. Because the cells did not adhere well, they screened additional inhibitors and growth factors that promote the expansion of epithelial stem cells, eventually arriving at a cocktail comprising the transforming growth factor β (TGF-β) inhibitors SB431542 and A83-01, the histone deacetylase (HDAC) inhibitor valproic acid, recombinant EGF, CHIR99021, and the Rho-associated kinase (ROCK) inhibitor Y27632 (termed SAVECY medium). Under these conditions, they were able to derive putative hTSCs from both primary first-trimester CTBs and TE outgrowths of attached blastocysts. These cells could be propagated for over 70 passages, and expressed trophoblast markers, such as GATA3, TP63, TEAD4, and C19MC miRNAs.

Okae et al. then examined whether their candidate hTSCs were capable of differentiating into functional placental cell types. Upon removal of the WNT-promoting factor, CHIR99021, they noticed that the cells differentiated into HLA-G-positive EVT-like cells. To further optimize their protocol for lineage-directed EVT differentiation, they also increased the concentration A83-01, provided an extracellular matrix (ECM) in the form of Matrigel, and added Neuregulin (NRG1), all of which have been shown to promote EVT differentiation in placental explant cultures [50–52]. These hTSC-derived EVTs completed key aspects of EMT and expansion of all trophoblast cell types, although migratory EVT-like cells were few in number.

Transcriptional analysis of hTSC lines derived from blastocysts and CTBs revealed strong similarities between each other and to primary CTBs [21]. Likewise, hTSC-derived EVTs and STBs most closely resembled their respective in vivo counterparts. Okae et al. then used whole genome bisulfite sequencing (WGBS) to assess the DNA methylation landscape of hTSCs in relation to first-trimester CTBs. Overall, hTSCs were hypomethylated compared to primary CTBs (33.6% and 52.3%, respectively), but critical patterns of methylation were conserved, such as the retention of placenta-specific germline differentially methylated regions (gDMRs) [54] and the presence of large partially methylated domains [55]. A potential explanation for the apparent hypomethylation of in vitro derived hTSCs is that these culture conditions may enrich for proliferative CTBs, which tend to be less methylated than other CTB populations [56]. Specific examples of placenta-specific hypomethylation observed
in hTSCs include the promoter regions of ELF5, INSL4, ZNF750, and DSCR4. Finally, hTSCs also maintained the expected intermediate methylation levels at gDMRs associated with placenta-specific imprinted genes. Altogether, these data suggest that the conditions for hTSC isolation developed by the Arima laboratory capture a reliable in vitro counterpart of human first-trimester CTBs.

The original report from Okae et al. stated that they were unable to derive hTSCs from term placenta. This suggested that bipotent proliferative CTBs, which are the likely source of hTSCs, may be lost during or after the second trimester of pregnancy. However, two recent reports indicate that it is possible to derive hTSCs from term placenta using modified protocols. Kessler and colleagues reported that non-integrating viral expression of five transcription factors (CDX2, ELF5, ETS2, TFAP2C, and TEAD4) can reprogram term villous CTBs into induced TSCs (iTSCs), which are capable of long-term self-renewal and display transcriptional similarity to hTSCs derived from first-trimester placental tissues [57]. In addition, Chen and colleagues reported that treatment with SAVECY media under hypoxic conditions facilitates the derivation of hTSCs from term placenta by targeting the GCM1-ΔNp63α antagonistic signaling axis [58]. The establishment of hTSC lines from term placenta enables the generation of trophoblast models from placental tissues with known pregnancy outcomes, but it is important to remember that term placenta is inherently programmed to stop functioning after 9 months due to accumulating DNA methylation and damage from reactive oxygen species, a decline in CTB proliferation, and fluctuations in placental hormone secretion during pregnancy and parturition [59–62]. Because of these accumulated phenomena over gestation, determining the cause of placental dysfunction is difficult without access to earlier stages of placental development.

**Generation of human trophoblast stem cells from naïve pluripotent stem cells**

Given limited access to human embryos, stem cell models have become an invaluable tool to study early human development. Self-renewing hESC lines were first derived from pluripotent cells in human blastocysts in 1998 [63]. However, experimentally observed molecular differences between mouse and human pluripotent stem cells raised the question whether there may be multiple pluripotent states. It is now well described that pluripotency exists in a continuum from the initial, preimplantation naïve pluripotent state to the primed pluripotent state, the last step before gastrulation [25]. These discrete hPSC states are isolated using different culture media. Whereas conventional hPSCs are commonly maintained in commercially available media containing high levels of FGF and Activin, naïve hPSCs can be established using small molecule inhibitor cocktails. In particular, the t2i/L/Gö [64] and 5i/L/A [65] cocktails, have been widely used to induce molecular signatures of naïve pluripotency in hPSCs, including transcriptional correspondence to the EPI compartment of the blastocyst [26, 66, 67], X chromosome reactivation in female cells [68, 69], and expression of blastocyst-specific cell surface markers [70, 71]. Detailed reviews on methods for inducing and maintaining naïve hPSCs and their properties are available elsewhere [72–75]. In addition, naïve hPSCs can also be reprogrammed directly from somatic cells [76–80] or derived de novo from human blastocysts [81, 82].

The TE vs. inner cell mass (ICM) cell fate decision is the first to occur in the mammalian embryo and is viewed as a strong barrier in mouse. Indeed, pluripotent cells in the mouse blastocyst have lost the ability to give rise to TE and mouse PSCs require genetic manipulation, such as overexpression of Cdx2 or downregulation of Oct4, to acquire TE fate [83, 84]. However, several lines of evidence suggested that naïve hPSCs may have an expanded fate potential compared to their murine counterparts. First, scRNA-seq profiling of human embryos has revealed more fluid lineage segregation compared to mouse embryos. The TE program is initiated by compaction in the human morula [85], but only becomes transcriptionally distinct from ICM cells 12 h later at the B2 blastocyst stage [86, 87]. Cdx2 expression is acquired in TE cells at the B3 stage before it is specifically lost in polar TE cells, which subsequently gain NR2F2. This stage of blastocyst development also coincides with onset of PrE markers [86, 88]. Although these observations still require validation by time-lapse staging, overall lineage segregation appears to be more fluid than the paced, stepwise lineage segregation seen in mouse embryos [89–91]. This change of pace could explain retention of plasticity in human blastocyst lineages, as demonstrated by the ability of TE cells to form ICM upon re-aggregation [92]. Since naïve hPSCs correspond to the early human blastocyst based on the gene and transposon expression profiling [67, 69], they may conceivably retain developmental plasticity associated with this stage of human embryogenesis.

Second, naïve hPSCs exhibit elevated expression of a subset of transcription factors and open chromatin sites associated with the human trophoblast lineage [69, 93], which raised the question whether they may have an enhanced potential for trophoblast differentiation. Indeed, we and others demonstrated that naïve cells can be directly converted into cells that closely resemble hTSCs upon application of the SAVECY media developed by Okae et al. [32–35] (Fig. 1). Some methodological differences were reported in these studies: the David and Pastor laboratories used naïve cells maintained in t2iLGöY [64], while Dong et al. primarily used naïve cells derived in 5i/L/A [65]. In addition, the David and Theunissen laboratories passaged the cells until...
homogeneous hTSC cultures were obtained [32, 34], while the Pastor laboratory performed FACS sorting based on the EpCAM and ITGA2 [33]. However, in all cases, cells derived from naïve hPSCs closely resemble primary hTSCs by morphology and surface marker profile.

The biological and molecular properties of naïve hPSC-derived hTSCs were evaluated by using a number of different assays. Bulk RNA-seq analyses revealed close transcriptional correspondence between naïve hPSC-derived hTSCs and primary hTSCs derived from blastocysts or first-trimester placental tissues [21]. The resulting hTSCs correspond to human post-implantation CTB-like cells [32–34].

Fig. 1 Derivation of human trophoblast stem cells (hTSCs) from naïve human pluripotent stem cells (hPSCs). Naïve hPSCs can be directly converted into hTSCs by treatment with SAVECY media, which were originally developed to isolate primary hTSCs from blastocysts and first-trimester placental tissues [21]. The resulting hTSCs correspond to human post-implantation CTB-like cells [32–34].

including ZFAT and PROSER-A1, resisted demethylation both in the naïve hPSC state and upon differentiation into hTSCs. Importantly, all three studies also evaluated the differentiation potential of naïve hPSC-derived hTSCs towards specialized trophoblast fates, demonstrating that the cells were capable of giving rise to invasive EVTs and hormone-producing STBs using the methods for lineage-directed hTSC differentiation described by Okae et al. [32–34].

Based on the alignment with single cell expression data from human embryos, naïve hPSC-derived hTSCs most closely correspond to post-implantation trophoblast cells around 10–12 dpf [32, 34]. This suggests that the hTSC conditions developed by Okae et al. promote the differentiation of naïve hPSCs into a post-implantation CTB-like state and that it may be possible to capture a pre-implantation TE-like state from naïve cells under appropriate culture conditions. Indeed, recent work from the Smith laboratory demonstrated that naïve hPSCs transiently acquire a pre-implantation TE identity upon treatment with the MEK inhibitor PD0325901 and the TGFβ inhibitor A83-01 (PDA83) before the cells are switched to SAVECY media to attain a post-implantation CTB-like state [30] (Fig. 2). This work was independently corroborated by Takashima and colleagues, who supplemented the PDA83 cocktail with BMP4 and a Janus kinase (JAK) inhibitor [29] (Fig. 2). These naïve-like TE cells share common trophoblast signatures with post-implantation hTSCs, such as expression of GATA3, TFAP2C, and KRT7, demethylation of the ELF5 promoter region, and expression of C19MC miRNAs, but show increased expression of early TE markers, such as CDX2 and HAVCR1. Naïve TE-like cells can be distinguished by a unique cell surface profile, including expression of ENPEP and TACSTD2, which are also expressed in TE in human pre-implantation blastocysts, but reduced levels of HLA-ABC and SIGLEC6 as compared to hTSCs [94]. The absence of classical HLA molecules is one of the hallmarks of human trophoblast cells as proposed by Lee et al. [38], and therefore the increased expression of these antigens in hTSCs was unexpected. Takashima et al. proposed a simplified formulation for promoting the transition from naïve TE-like cells into a post-implantation CTB-like state by culture in the presence of A83-01, the GSK3 inhibitor CHIR99021, and EGF (ACE), which prevents aberrant HLA-ABC activation [94]. These conditions also support the derivation of hTSCs from primary chorionic villi of first-trimester placental tissues. By benchmarking their expression data to single cell studies in human and monkey embryos, these papers showed that the differentiation of naïve hPSCs into hTSCs via a pre-implantation TE intermediate recapitulates key steps during primate trophoblast development, offering an accessible model system of human trophoblast specification. Furthermore, the Smith laboratory demonstrated that EPI cells from human blastocysts harbor the intrinsic potential to generate TE [30], which reveals that
the human EPI retains an expanded extraembryonic plasticity as compared to the mouse EPI, in accordance with in vitro studies using naïve hPSCs. In addition to providing a source of 2D models of trophoblast development, naïve hPSCs can also be used to generate 3D organoids that encompass a diversity of trophoblast cell types. The isolation of trophoblast organoids was first reported by the Moffett and Knöfler laboratories in 2018 \[95, 96\], but required the isolation of CTBs from first-trimester placental tissue, which is not readily available in many jurisdictions due to ethical and practical restrictions. Given the transcriptional and epigenetic similarities between hTSCs and CTBs, we and others postulated that it should be possible to generate trophoblast organoids from naïve hPSCs via an hTSC intermediate. Indeed, when transferred to Matrigel droplets in the presence of trophoblast organoid medium (TOM), naïve hPSC-derived hTSCs self-organize into 3D organoids that exhibit a similar architecture as primary trophoblast organoids with an outer CTB layer and an inner STB compartment \[29, 34, 97\]. This marks a reversal of the architecture of the primary placental villi in which STBs are located towards the periphery. We recently performed single cell transcriptome profiling on stem-cell-derived trophoblast organoids (SC-TOs) generated from various hTSC lines, which revealed a highly concordant cellular distribution of progenitor and specialized trophoblast states \[97\]. This suggests that trophoblast organoid culture represents a powerful attractor state in which the influence of subtle epigenetic differences between hTSCs obtained from naïve hPSCs and those obtained from first-trimester placental tissues is mitigated. In addition, the proportion of EVTs can be enhanced by transfer to a WNT-reduced environment, similar to observations in primary trophoblast organoids \[98\]. We further showed that SC-TOs generated from naïve hPSCs recapitulate placental X chromosome inactivation patterns and model early placental susceptibility to emerging pathogens, including SARS-CoV-2 and Zika virus \[97\]. Like hTSCs in 2D culture, the CTB subpopulation within SC-TOs is most closely aligned with trophoblast progenitors in early post-implantation embryos and consequently these two culture systems exhibit many shared markers, such as GATA3, TFAP2C, KRT7, and C19MC miRNAs. However, the reduced expression of classical HLA molecules in trophoblast organoids and their 3D microenvironment present advantages for modeling placental organogenesis.

Is it possible to derive human trophoblast stem cells from primed and other pluripotent states?

The initial studies that directly compared the response of naïve and primed hPSCs to hTSC media all concluded that primed hPSCs fail to acquire an hTSC-like identity when directly transferred to the SAVECY media developed by Okae et al. \[32–34\]. In fact, primed hPSCs showed upregulation of neuroectoderm-related genes as opposed to trophoblast-related genes under these conditions \[32\]. Nevertheless, several other studies have reported that primed hPSCs are capable of generating hTSC-like cells under modified culture conditions. The generation of hTSCs from primed hPSCs was reported via micromesh culture \[99\] or by pre-treatment with BMP4, a combination of BMP4 and the WNT inhibitor IWP2, or the TGFβ inhibitor A83-01 before applying hTSC media (Fig. 3) \[100–104\]. The resulting cells share several global features with primary hTSCs, including gene expression, differentiation potential towards EVT and STB lineages, and increased chromatin accessibility at trophoblast-specific genes. These data indicate that it is possible to direct primed hPSCs towards an hTSC-like state. However, there may yet be subtle but significant differences between hTSC-like cells derived from naïve and primed states. For example, a recent study from the Arima laboratory reported that hTSCs derived from naïve, but not...
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primed, hPSCs display complete activation of placenta-imprinted C19MC miRNAs [105], which is one of the original trophoblast criteria proposed by Lee et al. [38]. Ectopic activation of C19MC miRNAs in primed hPSCs enhanced their potential to differentiate into proliferative hTSCs with full differentiation potential towards specialized trophoblast lineages [105]. Consistent with these findings, incomplete activation of C19MC miRNAs was also observed in two other studies that derived hTSCs from primed hPSCs [101, 103]. Therefore, an important objective in future studies will be to further compare the biological and molecular properties of hTSC-like cells isolated from isogenic naïve and primed states side-by-side.

A complicating factor in these studies is that trophoblast cells have substantially overlapping transcriptional signatures with amnion, which is another extraembryonic lineage that arises from the primate EPI following implantation [106]. Both the Pastor and Smith laboratories noted upregulation of amnion markers in hTSC-like cells generated from extended pluripotent stem cells (EPSCs), which lack clear alignment with a developmental equivalent in vivo [33, 107, 108]. On the other hand, the David laboratory reported that hTSCs can be derived from a different type of EPSC following extended culture in SAVECY media [34]. As noted earlier, there also remains a significant debate whether BMP4 treatment of primed hPSCs may give rise to amnion [27–30]. Therefore, careful analysis of amnion and trophoblast markers will be essential when assessing the quality of putative hTSCs and a recent scRNA-seq analysis of amnion development in primate embryos provides a helpful set of distinctive markers [109]. Adding further complexity, this study also demonstrated that early amnion differentiation from EPI cells follows a TE-like trajectory, including the activation of a broad range of STB-associated markers. Consequently, it may be instructive to reassess whether primed hPSCs or EPSCs acquire an amnion fate before being redirected towards an hTSC-like identity.

**Direct reprogramming of human somatic cells into induced trophoblast stem cells**

The derivation of hTSCs offers an in vitro model system of placental organogenesis and dysfunction during differentiation of villous CTB progenitors into specialized trophoblast cell types. However, human stem cell models are limited in terms of their genetic background, since they are mainly generated from embryos, or in the case of hTSCs, first-trimester placental tissues [21]. The David and Polo laboratories have developed methods to directly reprogram patient samples into an array of stem cells. Their initial focus was to directly reprogram somatic cells into naïve iPSCs, assessing multiple culture conditions in parallel. They showed that somatic cells could be directly reprogrammed using the canonical OCT4, SOX2, KLF4, and c-Myc (OSKM) transcription factor cocktail and two naïve-specific culture media, t2iLGö and 5iLAF, yielding cells that closely corresponded to the human pre-implantation EPI based on the transcriptional and epigenetic criteria [76, 77].

The David laboratory noticed that cells co-expressing the EPI marker NANOG and the TE marker GATA3 emerged transiently at an early stage of naïve reprogramming in reprogramming intermediates that retained high transgene expression levels [34]. The Polo laboratory came to the same conclusion by mapping the reprogramming routes from somatic cells to primed and naïve iPSCs [36]. These observations were concomitant with the publication of SAVECY media that support hTSC derivation and growth [21], prompting both laboratories to fine tune their reprogramming conditions to generate hiTSCs. Technically, the protocols established by both groups are similar, with minor differences (described in soon-to-be-published Nature Protocol papers). Both groups demonstrated that the hiTSCs were *bona fide* hTSCs: they reported in vivo differentiation into trophoblast-like tissues, the ability to differentiate into hCG-secreting syncytia and EVTs, and high expression levels of C19MC miRNAs as compared to both fibroblasts and iPSCs,
a unique feature of primary trophoblast [38]. Building on their scRNA-seq analysis, Liu and colleagues [36] identified GATA2 and TFAP2C as important regulators during somatic cell reprogramming into naïve iPSCs. Indeed, naïve reprogramming was greatly impaired when TFAP2C was knocked down, while reprogramming to primed pluripotency was less impacted. Moreover, GATA2 knockdown resulted in impaired reprogramming to both naïve and primed states, suggesting an earlier role of GATA2 during reprogramming. Further analysis of the scRNA-seq dataset revealed that during naïve reprogramming under both 5iLAF and t2iLGö conditions some subpopulations of cells were enriched in TE signatures (Fig. 4). Building on their expertise in human peri-implantation development [86], Castel and colleagues matched hiTSCs with 8 dpf trophoblast cells, as demonstrated by precise gene-set enrichment of developmental stages and notably the presence of NR2F2, a transcription factor that is expressed upon progression of CDX2-positive TE cells into CDX2-negative trophoblast cells [88].

**Concluding remarks**

Since the elucidation of culture conditions for derivation of self-renewing hTSCs from blastocysts and first-trimester placental tissues by Okae et al. [21], a number of groups have explored alternative avenues for accessing the hTSC state. Here we reviewed three sources for creating hTSCs, each of which offers their own relative advantages and disadvantages for basic and applied research.

First, the derivation of hTSCs from blastocysts and first-trimester placental tissues continues to provide a gold standard for the field, although ethical, legal, and practical constraints on accessing such tissues present a barrier for many researchers. The recent success in deriving hTSCs from placental tissues at term may offer a more accessible option for deriving primary hTSCs [57, 58]. Second, the generation of hTSCs from naïve hPSCs offers a renewable source of hTSCs and 3D trophoblast organoids that can be applied to a broad spectrum of hPSC lines, including patient-specific iPSCs [29, 30, 32–34, 97]. Furthermore, the ability to reconstitute the trophoblast lineage from naïve hPSCs, which correspond to pluripotent cells in the pre-implantation embryo, presents exciting possibilities to investigate the genetic and epigenetic mechanisms of human trophoblast specification in vitro. A potential limitation of the use of naïve hPSCs is the fact that they undergo erasure of parent-specific imprinting during extended culture [69, 110]. Nevertheless, most placenta-imprinted genes are activated during the naïve-to-hTSC transition [33, 97]. Third, direct reprogramming of somatic cells into hiTSCs presents a more efficient route towards the creation of patient-specific hTSCs for modeling placental diseases but may be less suitable for modeling the process of trophoblast specification during human pre-implantation development.

Several recent studies indicate that it is possible to generate hTSC-like cells directly from primed hPSCs under modified culture conditions. A short pulse of BMP4, BMP4 and IWP2, or A83-01 can redirect primed hESCs away from a neural fate and towards a trophoblast identity [99–104]. However, there may be subtle but significant differences between hTSC-like cells derived from naïve and primed cells, such as the expression level of placenta-imprinted C19 miRNAs [105], which is one of the original trophoblast criteria proposed by Lee et al. [38]. It is important to bear in mind that extensive single cell expression profiling of primate embryogenesis has not revealed evidence that post-implantation EPI cells contribute to the trophoblast lineage [26, 45, 111]. Therefore, we surmise that the generation of hTSC-like cells from primed hPSCs may represent a culture-induced trans-differentiation event. According to the recent work from Rugg-Gunn and
colleagues, primate EPI cells coopt the transcriptional program of TE specification to initiate the amniotic cavity just after implantation [109]. Cooptation of the pre-existing TE differentiation program during the early wave of amnion differentiation may present a window of opportunity for a subset of EPI cells to be redirected towards an hTSC-like identity under pressure from strong external stimuli. It has also been suggested that amnion may provide an independent source of STB-like cells [112], but this hypothesis requires validation using lineage tracing.

The SAVECY media developed by the Arima laboratory have had a transformative impact on the field of trophoblast biology, but the culture conditions for hTSC derivation and differentiation require further refinement. Current hTSCs are hypomethylated and exhibit increased expression of HLA class I surface molecules compared to CTBs in vivo [21]. Furthermore, it remains unclear why hTSC derivation requires the use of two distinct TGFβ inhibitors, A83-01 and SB435412. In fact, the ACE formulation developed by the Takashima laboratory uses only one of these inhibitors and yields hTSCs with reduced HLA-ABC expression [94], as does transfer to trophoblast organoid media [37, 97]. Additional refinements to these conditions and those used for lineage-directed differentiation may enable the isolation of hTSCs that more readily transition into functional EVT and STB cells. Alongside this effort, a more complete in vivo reference for TE and CTB progenitors is needed to stage-match hTSCs derived under various conditions to their counterparts in the human embryo and placenta. New embryo models, such as blastoids [88, 113, 114], offer an opportunity to better understand TE fate progression from its initiation at the morula stage [85] until the appearance of the first STB and EVT cells [45].

While the culture conditions developed by Okae et al. capture hTSCs in a post-implantation trophoblast identity, a pre-implantation TE state can be transiently accessed by treating naïve hPSCs with MEK and TGFβ inhibitors in monolayer culture [29, 30] or by promoting their self-organization into blastoids. An important question for future research will be to investigate whether culture conditions can be devised to capture a self-renewing pre-implantation TE state in human cells. Mouse TSCs, in fact, more closely resemble the pre-implantation TE at the level of marker expression and based on their ability to colonize the placenta following injection into E3.5 mouse embryos [17], although significant heterogeneity has been reported within mouse TSC culture [115, 116]. Rivron and colleagues recently reported progress in capturing mouse TSCs with more specific features of pre-implantation polar TE by applying inductive signals originating from the inner embryonic cells of the blastocyst [117]. Conceivably, a similar approach may be effective for stabilizing a self-renewing pre-implantation TE state from naïve hPSCs. As an alternative approach, Mischler et al. reported that primed hPSCs can give rise to CDX2-positive hTSC-like cells in the presence of a spherogosine-1 phosphate agonist, a GSK3 inhibitor, a TGFβ inhibitor, and recombinant FGF10 [100], but it remains to be determined whether these cells correspond to TE cells in the human pre-implantation embryo.

What are the potential biomedical applications of hTSCs derived from pluripotent and somatic sources? The development of the placenta remains an understudied aspect of human embryology, but placental complications during the first trimester are associated with pregnancy complications such as preeclampsia, miscarriage, and fetal growth restriction [118]. In addition, there is increasing recognition that variations in the supply of nutrients to the developing fetus can manifest in disease during postnatal life, such as cardiovascular disease [119]. Therefore, it would be instructive to generate hTSCs from patients who suffered pre-eclampsia or fetal growth restriction using the various approaches discussed in this review and rigorously evaluate their phenotype relative to healthy controls based on the molecular profiling (transcriptome, DNA methylome, and chromatin accessibility) and differentiation towards specialized trophoblast fates using 2D monolayer and 3D organoid culture. More advanced phenotyping could involve the use of co-culture assays between trophoblasts and human endometrial cells to model the process of trophoblast implantation and invasion, as recently demonstrated using blastoids and stem-cell-derived trophoblast organoids [88, 97]. These studies could identify molecular targets for early detection and potential pharmacological intervention. In addition, an improved understanding of the interplay between placental genes and endometrial environmental cues that are essential for hTSC specification and differentiation may aid in understanding the genetic and environmental basis of placental pathologies leading to fetal growth restriction, preterm birth, and recurrent pregnancy loss [120–122], while genes that suppress the proliferation and invasion of hTSCs may provide candidates for treatment of choriocarcinoma, a highly malignant tumor of trophoblastic origin [123]. From this standpoint, recent candidate-based and genome-wide approaches to identify essential and growth-restricting genes in hTSCs present an important foundation for future studies [124–128]. We anticipate that the ensuing decade will witness unprecedented advances in our understanding of placental development and women’s health through the concerted efforts of reproductive scientists, stem cell biologists, clinicians, and biomedical engineers.

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