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# 1 Origin and segregation of the human germline

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#### 2

#### 28 Abstract

#### 29

Human germline-soma segregation occurs during weeks 2-3 in gastrulating embryos. While direct 30 31 studies are hindered, here we investigate the dynamics of human primordial germ cell (PGCs) 32 specification using in vitro models with temporally resolved single-cell transcriptomics and in-depth 33 characterisation to in vivo datasets from human and non-human primates, including a 3D marmoset 34 reference atlas. We elucidate the molecular signature for the transient gain of competence for germ cell 35 fate during peri-implantation epiblast development. Further, we show that both the PGCs and amnion 36 arise from transcriptionally similar TFAP2A positive progenitors at the posterior end of the embryo. 37 Notably, genetic loss of function experiments show that TFAP2A is crucial for initiating the PGC fate 38 without detectably affecting the amnion, and its subsequently replaced by TFAP2C as an essential 39 component of the genetic network for PGC fate. Accordingly, amniotic cells continue to emerge from 40 the progenitors in the posterior epiblast, but importantly, this is also a source of nascent PGCs.

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# 42 Introduction

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Human primordial germ cells (PGCs) are among the first lineages to emerge in the developing
gastrulating peri-implantation embryo at weeks (Wks) 2-3, eventually developing into sperm or eggs.
The parental gametes generate the totipotent state at fertilisation and transmit genetic and epigenetic
information necessary for development.

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49 The specification of PGCs is linked with the initiation of the unique germ cell transcriptomic and 50 epigenetic program. Aberrant specification and development of germ cells can lead to sterility, germ-51 cell-derived cancers, and other human diseases with long term consequences across generations. Ethical 52 and technical reasons restrict direct studies on nascent human PGCs, necessitating in vitro models, 53 which are, however, experimentally tractable for mechanistic insights (Hirate et al., 2013; Irie et al., 54 2015; Sasaki et al., 2015; Tang et al., 2016; Kobayashi et al., 2017; Irie, Sybirna and Surani, 2018). 55 Due to the in vitro nature of these models, comprehensive comparisons with rare human embryos and 56 animal proxies, including in vivo development of non-human primates such as marmosets can be 57 significantly informative for germline biology.

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59 The induction of PGC-competent cells from cultured pluripotent stem cells (PSCs) is possible using

60 self-renewing or transient pre-mesendoderm (PreME) populations (Irie *et al.*, 2015; Sasaki *et al.*, 2015;

61 Kobayashi *et al.*, 2017). The timing and regulation of the transient state of competence for PGC-fate in

62 human embryos are not yet fully defined but likely determine the number of founder PGCs in vivo. If

63 aggregated into 3D embryoid bodies, these competent cells give rise to 10 - 40% PGC-like cells

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64 (PGCLCs) in response to BMP and other cytokines (Irie *et al.*, 2015; Sasaki *et al.*, 2015; Kobayashi *et al.*, 2017). The remaining cells adopt somatic fates, but their relationship with the emerging PGCLCs remains unclear (Irie *et al.*, 2015; Sasaki *et al.*, 2015; Kobayashi *et al.*, 2017). Defining the characteristics of the somatic lineages in embryoid bodies may help identify soma-PGC interactions and reveal the context of how PGCs form in experimental models concerning the lineages in the embryo.

In vitro models identified SOX17, PRDM1, and TFAP2C as the core regulators of human PGC fate (Irie *et al.*, 2015; Kobayashi *et al.*, 2017; Kojima *et al.*, 2017; Tang *et al.*, 2022). This tripartite network for
PGC fate has also been observed in vivo in other species that develop as a bilaminar disc, including
cynomolgus, marmoset, rabbit, and pig (Sasaki *et al.*, 2016; Sybirna, Wong and Surani, 2019; Alberio,
Kobayashi and Surani, 2021; Kobayashi *et al.*, 2021; Zhu *et al.*, 2021; Bergmann *et al.*, 2022).

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On the other hand, *Sox17* is not a critical regulator of PGC specification in rodents where the embryos
develop as egg cylinders (Kanai-Azuma *et al.*, 2002). Notably, when *SOX17* is the critical regulator for
PGC specification as in humans and non-human primates, there is concomitant repression of SOX2, but

not in mice, where *Sox2* has a crucial role in PGC development (Campolo *et al.*, 2013).

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81 The site of human PGC specification remains unclear. In cynomolgus and marmosets, PGCs are first 82 observed in the amnion prior to gastrulation (Sasaki et al., 2016; Bergmann et al., 2022). At later stages, 83 PGCs are detected in the posterior epiblast, with the possibility of a dual origin (Sasaki et al., 2016; 84 Kobayashi et al., 2017; Kobayashi and Surani, 2018). Note that in humans and non-human primates, 85 the nascent amnion is among the first lineages to form from the epiblast (Bergmann et al., no date; 86 Xiang et al., 2019). In some non-primate embryos, including bilaminar disc forming species such as 87 rabbit and pig, PGC specification precedes amnion development (Alberio, Kobayashi and Surani, 2021; 88 Kobayashi et al., 2021; Zhu et al., 2021). In the pig, at least, PGCs arise from pre-primitive streak (PS) 89 and early-PS stage competent epiblast(Kobayashi et al., 2017), and in a rare Wk3 (Carnegie stage 7) 90 human embryo, PGCs are associated with the primitive streak (Tyser et al., 2021). Here we used our 91 PSC-based model for PGC specification (Irie et al., 2015; Kobayashi et al., 2017) in conjunction with 92 highly resolved single-cell transcriptome sequencing and integrative analysis with existing human and 93 primate datasets to document the nature of the somatic components of the models and provide the 94 context for PGCLC specification. Notably, we identified TFAP2A, considered an amnion marker 95 (Shao, Taniguchi, Gurdziel, et al., 2017; Shao, Taniguchi, Townshend, et al., 2017), as an essential and 96 thus far the earliest regulator of PGC fate. Loss of TFAP2A leads to an almost complete abrogation of 97 PGCLCs, in favour of a population of cells displaying SOX2 expression but no significant effect on 98 somatic lineages. The observations also provide insights into the likely origin of human PGCs. 99

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#### 101 **Results**

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# 103 A highly resolved transcriptional characterisation of PGC specification in embryoid 104 bodies

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106 Human pluripotent stem cells (PSCs) in a primed state represent non-gastrulating postimplantation 107 epiblast cells (Yu et al., 2021) with a low competence for PGCLC fate (<5%) (Irie et al., 2015). PSCs 108 can, however, acquire competence for PGC fate as self-renewing populations in media containing four 109 inhibitors (henceforth called 4i conditions) (Gafni et al., 2013; Irie et al., 2015). Secondly, in response 110 to WNT and Activin signalling, PSC can transiently acquire competence for PGC-fate at 12h, known 111 as pre-mesendodermal cells (henceforth called Pre-ME) (Kobayashi et al., 2017). Pre-ME progress to 112 mesendoderm (ME) fate at 24h when they lose competence for PGCLC specification and instead gain 113 competence for definitive endoderm (DE; 60-80%) and mesoderm fates (Fig. 1a).

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115 The efficiency of PGCLC induction ranges from  $\sim 10-40\%$  of cells in the embryoid body (EB), 116 depending on the cell line (Chen et al., 2017); the remaining non-PGCLCs cells acquire somatic fates. 117 Using our in vitro model (Kobayashi et al., 2017), we elucidate the transcriptional dynamics as the Pre-118 ME cells undergo specification to PGCLCs in response to BMP. To discern the changes in 119 transcriptional states, we analysed the embryoid body at the resolution of single cells using 10X 120 Genomics single-cell RNA-sequencing. We sampled EBs over a highly-resolve time series between 121 12h-96h post-induction with additional comparative samples of conventional PSCs, PGC-competent 122 populations (4i and PreME), DE and ME populations, and Wk7 human gonadal PGCs for in vivo 123 reference (Fig. 1a).

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125 We first sought to establish the identity of detectable lineages using droplet single cell RNA sequencing 126 in the embryoid bodies, which fell into 15 main clusters (Supplementary Fig. 1a). Pseudo-bulk 127 correlation showed a high degree of correlation between these clusters and amnion-like cells (AMLC), 128 primordial germ-cell like cells (PGCLCs), or mesoderm-like cells (MELCs) (Zheng et al., 2019) 129 (Supplementary Fig. 1b). Pseudo-bulk comparison with a human in vitro embryo culture (Xiang et al., 130 2019) and the in vivo CS7 human gastrula (Tyser et al., 2021) corroborates these observations, showing 131 a higher degree of correlation between EBs and embryonic disc or amnion but a substantially reduced 132 correlation with other extraembryonic-tissues and pre-implantation lineages (Supplementary Fig. 1c), 133 and a comparatively low correlation with human syncytiotrophoblast (SCT) and extravillous 134 trophoblast (EVT) (Vento-Tormo et al., 2018). Together these results suggest that in response to BMP, 135 EBs progress to lineages of the peri-gastrulation embryo but not the extraembryonic tissues except for 136 the amnion.

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- 138 We aligned our data to a comprehensive range of existing embryonic datasets to refine cell annotations 139 and create a human primate gastrulation and PGC atlas (Fig. 1b). We included embryonic and amniotic 140 lineages from human and cynomolgus in vitro cultured embryos (Ma et al., 2019; Xiang et al., 2019; 141 Zhou et al., 2019), in vivo human and marmoset gastrula (Bergmann et al., no date; Tyser et al., 2021), 142 and human gonadal primordial germ cells (Guo et al., 2015; Li et al., 2017). We also include three in 143 vitro models of human PGCLC induction based on the microfluidic amnion model (Zheng et al., 2019), 144 micropatterned gastruloids (Minn et al., 2020) and embryoid bodies from two other cell lines(Chen et 145 al., 2019) (Supplementary table 1). We show a representation of our aligned dataset as a 2D UMAP 146 projection in Fig. 1c, with cells coloured by sampling time. For comparison, we also provide aligned 147 samples from the human gastrula dataset (Fig. 1d), with the remaining datasets shown in Supplementary 148 Fig. 1d-1j. Clustering across all the datasets grouped cells into approximately 30 clusters, with the key 149 clusters visualised for our data in Fig. 1e. Initial assessment suggests a low number of doublets 150 throughout (Supplementary Fig. 1k).
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A heatmap of gene expression of relevant lineage markers confirms the presence of a primitive streaklike population (cluster 1), as well as mesoderm-like cells (MELCs) (cluster 2-3), definitive endoderm-

- 154 like cells (DELCs) (cluster 11), amnion-like cells (AMLCs) (cluster 4, 7-10), and PGC-like cells
- 155 (PGCLCs) (cluster 5-6) within the EBs (Fig. 1f). We show key differentially expressed transcription
- 156 factors during the formation of individual cell types in Supplementary Fig. 2. To visualise expression 157 heterogeneity, we depict gene expression of six key lineage markers that, in combination, can be used
- to identify the cell fates in the EB (Fig. 1g); these findings were also confirmed at the protein level by immunofluorescence (IF) staining (Fig. 1h). Notably, our detailed integrated roadmap and characterisation show that at early stages, embryoid bodies contain subpopulations with molecular
- 161 signatures similar to the PS, with cells at later time points showing transcriptional profiles associated
- 162 with embryonic somatic fates (mesoderm and endoderm), PGCLCs, and amnion.
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## 164 **Detection of PGC competent population**

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166 Currently, there is no clear indication of what constitutes a PGC-competent population. We investigate
167 how the precursor PreME cells gain competence for PGC-fate to address this. We also analysed the
168 PGC-competent 4i cells against the non-competent populations (PSCs and ME) (Tang *et al.*, 2022).
169 Comparisons with existing datasets suggested that our PSCs are transcriptionally similar to other
170 PSCs(Chen *et al.*, 2019; Zheng *et al.*, 2019; Minn *et al.*, 2020) (Supplementary Fig. 1d-f) and align with

- 171 a subset of cells from in vitro cultured human embryos labelled as EmDisc (Xiang et al., 2019; Zhou et
- 172 al., 2019) (Supplementary Fig. 1g-h) and in vivo postimplantation epiblast (Tyser et al., 2021) (Fig. 2a-

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b). Conversely, PreME cells cluster with pluripotent embryonic disc sample (Xiang *et al.*, 2019; Tyser *et al.*, 2021) and cells labelled as epiblast and primitive streak and mesoderm in a human CS7 gastrula
(Tyser *et al.*, 2021) (Fig. 2c).

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177 Since the interpretation of distances in UMAP representations remains difficult (Chari, Banerjee and

178 Pachter, 2021), we also chose to visualise cells using diffusion maps (DM) to gauge the behaviour of

these precursor populations compared to non-competent PSCs, ME populations, and terminal stage

180 PGCLCs (Fig. 2d). These populations exist as a continuum of transcriptional states extending from

181 PSCs to ME, and diffusion components 2 (DC2) and DC3 with PGCLCs extending out along DC1 (Fig.

182 2e).

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184 Visualisation of the fraction of cell types in each subcluster identified a PSC-dominant (subcluster 1) 185 and a ME-dominant subpopulation (subcluster 22), with three other subpopulations (subclusters 3, 14, 186 and 19) comprised primarily of PGC-competent populations (4i and PreME) (Fig. 2f, Supplementary 187 Fig. 3a). Pairwise differential expression analyses of PreME cells in (competent) subcluster 3 and PSCs 188 in (non-competent) subcluster 1 identified several likely regulators of competence, including EOMES, 189 which has an identified role in PGC-competence (Chen et al., 2017; Kojima et al., 2021), and 190 mesodermal markers SP5 and MIXL1 (Fig. 2g). Additional pairwise comparison of the other competent-191 enriched subpopulations, e.g., subclusters 14 and 19, against cluster 1 identified similar markers, 192 including OTX2, SOX11, TERF1, TCF7L1, SALL2, LIN28A and TET1 (Supplementary Fig. 3b-f). 193 Comparison of competent clusters against cluster 22 showed further upregulation of mesoderm related 194 genes, MIXL1, GATA6, GSC, MESP1, ZIC2, EOMES in ME-dominated cluster and concomitant 195 reduction of pluripotency factor expression (SOX2, SOX3, NANOG) and MYC in PGC-competent 196 cluster (subcluster 3, 14, 19).

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198 Since competent subclusters 3, 14 and 19 showed similar marker expression (Supplementary Fig. 3b-199 f), we focused on subcluster 3 for simplicity (Fig. 2f). The signalling dynamics of competence gain and 200 loss were examined by expressing key genes between clusters representing PGC-competent and non-201 competent cells (Fig. 2f). We observed a progressive activation of NODAL and WNT signalling 202 together with the expression of BMP inhibitors, with the highest levels shown in the ME-dominated 203 subcluster (subcluster 22). BMP is the inductive signal for PGC-fate, and accordingly, the PGC-204 competent subcluster (subcluster 3) shows a reduced expression of BMP inhibitors CER1 ( $p < 2.17e^{-19}$ ) compared to the non-competent subcluster 22. Furthermore, we observed upregulation of NANOG (p < 205  $9.26e^{-48}/1.62e^{-6}$ ) and concomitant downregulation of OTX2 (p <  $6.48e^{-10}/0.045$ ) in subcluster 3 206 207 compared to non-competent clusters 1 and 22 respectively. Notably, OTX2 negatively regulates PGCLC 208 competence in mice (Zhang et al., 2018) and we recently found that OTX2 has a similar function in the 209 human germline (Tang et al., 2022).

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211 Together these analyses have identified molecular signatures that may underlie the transition from 212 primed pluripotency to a competent state for PGC fate.

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# 214 Specification of PGCLCs in EBs represents a primitive-streak-like stage

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Based on the expression of marker genes, EBs first transition through a primitive-streak-like stage before diversifying into mesoderm-like (MELC), definitive-endoderm-like (DELC), and primordial germ cell-like states, with the additional formation of amnion-like cells but with a notable lack of neural ectoderm populations (Supplementary Fig. 2e). Strikingly, these are lineages expected to arise at the posterior region of the developing embryo around the time of gastrulation.

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222 To test this hypothesis further, we sought to map cells found in vitro to existing spatial transcriptomics 223 datasets. Although spatially resolved human gastruloid datasets exist (Moris et al., 2020), these models 224 capture the onset of somitogenesis (CS9) and are therefore more developmentally advanced than our 225 model, which aligns well with data from CS5-7 embryos relevant to the emergence of PGCLCs. In this 226 regard, we note recent comprehensive spatially resolved transcriptional datasets of marmoset embryos 227 at CS5 and CS6 (Bergmann et al., 2022), where the peri-implantation development strongly resembles 228 that of human embryo development at the morphological and transcriptional level (Bergmann et al., 229 2022), including conserved expression of SOX17, PRDM1, TFAP2C and NANOS3 in PGCs. 230 Notwithstanding the differences in human and marmoset development timing, archival embryo 231 collections allow consistent staging between species based upon Carnegie staging (Strachan, Lindsay 232 and Wilson, 1997; O'Rahilly and Müller, 2010).

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234 To evaluate possible anterior-posterior bias, we mapped cells from our in vitro model to an existing 3D 235 spatially resolved depiction of a CS6 marmoset embryo in which laser capture microdissection was 236 used to generate a 3D spatially resolved transcriptome (see Materials and Methods). Together, they 237 capture the critical cell types for comparison (Fig. 3a) with gene expression patterns of critical markers 238 shown in Fig. 3b. We found that pluripotent stem cell populations mapped best to the anterior 239 compartment (Fig. 3c), in agreement with earlier studies (Tyser et al., 2021; Bergmann et al., 2022), 240 although we could not rule out that these cells might have a better mapping to earlier stages, e.g., CS4 241 bilaminar disc embryo since no data for this stage is available. We found that the PreME population 242 shifted towards the posterior end of the embryo, with amnion-like cells primarily mapping to the 243 posterior amnion (Fig. 3c). The basal cluster, which represents the 12h embryoid body mapped to the 244 posterior end of the embryonic disc to a region expression TBXT and other primitive streak markers. 245 Other cell lineages, including PGCLCs, showed an even stronger bias to the posterior end of the embryo,

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with PGCLC mapping to a distinct *NANOS3*-expressing region between the posterior-most embryonic
disc and amnion (Fig. 3a). Together, these results provide further evidence that our model represents
the development of the posterior end of the embryo during gastrulation and suggests ongoing
specification of both amnion and primordial germ cells.

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## 251 Highly resolved time series reveal dynamics of cell trajectories

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253 Having established the identity and spatial correspondence of key lineages, we next investigated the 254 dynamics of individual cell fate decisions within the EB. We performed a label transfer from the human 255 CS7 gastrula dataset (Tyser et al., 2021) to our data and separated EBs by collection time to visualise 256 the emergence of cell types (Fig. 4a). Twelve hours after inductive BMP cues, cells aligned primarily 257 to the primitive streak (PS) with a limited pool of epiblast-like cells. Primitive streak-like cells (PSLCs) persisted in limited numbers until ~24-32h, with sustained expression of NODAL (Supplementary Fig. 258 259 4). Nascent and emergent mesoderm-like cells (denoted nMELC, eMELC) appeared as early as 12h, 260 becoming more pronounced by 18h, with these lineages roughly corresponding to cluster 2. The earliest 261 PGCLCs arose around the 18h mark, with amnion-like cells and definitive endoderm-like cells around 262 24h.

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264 We visualised the segregation of early mesoderm from precursors with primitive-streak-like identity 265 using a diffusion map (Fig. 4b). Cells not committed to mesoderm-fate are instead predominately 266 directed towards PGCLC or AMLC. Both UMAP and DM representations suggest that PGCLCs and 267 AMLCs stem from highly similar progenitor cells (Fig. 4b). Interestingly, there remains some 268 association between the PGCLC and AMLC branches until around 48h, with a number of cells falling 269 between the two main branches. Visualisation of the PGCLC branch alongside samples from the CS7 270 human gastrula shows an overlap between the gastrula samples and our Wk7 in vivo PGCs and late in 271 vitro PGCLCs (Fig. 4c). It is also worth noting that four other cells, initially labelled as PS in the human 272 gastrula dataset, were also found to align to early PGCLCs and were reannotated accordingly. Together 273 these observations strongly suggest that the CS7 gastrula contains samples of PGCs at different stages 274 of specification and that our in vitro model captures the dynamics of this developmental trajectory at a 275 much finer resolution. Cross comparison of CS7 PGCs with PGCLCs from various other in vitro models 276 confirms a robust and conserved program of PGCLC-specification centred around the 277 SOX17/TFAP2C/PRDM1 network with consistent up-regulation of TFAP2A and other genes 278 (Supplementary Fig. 5).

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We quantified the dynamics of individual bifurcations by inferring lineage trajectories with Waddington-OT (Schiebinger *et al.*, 2019), an optimal transport-based approach that allowed us to infer 282 progenitor-progeny relationships between groups of cells statistically. By integrating these results with 283 reduced dimensional representations of our time-course data, such as UMAP, DM, or PCA, we sought 284 to identify the most likely earliest progenitors of PGC specification in our data. Using the ancestor-285 progeny relationships computed by WOT we inferred the broader lineages by first constructing a sparse 286 network of clusters (Supplementary Fig. 6a,b) which were further grouped using a community-detection 287 algorithm (see Methods). We assigned the inferred lineage identities to the single cells in these groups 288 using broad marker gene expression patterns. As an initial check, we overlaid these WOT-inferred 289 lineages onto our UMAP in Supplementary Fig. 6c, which demonstrated a good agreement with our 290 earlier annotation-based lineage assignments with a high degree of correlation to our earlier cluster and 291 marker-based annotations (Supplementary Fig. 6d). Using Waddington-OT inference, most terminal 292 cell fates were effectively traced to 24h, with some cell groups traced to earlier stages.

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294 Early mesoderm populations progressed from a PS-like state through a nascent-mesoderm-like state 295 (nMELC) expressing MESP1/2 and T to an emergent mesoderm-like state (eMELC), representing the 296 highest levels of MESP1/2 and downregulation of T (Supplementary Fig. 6e-f). Between 24 to 32h, a 297 PDGFRA positive population emerged, aligned to advanced mesoderm of the human gastrula (denoted 298 advanced mesoderm-like cells; aMELC), concomitant with the gradual loss of nMLC and eMLC 299 subpopulations. By ordering gene expression along a diffusion pseudotime analysis, we observed the 300 late up-regulation of several advanced mesoderm markers, HAND1, SNAI2 and GATA6 (Supplementary 301 Fig. 6e-g). As the earliest specified fate, nascent and emergent mesoderm cells express several genes 302 that may influence the balance of fates within the embryoid body, including BMP4, WNT5A, and CER1, 303 and extra-cellular matrix genes (see e.g., Supplementary Fig. 4).

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From 24h to 32h, a limited pool of *SOX17*-positive endoderm-like cells bifurcated from the PS-like subpopulation and showed sustained *NODAL* expression with subsequent upregulation of endoderm markers *FOXA1/2* (Supplementary Fig. 6h,i). Although the number of cells in this population appeared to be fewer than that of other cell lineages, it was nevertheless a conserved feature across in vitro models.

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Around the 18-hour mark, the earliest PGCLCs bifurcated from a progenitor population with strong up regulation of *SOX17*, *TFAP2C*, and *PRDM1* (see Supplementary Fig. 2g-h) and subsequent expression
 of *NANOS3*. PGCLCs also showed up-regulation of *WNT2* with early PGCLCs expressing *NODAL*

- 314 (Supplementary Fig. 4).
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Indeed, a comparison of PGCLC-precursor cells in high and low PGC-competence cell lines (Chen *et al.*, 2019) revealed *NODAL* to be differentially expressed, consistent with a recently observed role for *NODAL* in PGCLC specification (Jo *et al.*, 2021). Slightly later, at 24h, an AMLC branch also became

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319 evident, expressing TFAP2A and, at later time points, ISL1, a LIM/homeodomain transcription factor 320 protein recently identified as an amnion marker (Guo et al., 2020; Yang et al., 2021) (Supplementary 321 Fig. 2c-d). This AMLC branch shows an expression of WNT6 (Supplementary Fig. 4). We identified 322 differentially expressed genes along the separate AMLC and PGCLC lineages using the diffusion 323 pseudotime ordering of single cells (Fig. 4d; see Supplementary Materials). Within these pseudotime 324 trajectories, we observed that both AMLC and PGCLC showed early coordinated expression of 325 EOMES, MIXL1 and ZIC, together with rapid downregulation of SOX2. Moreover, we observed late 326 expression of VTCN1, GATA3, GATA2, ISL1 and HAND1 in AMLCs, while the PGCLC trajectory 327 showed late expression of PGC markers SOX17, PRDM1, TFAP2C, SOX15, KLF4, LIN28, and 328 *POU5F1*. Fig. 4e shows the divergent expression patterns of crucial TFs over pseudotime to trace their 329 rise and fall to AMLC versus PGCLC trajectories. We note an initial up-regulation of SOX17 in AMLC 330 and PGCLCs that is transient in AMLC but sustained in PGCLCs. Surprisingly, TFAP2A, which is 331 generally considered a trophoblast or amnion marker (Krendl et al., 2017; Zheng et al., 2019; Minn et 332 al., 2020), precedes SOX17 expression and is transiently co-expressed with SOX17 in the PGCLC 333 trajectory. While AMLCs maintain TFAP2A expression, there is downregulation in PGCLCs, which 334 was confirmed by immunofluorescence staining at the protein level (Fig. 4f). Staining of EBs for 335 TFAP2A and SOX17 confirmed their co-expression at early time points, whereas, in the 96h EB, 336 TFAP2A expression is exclusive to AMLC and SOX17 to PGCLCs and DELCs. These results, taken 337 together, highlight the complex dynamics of PGCLC specification within our model system and identify 338 several putative markers of specification. The most interesting was the early and transient expression 339 of TFAP2A in PGCLCs. TFAP2A is an early BMP response gene that shares the TF binding site with 340 TFAP2C (Krendl et al., 2017). Given that we previously found TFAP2 motifs around PGC-related 341 genes (Tang et al., 2022), and that the TFAP2 family can play complementary roles, it is interesting to 342 see if TFAP2A plays a role in PGCLC specification before the onset of TFAP2C expression.

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# 344 TFAP2A is the most upstream crucial regulator of PGC specification

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346 To determine whether the transient *TFAP2A* expression has a role in PGC fate, we induced PGCLCs 347 via PreME states using PSCs with a knockout mutation in TFAP2A and compared the outcome with the 348 parental PSC line (Krendl et al., 2017) (Fig. 5a). We observed a reduction in PGCLCs in TFAP2A 349 mutant cells compared to parental controls by FACS using antibodies for PGC-surface markers PDPN 350 and AP (2.78% vs 9.28%) (Fig. 5b). Quantification of four independent experiments showed a 351 consistent and statistically significant reduction in PGCLC specification in TFAP2A KO EBs (Fig. 5c) 352 confirmed by immunofluorescence staining of d4 EBs generated from TFAP2A knockout cells, 353 compared to parental control (Fig. 5d).

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To characterise the phenotype due to TFAP2A loss of function further, we generated 10X scRNA-seq
datasets for two-time points: 18h, just before the diversification of distinct lineages in embryoid bodies,
and at 96h, when terminal cell fates have been established. We integrated these time points with our
existing EB dataset containing all cell types for reference using Seurat. For this alignment, we generated
a new clustering visualised on a UMAP in Fig. 5e.
Embryoid bodies in parental lines showed the precise formation of a MELC expressing *PDGFRA*,
amnion expressing *VTCN1*, and PGCLCs expressing *NANOS3* by 96h (Fig. 5e), suggesting conserved

364 TFAP2A KO line by scRNA-seq, likely due to their limited cell numbers, immunofluorescence analysis 365 shows that rare SOX17, FOXA2 double-positive cells were present in the EB (Supplementary Fig. 7a). 366 On the other hand, in EBs with TFAP2A KO cells, PGCLC lineages were virtually absent (Fig. 5E, 367 Supplementary Fig. 7a-b), but aMELC and AMLC populations were present. While the TFAP2A KO 368 appeared to lack PGCLCs, we observed a new subpopulation of cells at 96h clustered alongside 369 pluripotent cells (Fig. 5e). This population, absent in the parental line and rare at the 18h mark in the

terminal behaviour with previous lines. While there were no detectable DELCs in either the parental or

- KO line, showed expression of *SOX2* and other pluripotency markers (hereafter referred to as SOX2+d4cells; Supplementary Fig. 7c).
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373 To help establish the authenticity of the other fates, we generated a cross-correlation heatmap 374 (Supplementary Fig. 7d). The SOX2+d4 cluster is most similar to PSCs in the reference population. 375 AMLCs in the KO cluster were highly similar to AMLCs in the parental line and the reference line, 376 with MELCs also showing consistency across all cell lines. Together these observations suggest no 377 significant effect of TFAP2A loss of function for MELCs or AMLCs specification. 378 Immunofluorescence analysis confirmed the presence of AMLCs (GATA3+ HAND1+), MELCs 379 (HAND1+) (Supplementary Fig. 7e), and DELC cells (SOX17+, FOXA2+) in TFAP2A KO in 380 TFAP2A KO EBs but with a minimal number of PGCLCs (SOX17+, OCT4+) (Supplementary Fig. 381 7a), confirming that TFAP2A had no significant effect on the other fates of the EB. We, therefore, 382 focused on PGCLCs and the SOX2+ population.

- 383
- Differential expression analysis of the SOX2+d4 population compared to parental-line PGCLCs showed that the SOX2+d4 cells expressed pluripotency and neural-plate factors, *ZIC2*, *ZIC5*, *SOX11*, *OTX2*, while PGCLCs showed expression of germ cell markers *SOX17*, *PRDM1*, *SOX15*, *ARID5B*, *TFCP2L1* and *VENTX* (Fig. 5g). We found upregulation of naïve markers of pluripotency and neuronal lineage-associated genes in the SOX2+d4 population compared to PSCs in the reference atlas; markers
- 389 included PRDM14, KLF4, KLF6, and TFAP2C, and neuro-related genes ZIC2, ZNF292, FOXN3,
- 390 POU3F1, SOX11, SOX4, ZIC5, and SALL3 (Supplementary Fig. 7f).
- 391

12

To validate our findings at the protein level, we performed immunofluorescence staining at d4 EBs and
found expression of *SOX2* in TFAP2A KO cells even after four days of cytokine exposure. There was
a rapid downregulation of *SOX2* upon BMP exposure (Supplementary Fig. 7c) in WT cells, which is
critical for efficient PGCLC specification(Lin *et al.*, 2014). SOX2+d4 cells also showed co-expression
of OCT4 and NANOG (Fig. 5h).
We investigated if TFAP2A could potentially target SOX2 for downregulation based on these results.
For this, we generated a stable dox-inducible TFAP2A PSCs line. Upon doxycycline induction of

400 *TFAP2A* in PSCs cultured in E8 medium by dox for two days (Fig. 5i), we observed a substantial 401 reduction in SOX2 levels after TFAP2A overexpression by immunofluorescence (Fig. 5j). *POU5F1* 

402 was also slightly reduced. Together, our results suggest that TFAP2A is a regulator of PGCLC fate and

- 403 may participate in the downregulation of SOX2 and other targets impeding PGCLC specification.
- 404

#### 405 **Discussion**

406

In vitro models have been of vital importance for unravelling the transcriptional network responsible
for human germ cell competence and specification (Teo *et al.*, 2011; Irie *et al.*, 2015; Sasaki *et al.*,
2015; Chen *et al.*, 2017; Kobayashi *et al.*, 2017; Kojima *et al.*, 2017, 2021; Pierson Smela *et al.*, 2019;
Sybirna *et al.*, 2020). In this study, we characterise in vitro models for the derivation of PGCLCs from
PSCs by highly resolved single-cell transcriptomics and comprehensive comparison to in vivo
references in human, non-human primates, and other in vitro models of gastrulation.

413

414 Notably, we found that PGC competent PreME cells exist transiently within a continuum of states 415 extending from PSCs to mesendoderm (ME). Our analysis showed that clusters enriched for PGC-416 competent populations present a particular signalling signature, characterised by active Nodal and WNT 417 signalling. There is low expression of BMP inhibitors (BAMBI and CER1) in competent cells compared 418 to the ME-dominated cluster with the highest levels in non-competent cluster 22, which likely impedes 419 PGC specification. BAMBI is a direct target of WNT signalling (Sekiya et al., 2004), while activation 420 of CER1 occurs via both WNT and Nodal signalling (Katoh and Katoh, 2006; Martyn, Brivanlou and 421 Siggia, 2019). PGCLC-competent clusters also show transient downregulation of OTX2 and higher 422 levels of NANOG compared to non-competent clusters, which we recently found is conducive to 423 transition to the PGCLC state (Tang et al., 2022). Concomitantly, there is an increase in the levels of 424 EOMES, which has a prominent role in human PGC-competence (Chen et al., 2017; Kojima et al., 425 2017), but further activation of mesoderm factors hinders PGC specification. The tight signalling axis, 426 transcription factor levels and intrinsic heterogeneity modulating competence are consistent with a

13

relatively small number (~100-200) of founder PGCs in vivo (Saitou, Barton and Surani, 2002;
Kobayashi *et al.*, 2017).

429

430 Specification of PGCLCs in vitro occurs within a 3D aggregate that consists of a hitherto poorly 431 characterized fraction of somatic components. Currently, PGCLCs can be induced in various 2D 432 aggregates but more efficiently in 3D embryoids, highlighting the importance of the structure, cell-cell 433 interactions or signalling from adjacent tissues (Minn et al., 2020, 2021). Here we have shown that 434 these somatic cells collectively represent those in the posterior region of the embryo during gastrulation. 435 Among the somatic cell types, we note the early formation of mesoderm-like cells, which display strong 436 expression of BMP, WNT, and ECM components that may be important for PGC-fate and potentially 437 play a similar role to that of extraembryonic mesoderm in the embryo, and endoderm-like cells that are 438 double positive for FOXA2/SOX17. Furthermore, we also observe the emergence of ISL1/VTCN1 439 expressing amnion cells, providing evidence that amnion formation continues from the posterior 440 epiblast during gastrulation, as recently suggested in a study on marmoset (Bergmann et al., 2022).

441

Mapping the cells to a 3D primate embryo showed that PSCs best correspond to the anterior region of the embryonic disc, while PreME cells shifted towards the posterior end. Conversely, cells within the newly formed embryoid body at 12h, which transcriptionally resemble a primitive streak, mapped best to the posterior end of the embryonic disc, with PGCLCs mapping to a *SOX17/TFAP2C/NANOS3* positive region at the boundary between the posterior-most epiblast and amnion.

447

448 The origin of human PGCs remains unresolved due to the inaccessibility of human embryos, but 449 bilaminar disc embryos from other species provide valuable information. In species such as the rabbit 450 and pig, PGCs originate from the posterior epiblast, but the amnion develops later, indicating that the 451 development of the amnion and PGCs in some cases are temporally unconnected. In humans and non-452 human primates, development of the amnion commences prior to PGC specification, but according to 453 our work and by others (Bergmann et al., 2022; Rostovskaya et al., 2022) amniotic cells continue to 454 emerge later from the posterior epiblast, co-incidentally with the specification of PGCs at the time of 455 primitive streak formation. In cynomolgus monkeys, the earliest PGCs have been reported in the 456 amnion, with the majority found later in the epiblast. One possibility is that these early PGCs may arise 457 from intermediate cells that are en route to the amnion but are but not fully committed as squamous 458 amniotic epithelium as observed in our data (Fig. 4c). To contribute to the founder PGC pool, PGCs 459 arising in the amnion would need to migrate against the continuing amnion growth. We posit that at this 460 stage of development in humans and non-human primates, amnion cells continue to be specified with 461 nascent PGCs arising at the posterior-most end of the epiblast during the early PS stage.

14

In our model, AMLC and PGCLC progenitors display early expression of *TFAP2A*, a pioneer factor previously associated with the amnion (Shao, Taniguchi, Townshend, *et al.*, 2017). Whilst there is subsequent downregulation of *TFAP2A* in PGCLCs, expression is sustained in the amnion. Surprisingly, the knockout of *TFAP2A* did not have a detectable effect on AMLCs, which merits further investigation, but notably resulted in an almost complete abrogation of PGCLCs.

468

469 In PGCLCs, TFAP2A is rapidly replaced by the expression of TFAP2C, suggesting otherwise mutually 470 exclusive expression after a brief window of co-expression. Interestingly, TFAP2A shares the same 471 transcription factor binding motif as TFAP2C (Krendl et al., 2017). TFAP2C is essential for PGC 472 development (Kojima et al., 2017) and acts as both an activator and a repressor during PGC 473 specification but it is not sufficient for PGC fate in the absence of cytokines (Kobayashi et al., 2017). 474 In the PGCLC pseudotime trajectories, we saw early upregulation of TFAP2A (12h), followed by 475 expression of SOX17 and TFAP2C (18h), and later, activation of PRDM1 (24-32h) by SOX17(Tang et 476 al., 2022) (Supplementary Fig. 8). In some instances, TFAP2A functions similarly to TFAP2C 477 (Hoffman et al., 2007; Li and Cornell, 2007). Our work suggests that TFAP2A expression is transient 478 but essential for initiating the PGC transcriptional network, and may directly or indirectly repress SOX2 479 and other factors.

480

481 TFAP2A KO EBs show an emergent population (SOX2+ d4 cells) found to align to pluripotent stem
482 cells, with the expression of the core pluripotency genes; *SOX2, POU5F1* and *NANOG*. Differential
483 gene expression between PSCs and SOX2+ d4 cells shows aberrant upregulation of naïve markers
484 *KLF4, TFAP2C* and *PRDM14* and genes associated with the neuronal lineage, including *ZNF292*,
485 *FOXN3, SALL3, ZIC2, POU3F1* in SOX2+ d4 cells.

486

There is rapid downregulation of SOX2 during human PGCLC-induction (Kobayashi *et al.*, 2017); indeed, sustained SOX2 expression prevents PGCLC specification due to elevated differentiation into the neuronal lineage (Lin *et al.*, 2014), which could in part explain the expression of related neuronal markers in the TFAP2A mutant cells. The combinatorial role of SOX17-OCT4 involved in human germ cell fate (Tang *et al.*, 2022) might benefit from a repression of SOX2 to favour the SOX17-OCT4 interaction on the compressed motif.

493

We provide insight into early human development with the transient emergence of the germ cell competent PreME cells in a model mimicking human gastrulation starting with PSC. Our study suggests continuing emergence of the amnion from the posterior epiblast at the time of PGC specification during early gastrulation; the amnion and PGC likely arise from highly similar progenitor exemplified by TFAP2A expression. The loss of function has a marked effect on PGC specification but without a detectable effect on the amnion. Accordingly, PGCs likely emerge from the posterior epiblast

15

500 predominantly, notwithstanding a sub-set in the early amnion (Fig. 6). Of great interest would be to test,

501 when possible, the predictions we make by direct observations in extended cultures of developing

- 502 human embryos.
- 503

# 504 Material and methods

505

# 506 Cell culture

H1 NANOS3-tdTomato PSC line was previously generated in the lab (Kobayashi et al., 2017). H9
parental and TFAP2A KO cells were kindly provided by Micha Drukker (Krendl et al., 2017). All cell
lines were confirmed as mycoplasma negative. PSCs were maintained on vitronectin-coated plates in
Essential 8 medium (Thermo Fisher Scientific) according to the manufacturer's protocol. Cells were
passaged every three to four days using 0.5 mM EDTA in PBS without breaking cell clumps.

For the 4i condition, undifferentiated PSC cells were maintained on irradiated mouse embryonic
fibroblasts (MEFs) (GlobalStem) in 4i medium (Irie et al., 2015). 4i were passaged every three to five
days using TrypLE Express (Gibco) quenching with MEF media and filtered with 50 μm cell filter

515 (PERTEC). ROCK inhibitor (10  $\mu$ M; Y-27632, TOCRIS Bioscience) was kept in the culture for 24 h 516 after passaging.

517 Mesendoderm induction was performed as reported in (Kobayashi et al., 2017). PSCs were dissociated

518 into single cells using TrypLE Express and seeded onto vitronectin coated plates at 500,000 cells per

519 well of 6-well plate and cultured in mesendoderm (ME) induction medium for 10 to 12 hours. ME

520 medium is based on aRB27

521 Primordial germ cells were induced as reported previously (Irie et al., 2015; Kobayashi et al., 2017). 522 For this PreME cells were disaggregated into single cell solution using TrypLE, then 4,000 cells per 523 well were seeded into ultra-low attachment 96-well plates (Corning Costar) in PGC induction medium. 524 Mesendoderm, PGCLC and definitive endoderm were induced from NANOS3-tdTomato reporter 525 PSCs as described before (Kobayashi et al., 2017) using the aRB27 basal medium, which was composed 526 of Advanced RPMI 1640 Medium (Thermo Fisher Scientific) supplemented with 1% B27 supplement 527 (Thermo Fisher Scientific), 0.1 mM NEAA, 100 U/ml penicillin, 0.1 mg/ml streptomycin, 2 mM L-528 glutamine. To induce mesendoderm, trypsinized hPSCs were seeded on vitronectin-coated dishes at 529 200,000 cells per well in a 12-well plates and cultured in mesendoderm induction medium for 12 530 (PreME) and 24 (ME) hours. Mesendoderm induction medium contained aRB27 medium supplemented 531 with 100 ng/ml activin A (Department of Biochemistry, University of Cambridge), 3 µM GSK3i 532 (Miltenyi Biotec) and 10 µM of ROCKi (Y-27632, Tocris bioscience). To induce definitive endoderm 533 from ME, mesendoderm induction medium was replaced with definitive endoderm induction medium 534 after washing with PBS once and cells were cultured for a further 2 days. Definitive endoderm induction 535 medium was composed of aRB27 medium supplemented with 100 ng/ ml activin A (Department of 536 Biochemistry) and 0.5 µM BMPi (LDN193189, Sigma).

#### 16

#### 537

To induce PGCLCs, PreME cells were trypsinized into single cells and harvested into Corning Costar
Ultra-Low attachment multiwell 96-well plate (Sigma) at 4,000 cells per well in hPGCLC induction
medium, which composed of aRB27 medium supplemented with 500 ng/ml BMP4,10 ng/ml human
LIF (Department of Biochemistry), 100 ng/ml SCF (R&D systems), 50 ng/ml EGF (R&D Systems), 10
µM ROCKi, and 0.25% (v/v) poly-vinyl alcohol (Sigma). Cells were cultured as floating aggregate for

- 543 2-4 days.
- 544

545 To collect PSCs, PreME, ME, DE, PGCLCs, cells were trypsinized with 0.25% trypsin/EDTA at 37 °C

546 for 5-15 min. DE was stained with PerCP-Cy5.5 conjugated anti-CXCR4 antibody (Biolegend). Cell

- 547 suspension was subjected to FACS by SH800Z Cell Sorter (Sony) and analyzed by FlowJo software.
- 548

# 549 Collection of human PGCs from human embryos

550

Human embryonic tissues were used under permission from NHS Research Ethical Committee, UK (REC Number: 96/085). Human embryonic samples were collected following medical or surgical termination of pregnancy carried out at Addenbrooke's Hospital, Cambridge, UK with full consent from patients. Crown-rump length, anatomical features, including limb and digit development, was used to determine developmental stage of human embryos with reference to Carnegie staging (CS). The sex of embryos were determined by sex determination PCR as previously described (Bryja and Konečný, 2003).

558

559 Human embryonic genital ridges from individual embryos (wk7) were dissected in PBS and separated 560 from surrounding mesonephric tissues. The embryonic tissues were dissociated with Collagenase IV 561 (2.6 mg/ml) (Sigma, C5138) and DNase I (10 U/ml) in DMEM-F/12 (Gibco) at 37°C for 15-30 minutes 562 (depending on tissue size). Tissues were pipette up and down for five times every 10 minutes to facilitate 563 dissociation into single cell suspension. After that, samples were diluted with 1 ml FACS medium (PBS 564 with 3% fetal calf serum & 5 mM EDTA) and centrifuged at 500 xg for 5 minutes. Cell pellet was 565 suspended with FACS medium and incubated with 5 µl of Alexa Fluor 488-conjugated anti-alkaline 566 phosphatase (BD Pharmingen, 561495) and 5 µl of APC-conjugated anti-c-KIT (Invitrogen, CD11705) 567 antibodies for 20 minutes at room temperature with rotation at 10 revolutions per minutes (rpm) in dark. 568 Cell suspension was then diluted in 1 ml FACS medium and centrifuged at 500 xg for 5 minutes. After 569 removing the supernatant, the cell pellet was resuspended in FACS medium and passed through a 35 570 µm cell strainer. FACS was performed with SH800Z Cell Sorter (Sony) and FACS plots were generated 571 by FlowJo software.

572

#### 573 Fluorescence-activated cell sorting (FACS)

17

574

575 PSCs, 4i, PreME and ME cells were harvested using TrypLE (GIBCO) at 37°C for 2-3 min. Embryoid 576 bodies were collected and dissociated into single cells using Trypsin-EDTA solution 0.25% at 37°C for 577 5 to 15 min. Dissociated cells were washed and resuspended in the FACS buffer (PBS 3% FCS). DE 578 samples were stained with PerCP-Cy5.5 conjugated anti-CXCR4 antibody (Biolegend) for 1h on ice. 579 Samples were washed with PBS, stained with DAPI (1:10,000) and sorted on a SONY SH800 sorter. 580

581 Human embryonic genital ridge or mesonephros from a week 7.0 male embryo were collected in 582 dissection medium (DMEM (Gibco), 10% FCS, 1 mM sodium pyruvate (Sigma)). Embryonic tissues 583 were dissociated with 300 µL collagenase IV (2.6 mg/mL in DMEM-F/12) supplemented with DNaseI 584 (10U/mL) per genital ridge and incubated for 10 minutes at 37°C with mixing by pipetting up and down. 585 Then, cells were washed with 1 mL FACS buffer (PBS with 3% FCS and 5 mM EDTA). Resuspended 586 with 75 µL FACS buffer and stained with 0.5 µL alexa Fluor 488-conjugated anti-alkaline phosphatase 587 (BD Pharmingen, 561495) and 25 µL of PerCP- Cy5.5-conjugated anti-CD117 (BD Pharmingen 588 333950) for 15 minutes at room temperature. Samples were washed with PBS and sorted on a SONY 589 SH800 cytometer. Flow cytometry data was analysed on FlowJo v10 (FlowJo LLC).

590

# 591 Immunofluorescence

592

Embryoid bodies (EBs) were fixed in 4% PFA for 2h at 4 °C and embedded in O.C.T. compound (Cellpath) for frozen sections. Each sample was incubated with primary antibodies for 1–2 h at room temperature or overnight at 4°C and then with fluorescent-conjugated secondary antibodies and DAPI (Sigma) for 1 h at room temperature. Samples were then imaged under a Leica SP8 upright or inverted scanning confocal microscope.

598 Cells were cultured on ibidi µ-Slide and fixed in 4% PFA for 30 minutes at 4°C. Embryoid bodies were 599 fixed in 4% PFA for 2 hours at 4°C and embedded in OCT compound for frozen sections. The samples 600 were incubated with primary antibodies overnight at 4°C and subsequently with fluorescence-601 conjugated secondary antibodies (Thermo Fisher Scientific) and DAPI for 1 hour at RT. The primary 602 antibodies used are: anti-GFP (abcam, ab13970), anti-PRDM1 (Cell Signaling Technology, 9115), anti-603 SOX17 (R&D, AF1924), anti-TFAP2C (Santa Cruz Biotechnology, sc-8977), and anti-OCT4 (BD 604 Biosciences, 611203). Samples were imaged under Leica SP8 upright or inverted scanning confocal 605 microscope.

606

#### 607 10X genomics

608

For each stage, 5,000 cells were sorted into an eppendorf tube containing PBS with 0.04%
weight/volume BSA (400 μg/mL). Samples collected are listed in table 2.6. During sorting, dead cells,

18

611	debris and doublets were gated out. Sorted cells were directly taken for 10x processing at Cancer
612	Research UK, Cambridge Institute and loaded into the 10x-Genomics Chromium using the single cell
613	3' reagents kit v2. Libraries were prepared as per the manufacturer's instructions and pooled for
614	sequencing so that all lines would include all samples. Libraries were sequenced, aiming at a minimum
615	coverage of 50,000 raw reads per cell, on an Illumina HiSeq 4000 (paired-end; read 1: 26 cycles; i7
616	index: 8 cycles, i5 index: 0 cycles; read 2: 98 cycles).
617	
618	Bioinformatics
619	
620	10X RNA sequencing processing
621	
622	Multiplexed single-cell libraries were processed using the 10X Genomics cell ranger pipeline. Reads
623	were aligned to a reference genome (Homo sapiens GrCh38) using STAR (Dobin et al., 2013), and
624	quantification of genes against an annotation reference (based on Ensembl GrCh38 v90).
625	
626	Analysis
627	
628	Initial analysis of our data was done using Seurat (v3.1.4) (Stuart et al., 2019). Count data was
629	normalised and scaled using NormalizeData based on log counts per 10000 (logCP10k) and scaled
630	using ScaleData. Clusters were generated using FindCluster with resolution of 0.1. Nearest neighbour
631	graphs and UMAP plots were calculated using the first 20 PCs.
632	
633	Heatmaps of gene expression were generated based on row-scaled values using pheatmap (v.1.0.12)
634	with cross-correlations calculated based on Pearson's correlation and visualised using pheatmap.
635	
636	Integrative analysis
637	
638	Individual datasets were first curated to remove pre-implantation and extraembryonic tissues. Datasets
639	were then integrated based on logCP10k using FindIntegrationMarkers with 5000 integration features
640	and k.filter=50. Data was integrated based on CCA with 5000 features and using the first 20 PCs. Joint
641	clustering was generated based on the integration-corrected gene expression matrices using the
642	FindClusters function with complexity parameter uniformly incremented form 0.1-0.9 in steps of 0.1.
643	For visualisation purposes we used parameter of 0.9 for the figures within the paper.
644	
645	For initially establishing cell fates expression of key marker genes were plotted as a heatmap using
646	pheatmap. To establish veracity of cell types between datasets, a scatter plot of differential expression

647 was used with the x-axis showing logFC of a specific cluster vs a reference cell type/cluster (e.g., cluster

19

648	0 vs PSCs) with the y-axis showing the same comparison (cluster 0 vs PSCs) in the second dataset.
649	Genes in the top right and bottom left quadrants represented conserved changes between the two
650	datasets, whilst genes to the top left or bottom right represented dataset specific changes.
651	
652	As a preliminary visualisation of individual bifurcations, diffusion maps were generated for selected
653	sets of subclusters using destiny (v2.12.0) (Angerer et al., 2016) based on integration-corrected
654	expression matrices.
655	
656	Differential expression analysis
657	
658	Unless otherwise indicated, differential expression between two groups was done in Seurat using MAST
659	(Finak et al., 2015). For volcano plots, genes were filtered to show genes with adjusted p-values <0.05
660	with a >1.2 FC.
661	
662	Mapping of cells from CS7 gastrula to embryoid bodies
663	
664	Carnegie stage 7 human gastrula annotations were projected onto our EB dataset based on statistically
665	enriched proximity in nearest neighbour graphs. Specifically, the aligned datasets were subsetted on the
666	human CS7 gastrula and EB dataset and used to calculate a KNN graph (using the FindNeighbours
667	function). For each cell within our EB dataset, the enrichment of individual CS7 gastrula annotations
668	was calculated using a hypergeometric test, and final annotations assigned based on adjusted p-values.
669	Cells that showed no significant overlap in KNN graphs were not assigned a lineage.
670	
671	Mapping of cells to the CS6 marmoset embryo
672	
673	Cells within our EB were mapped to the marmoset embryo based on proximity in KNN-graphs in the
674	CCA aligned datasets. Aligned datasets were first subsetted on the marmoset dataset and EB dataset.
675	For a cell, <i>j</i> , in the EB dataset, we calculated the KNN from the CS6 embryo, with positions at positions
676	$\{r_1, r_2,, r_k: r_i \in \mathbb{R}^3\}$ , and calculated the shared nearest neighbour (SNN) vector $\boldsymbol{\theta}^{(j)} =$
677	$\{\theta_1, \theta_2, \dots, \theta_K\}$ . Weights were normalised $\widehat{\theta}^{(j)} = \theta^{(j)}/c$ , $c = \sum_i \theta_i$ and a projection of cell <i>j</i> calculated
678	as: $\mathbf{R} = \sum_i \mathbf{r}_i \hat{\theta}_i$ , where $\mathbf{r}_j \in \mathbb{R}^3$ denotes a 3-dimensional position vector of marmoset cell <i>j</i> . After
679	mapping of individual cells, the density of specific groups e.g., PGCLCs (cluster 5 and 6), AMELC
680	(clusters 7, 9 and 10), basal (cluster 1), was calculated using the MATLAB function mvksdensity.
681	
682	Doublet detection

20

To minimise doublets in our analyses, we limited the number of cells loaded into each chip, with each sample capturing around 1000-2000 cells. Potential doublets were identified computationally for each individual sample using the R package DoubletFinder(McGinnis, Murrow and Gartner, 2019). For samples with ~1000 captured cells we assumed a doublet rate of 1%, and for samples with ~2000 cells we assumed a 2% doublet rate. No cluster analysed in this paper was found to contain a high level of doublets.

690

#### 691 Waddington Optimal Transport analysis

692

693 Highly variable genes were computed across all single PSCs, PreME and EB cells, and used as input to 694 PCA, with the first 50 PCs computed using irlba. Cells were assigned to clusters as described above, 695 which were used as the basis for WOT. Transport maps were computed with parameters ( $\lambda 1=1$ ,  $\lambda 2=50$ , 696  $\varepsilon$ =0.01) between all pairs of time points using the PSCs as 0hours, PreME as 12 hours, and all subsequent time points as 12+ti for  $i \in \{1, 2, ..., T\}$  and  $T = \{12, 18, 24, 32, 40, 48, 96\}$ . Ancestor 697 698 contributions to populations at subsequent time points were estimated from these transport maps using 699 the OT trajectory command-line interface (CLI) function. Cell mass contributions between clusters 700 across time points were concatenated into a cluster: timepoint X cluster: timepoint matrix, where the 701 rows denote the contribution of clusteri timepointi to clusteri timepointi+1. A power threshold (p=30) 702 was used to enforce sparsity on this matrix with values  $\leq 0.1$  censored to 0. This sparse matrix was then 703 used as a weighted adjacency matrix to compute a directed KNN graph (k=5), as shown in 704 Supplementary Figure 6. Meta-clusters were defined on this graph using the Walktrap community 705 detection algorithm implemented in igraph, which were annotated based on the mean expression level 706 of single-cells that contribute to each original cluster (Supplementary Figure 6). These annotations were 707 then mapped back onto the original constituent single-cells based on their cluster identity.

708

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710

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2	Τ.

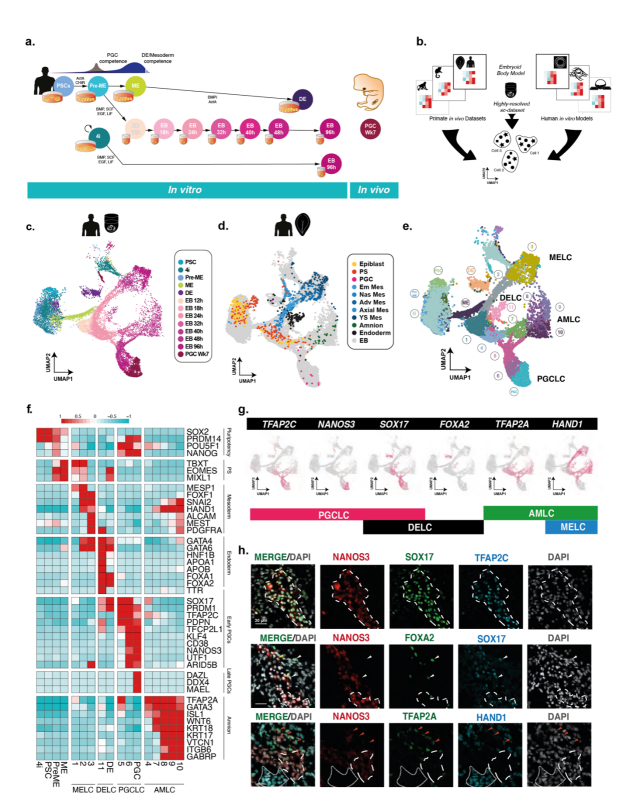
721	
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728	
729	Availability of materials
730	
731	Any enquiries on reagents and cell lines can be directed to (a.surani@gurdon.cam.ac.uk). Plasmids
732	generated in this study will be made freely available upon request. Modified human embryonic stem
733	cell lines generated in this study will be made available on request upon completion of a Materials
734	Transfer Agreement.
735	Single cell RNA-seq (10X) data has been deposited at ArrayExpress under accession numbers E-
736	MTAB-11283 and E-MTAB-11305. Code for repeating analyses will be available via a GitHub
737	repository https://github.com/cap76/PGCLC.
738	
739	Author contributions
740	
741	ACV, CAP, MAS wrote the manuscript with input from all authors. ACV, CAP, MDM designed
742	experiments and performed analysis. ACV generated human data. SB and ES generated marmoset data.
743	ACV, WWCT, TK, FCKW performed experiments. MAS, JCM, and TEB supervised.
744	
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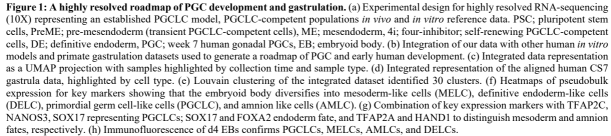
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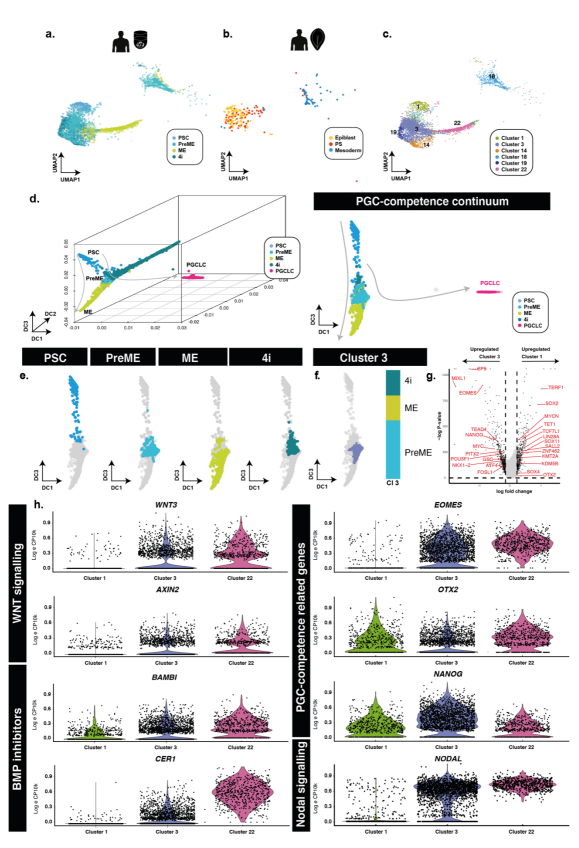
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**Figure 2: PGCLC competent populations form a continuum of states.** (a) Aligned UMAP representations of pluripotent and PGCLCcompetent populations, alongside (b), human *in vivo* samples shows that PSCs align best to pluripotent epiblast cells while competent (PreME and 4i) align to both epiblast-like and primitive-streak-like populations. (c) Sub clustering of competent and non-competent cells identified six main populations; a PSC-dominated \*cluster 1, a mesendoderm-dominated cluster 22, and competence-dominant clusters 3, 14, 19. (d) Diffusion map representations shows samples lie along a continuum of overlapping states. (e-f) The fractional makeup of competencedominated subclusters 3 showed almost equal contribution from 4i and PreME cells. (g) Differential expression of competence-dominated subcluster 3 versus PSC-dominant, non-competent subcluster 1 identifies putative regulators of competence. (h) Violin plots of putative competence related genes and markers for WNT and BMP signalling reveal heterogeneous signalling response

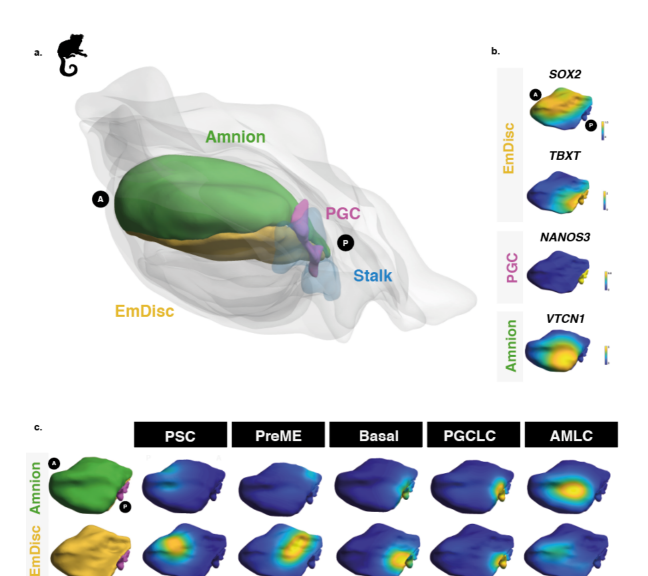
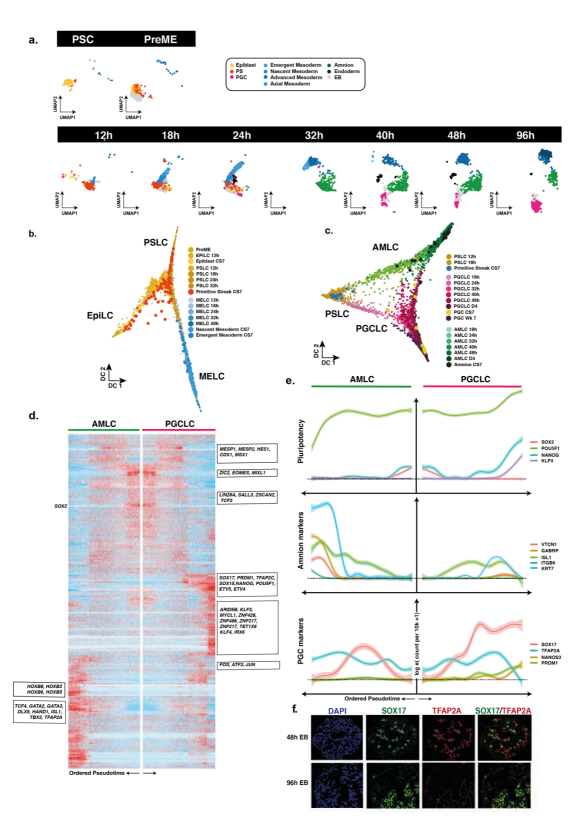
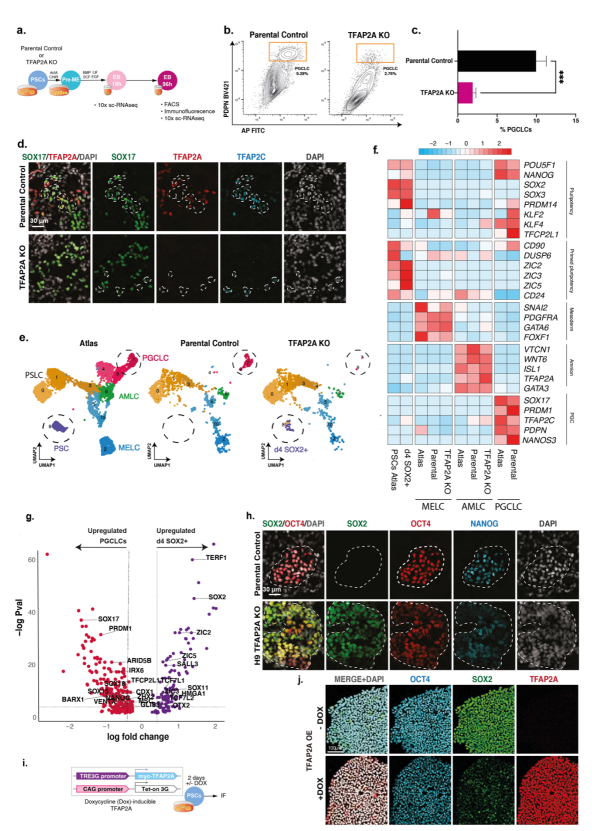


Figure 3: Spatial mapping of embryoid body models to gastrulating marmoset embryos reveals posterior bias. (a) Spatially resolved marmoset embryos at CS6 with the embryonic disc in yellow, amnion in green, PGCs in pink, and stalk in blue. Extraembryonic tissues are shown in grey. (b) Expression analysis in the marmoset embryo shows the anterior embryonic disc is SOX2 and the posterior, T positive, respectively. Specified PGCs with NANOS3 expression, amnion with partial VTCN1 expression. (c) Mapping *in vitro* cells shows PSCs map best to the anterior embryonic disc. Competent populations show a distinct posterior bias, with PGCLCs showing strong localisation to posterior-most PGC region and AMLCs mapping to the amnion.

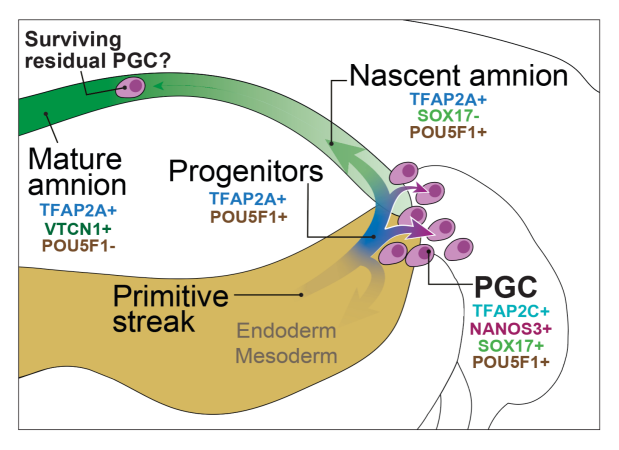


**Figure 4: Resolving the dynamics of bifurcations in embryoid bodies.** (a) Visualisation of data separated by sample time with cells annotated by transfer of labels from the human CS7 gastrula<sup>1</sup>; representation suggests EBs develops first through a primitive streak-like stage, early emergence of mesoderm-like cells and primordial germ cell-like cells, followed by amnion-like cells. (b) Diffusion map representation of specific clusters reveals strong bifurcation of mesoderm from the PS-like progenitors, with the remaining PS-like cells destined for other lineages. (c) Diffusion map representation of AMLC and PGCLCs shows bifurcation from common progenitor cells, with the continued association until 48h. Superimposition of cells from the CS7 gastrula labelled as PS, amnion or PGCs shows early alignment of hPGCs to PGCLCs. (d) WOT analysis to infer progenitor-descendent relationships, identifying bifurcations of individual lineages. Heatmap representants differentially expressed genes between AMLC and PGCLC ordered by pseudotime. (e) Line plot representations of essential genes ordered by pseudotime shows early up-regulation of TFAP2A in both PGCLC and AMLCs, which is sustained in AMLC. (f) IF shows TFAP2A in early PGCLCs at 48h (SOX17/TFAP2A double-positive) is lost by 96h.

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**Figure 5: TFAP2A is a regulator of PGCLC fate.** (a) Testing the role of TFAP2A in PGC specification. (b) FACS plot reveals % PGCLCs in TFAP2A KO EBs and WT parental control. (c) Immunofluorescence shows co-expression for SOX17, TFAP2A and TFAP2C in d4 EB. (d) Quantification of PGCLC (%) in WT and TFAP2A KO. (e) Aligned UMAPs for the reference data versus parental control and H9 TFAP2A KO. (f) Row-normalised gene expression demonstrate consistent expression in AMLC and MELC in the TFAP2A KO line. D4 SOX2+ cells shows expression of pluripotency genes. (g) Immunofluorescence of d4 parental EBs shows OCT4 NANOG double-positive cells (PGCLCs) but not in TFAP2A KO EBs; instead, there are OCT4, NANOG, and SOX2 triple positive cells. (h) Volcano plot for differentially expressed genes between the d4 SOX2+ cluster in TFAP2A KO vs PGCLCs in parental control) (i) Testing the role of TFAP2A overexpression in PSCs. (j) Immunofluorescence for OCT4, SOX2, TFAP2A in PSCs.



**Figure 6:** A unifying model of PGC specification in bilaminar disc embryos. PGCs are specified from *TFAP2A*-positive progenitors at the posterior end of the embryonic disc, which also give rise to nascent amnion. PGCs in the amnion specified at an earlier stage might only contribute to the founder PGC pool if they can migrate against the flow of nascent amnion expansion.